

Computational Physics, an Introduction

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So, ultimately, in order to understand nature it may be necessary to have a deeper understanding of mathematical relationships. But the real reason is that the subject is enjoyable, and although we humans cut nature up in different ways, and we have different courses in different departments, such compartmentalization is really artificial, and we should take our intellectual pleasures where we find them. *Richard Feynman, The Laws of Thermodynamics.*

Why a preface you may ask? Isn't that just a mere exposition of a *raison d'être* of an author's choice of material, preferences, biases, teaching philosophy etc.? To a large extent I can answer in the affirmative to that. A preface ought to be personal. Indeed, what you will see in the various chapters of these notes represents how I perceive computational physics should be taught.

This set of lecture notes serves the scope of presenting to you and train you in an algorithmic approach to problems in the sciences, represented here by the unity of three disciplines, physics, mathematics and informatics. This trinity outlines the emerging field of computational physics.

Our insight in a physical system, combined with numerical mathematics gives us the rules for setting up an algorithm, viz. a set of rules for solving a particular problem. Our understanding of the physical system under study is obviously gauged by the natural laws at play, the initial conditions, boundary conditions and other external constraints which influence the given system. Having spelled out the physics, for example in the form of a set of coupled partial differential equations, we need efficient numerical methods in order to set up the final algorithm. This algorithm is in turn coded into a computer program and executed on available computing facilities. To develop such an algorithmic approach, you will be exposed to several physics cases, spanning from the classical pendulum to quantum mechanical systems. We will also present some of the most popular algorithms from numerical mathematics used to solve a plethora of problems in the sciences. Finally we will codify these algorithms using some of the most widely used programming languages, presently C, C++ and Fortran and its most recent standard Fortran 2008¹. However, a high-level and fully object-oriented language like Python is now emerging as a good alternative although C++ and Fortran still outperform Python when it comes to computational speed. In this text we offer an approach where one can write all programs in C/C++ or Fortran. We will also show you how to develop large programs in Python interfacing C++ and/or Fortran functions for those parts of the program which are CPU intensive. Such an approach allows you to structure the flow of data in a high-level language like Python while tasks of a mere repetitive and CPU intensive nature are left to low-level languages like C++ or Fortran. Python allows you also to smoothly interface your program with other software, such as plotting programs or operating

¹Throughout this text we refer to Fortran 2008 as Fortran, implying the latest standard.

system instructions. A typical Python program you may end up writing contains everything from compiling and running your codes to preparing the body of a file for writing up your report.

Computer simulations are nowadays an integral part of contemporary basic and applied research in the sciences. Computation is becoming as important as theory and experiment. In physics, computational physics, theoretical physics and experimental physics are all equally important in our daily research and studies of physical systems. Physics is the unity of theory, experiment and computation². Moreover, the ability "to compute" forms part of the essential repertoire of research scientists. Several new fields within computational science have emerged and strengthened their positions in the last years, such as computational materials science, bioinformatics, computational mathematics and mechanics, computational chemistry and physics and so forth, just to mention a few. These fields underscore the importance of simulations as a means to gain novel insights into physical systems, especially for those cases where no analytical solutions can be found or an experiment is too complicated or expensive to carry out. To be able to simulate large quantal systems with many degrees of freedom such as strongly interacting electrons in a quantum dot will be of great importance for future directions in novel fields like nano-technology. This ability often combines knowledge from many different subjects, in our case essentially from the physical sciences, numerical mathematics, computing languages, topics from high-performance computing and some knowledge of computers.

In 1999, when I started this course at the department of physics in Oslo, computational physics and computational science in general were still perceived by the majority of physicists and scientists as topics dealing with just mere tools and number crunching, and not as subjects of their own. The computational background of most students enlisting for the course on computational physics could span from dedicated hackers and computer freaks to people who basically had never used a PC. The majority of undergraduate and graduate students had a very rudimentary knowledge of computational techniques and methods. Questions like 'do you know of better methods for numerical integration than the trapezoidal rule' were not uncommon. I do happen to know of colleagues who applied for time at a supercomputing centre because they needed to invert matrices of the size of $10^4 \times 10^4$ since they

²We mentioned previously the trinity of physics, mathematics and informatics. Viewing physics as the trinity of theory, experiment and simulations is yet another example. It is obviously tempting to go beyond the sciences. History shows that trinities, trinities and for example triple deities permeate the Indo-European cultures (and probably all human cultures), from the ancient Celts and Hindus to modern days. The ancient Celts revered many such trinities, their world was divided into earth, sea and air, nature was divided in animal, vegetable and mineral and the cardinal colours were red, yellow and blue, just to mention a few. As a curious digression, it was a Gaulish Celt, Hilary, philosopher and bishop of Poitiers (AD 315-367) in his work *De Trinitate* who formulated the Holy Trinity concept of Christianity, perhaps in order to accommodate millennia of human divination practice.

were using the trapezoidal rule to compute integrals. With Gaussian quadrature this dimensionality was easily reduced to matrix problems of the size of $10^2 \times 10^2$, with much better precision.

More than a decade later most students have now been exposed to a fairly uniform introduction to computers, basic programming skills and use of numerical exercises. Practically every undergraduate student in physics has now made a Matlab or Maple simulation of for example the pendulum, with or without chaotic motion. Nowadays most of you are familiar, through various undergraduate courses in physics and mathematics, with interpreted languages such as Maple, Matlab and/or Mathematica. In addition, the interest in scripting languages such as Python or Perl has increased considerably in recent years. The modern programmer would typically combine several tools, computing environments and programming languages. A typical example is the following. Suppose you are working on a project which demands extensive visualizations of the results. To obtain these results, that is to solve a physics problems like obtaining the density profile of a Bose-Einstein condensate, you need however a program which is fairly fast when computational speed matters. In this case you would most likely write a high-performance computing program using Monte Carlo methods in languages which are tailored for that. These are represented by programming languages like Fortran and C++. However, to visualize the results you would find interpreted languages like Matlab or scripting languages like Python extremely suitable for your tasks. You will therefore end up writing for example a script in Matlab which calls a Fortran or C++ program where the number crunching is done and then visualize the results of say a wave equation solver via Matlab's large library of visualization tools. Alternatively, you could organize everything into a Python or Perl script which does everything for you, calls the Fortran and/or C++ programs and performs the visualization in Matlab or Python. Used correctly, these tools, spanning from scripting languages to high-performance computing languages and visualization programs, speed up your capability to solve complicated problems. Being multilingual is thus an advantage which not only applies to our globalized modern society but to computing environments as well. This text shows you how to use C++ and Fortran as programming languages.

There is however more to the picture than meets the eye. Although interpreted languages like Matlab, Mathematica and Maple allow you nowadays to solve very complicated problems, and high-level languages like Python can be used to solve computational problems, computational speed and the capability to write an efficient code are topics which still do matter. To this end, the majority of scientists still use languages like C++ and Fortran to solve scientific problems. When you embark on a master or PhD thesis, you will most likely meet these high-performance computing languages. This course emphasizes thus the use of programming languages like Fortran, Python and C++ instead of interpreted ones like Matlab or Maple. You

should however note that there are still large differences in computer time between for example numerical Python and a corresponding C++ program for many numerical applications in the physical sciences, with a code in C++ or Fortran being the fastest.

Computational speed is not the only reason for this choice of programming languages. Another important reason is that we feel that at a certain stage one needs to have some insights into the algorithm used, its stability conditions, possible pitfalls like loss of precision, ranges of applicability, the possibility to improve the algorithm and tailor it to special purposes etc etc. One of our major aims here is to present to you what we would dub 'the algorithmic approach', a set of rules for doing mathematics or a precise description of how to solve a problem. To device an algorithm and thereafter write a code for solving physics problems is a marvelous way of gaining insight into complicated physical systems. The algorithm you end up writing reflects in essentially all cases your own understanding of the physics and the mathematics (the way you express yourself) of the problem. We do therefore devote quite some space to the algorithms behind various functions presented in the text. Especially, insight into how errors propagate and how to avoid them is a topic we would like you to pay special attention to. Only then can you avoid problems like underflow, overflow and loss of precision. Such a control is not always achievable with interpreted languages and canned functions where the underlying algorithm and/or code is not easily accessible. Although we will at various stages recommend the use of library routines for say linear algebra³, our belief is that one should understand what the given function does, at least to have a mere idea. With such a starting point, we strongly believe that it can be easier to develop more complicated programs on your own using Fortran, C++ or Python.

We have several other aims as well, namely:

- We would like to give you an opportunity to gain a deeper understanding of the physics you have learned in other courses. In most courses one is normally confronted with simple systems which provide exact solutions and mimic to a certain extent the realistic cases. Many are however the comments like 'why can't we do something else than the particle in a box potential?'. In several of the projects we hope to present some more 'realistic' cases to solve by various numerical methods. This also means that we wish to give examples of how physics can be applied in a much broader context than it is discussed in the traditional physics undergraduate curriculum.
- To encourage you to "discover" physics in a way similar to how researchers learn in the context of research.

³Such library functions are often tailored to a given machine's architecture and should accordingly run faster than user provided ones.

- Hopefully also to introduce numerical methods and new areas of physics that can be studied with the methods discussed.
- To teach structured programming in the context of doing science.
- The projects we propose are meant to mimic to a certain extent the situation encountered during a thesis or project work. You will typically have at your disposal 2-3 weeks to solve numerically a given project. In so doing you may need to do a literature study as well. Finally, we would like you to write a report for every project.

Our overall goal is to encourage you to learn about science through experience and by asking questions. Our objective is always understanding and the purpose of computing is further insight, not mere numbers! Simulations can often be considered as experiments. Rerunning a simulation need not be as costly as rerunning an experiment.

Needless to say, these lecture notes are upgraded continuously, from typos to new input. And we do always benefit from your comments, suggestions and ideas for making these notes better. It's through the scientific discourse and critics we advance. Moreover, I have benefitted immensely from many discussions with fellow colleagues and students. In particular I must mention Hans Petter Langtangen, Anders Malthe-Sørenssen, Knut Mørken and Øyvind Ryan, whose input during the last fifteen years has considerably improved these lecture notes. Furthermore, the time we have spent and keep spending together on the Computing in Science Education project at the University, is just marvelous. Thanks so much. Concerning the Computing in Science Education initiative, you can read more at <http://www.mn.uio.no/english/about/collaboration/cse/>.

Finally, I would like to add a petit note on referencing. These notes have evolved over many years and the idea is that they should end up in the format of a web-based learning environment for doing computational science. It will be fully free and hopefully represent a much more efficient way of conveying teaching material than traditional textbooks. I have not yet settled on a specific format, so any input is welcome. At present however, it is very easy for me to upgrade and improve the material on say a yearly basis, from simple typos to adding new material. When accessing the web page of the course, you will have noticed that you can obtain all source files for the programs discussed in the text. Many people have thus written to me about how they should properly reference this material and whether they can freely use it. My answer is rather simple. You are encouraged to use these codes, modify them, include them in publications, thesis work, your lectures etc. As long as your use is part of the dialectics of science you can use this material freely. However, since many weekends have elapsed in writing several of these programs, testing them, sweating over bugs, swearing in front of a `f*%@?%g` code which didn't compile

properly ten minutes before monday morning's eight o'clock lecture etc etc, I would dearly appreciate in case you find these codes of any use, to reference them properly. That can be done in a simple way, refer to M. Hjorth-Jensen, *Computational Physics*, University of Oslo (2013). The weblink to the course should also be included. Hope it is not too much to ask for. Enjoy!

Symbols

α	Temperature coefficient of linear expansion (K^{-1})
β	Temperature coefficient of volume expansion (K^{-1})
γ	Ratio of heat capacities
ε	Permittivity
κ	Dielectric constant
λ	Wavelength (m)
ρ	Density (kg/m^3)
B	Magnetic field (T)
C	Molar heat capacity ($\text{JKg}^{-1}\text{K}^{-1}$)
f	Frequency
k	Thermal conductivity ($\text{Wm}^{-1}\text{K}^{-1}$)
R	Ideal gas constant ($8.31 \text{ Jmol}^{-1}\text{K}^{-1}$)

Part I

Introduction to programming and numerical methods

The first part of this text aims at giving an introduction to basic C++ and Fortran programming, including numerical methods for computing integrals, finding roots of functions and numerical interpolation and extrapolation. It serves also the aim of introducing the first examples on parallelization of codes for numerical integration.

Chapter 1

Introduction

In the physical sciences we often encounter problems of evaluating various properties of a given function $f(x)$. Typical operations are differentiation, integration and finding the roots of $f(x)$. In most cases we do not have an analytical expression for the function $f(x)$ and we cannot derive explicit formulae for derivatives etc. Even if an analytical expression is available, the evaluation of certain operations on $f(x)$ are so difficult that we need to resort to a numerical evaluation. More frequently, $f(x)$ is the result of complicated numerical operations and is thus known only at a set of discrete points and needs to be approximated by some numerical methods in order to obtain derivatives, etc etc.

The aim of these lecture notes is to give you an introduction to selected numerical methods which are encountered in the physical sciences. Several examples, with varying degrees of complexity, will be used in order to illustrate the application of these methods.

The text gives a survey over some of the most used methods in computational physics and each chapter ends with one or more applications to realistic systems, from the structure of a neutron star to the description of quantum mechanical systems through Monte-Carlo methods. Among the algorithms we discuss, are some of the top algorithms in computational science. In recent surveys by Dongarra and Sullivan [3] and Cipra [2], the list over the ten top algorithms of the 20th century include

1. The Monte Carlo method or Metropolis algorithm, devised by John von Neumann, Stanislaw Ulam, and Nicholas Metropolis, discussed in chapters ??-??.
2. The simplex method of linear programming, developed by George Dantzig.
3. Krylov Subspace Iteration method for large eigenvalue problems in particular, developed by Magnus Hestenes, Eduard Stiefel, and Cornelius Lanczos, discussed in chapter ??.

4. The Householder matrix decomposition, developed by Alston Householder and discussed in chapter ??.
5. The Fortran compiler, developed by a team lead by John Backus, codes used throughout this text.
6. The QR algorithm for eigenvalue calculation, developed by Joe Francis, discussed in chapter ??
7. The Quicksort algorithm, developed by Anthony Hoare.
8. Fast Fourier Transform, developed by James Cooley and John Tukey.
9. The Integer Relation Detection Algorithm, developed by Helaman Ferguson and Rodney
10. The fast Multipole algorithm, developed by Leslie Greengard and Vladimir Rokhlin; (to calculate gravitational forces in an N-body problem normally requires N^2 calculations. The fast multipole method uses order N calculations, by approximating the effects of groups of distant particles using multipole expansions)

The topics we cover start with an introduction to C++ and Fortran programming (with digressions to Python as well) combining it with a discussion on numerical precision, a point we feel is often neglected in computational science. This chapter serves also as input to our discussion on numerical derivation in chapter ??. In that chapter we introduce several programming concepts such as dynamical memory allocation and call by reference and value. Several program examples are presented in this chapter. For those who choose to program in C++ we give also an introduction to how to program classes and the auxiliary library Blitz++, which contains several useful classes for numerical operations on vectors and matrices. This chapter contains also sections on numerical interpolation and extrapolation. Chapter ?? deals with the solution of non-linear equations and the finding of roots of polynomials. The link to Blitz++, matrices and selected algorithms for linear algebra problems are dealt with in chapter ??.

Therafter we switch to numerical integration for integrals with few dimensions, typically less than three, in chapter ??. The numerical integration chapter serves also to justify the introduction of Monte-Carlo methods discussed in chapters ?? and ??. There, a variety of applications are presented, from integration of multi-dimensional integrals to problems in statistical physics such as random walks and the derivation of the diffusion equation from Brownian motion. Chapter ?? continues this discussion by extending to studies of phase transitions in statistical physics. Chapter ?? deals with Monte-Carlo studies of quantal systems, with an emphasis on

variational Monte Carlo methods and diffusion Monte Carlo methods. In chapter ?? we deal with eigensystems and applications to e.g., the Schrödinger equation rewritten as a matrix diagonalization problem. Problems from scattering theory are also discussed, together with the most used solution methods for systems of linear equations. Finally, we discuss various methods for solving differential equations and partial differential equations in chapters ??-?? with examples ranging from harmonic oscillations, equations for heat conduction and the time dependent Schrödinger equation. The emphasis is on various finite difference methods.

We assume that you have taken an introductory course in programming and have some familiarity with high-level or low-level and modern languages such as Java, Python, C++, Fortran 77/90/95, etc. Fortran¹ and C++ are examples of compiled low-level languages, in contrast to interpreted ones like Maple or Matlab. In such compiled languages the computer translates an entire subprogram into basic machine instructions all at one time. In an interpreted language the translation is done one statement at a time. This clearly increases the computational time expenditure. More detailed aspects of the above two programming languages will be discussed in the lab classes and various chapters of this text.

There are several texts on computational physics on the market, see for example Refs. [5, 6, 7, 8, 11? ? ?], ranging from introductory ones to more advanced ones. Most of these texts treat however in a rather cavalier way the mathematics behind the various numerical methods. We've also succumbed to this approach, mainly due to the following reasons: several of the methods discussed are rather involved, and would thus require at least a one-semester course for an introduction. In so doing, little time would be left for problems and computation. This course is a compromise between three disciplines, numerical methods, problems from the physical sciences and computation. To achieve such a synthesis, we will have to relax our presentation in order to avoid lengthy and gory mathematical expositions. You should also keep in mind that computational physics and science in more general terms consist of the combination of several fields and crafts with the aim of finding solution strategies for complicated problems. However, where we do indulge in presenting more formalism, we have borrowed heavily from several texts on mathematical analysis.

1.1 Choice of programming language

As programming language we have ended up with preferring C++, but all examples discussed in the text have their corresponding Fortran and Python programs on the webpage of this text.

¹With Fortran we will consistently mean Fortran 2008. There are no programming examples in Fortran 77 in this text.

Fortran (FORmula TRANslation) was introduced in 1957 and remains in many scientific computing environments the language of choice. The latest standard, see Refs. [9? ? ?], includes extensions that are familiar to users of C++. Some of the most important features of Fortran include recursive subroutines, dynamic storage allocation and pointers, user defined data structures, modules, and the ability to manipulate entire arrays. However, there are several good reasons for choosing C++ as programming language for scientific and engineering problems. Here are some:

- C++ is now the dominating language in Unix and Windows environments. It is widely available and is the language of choice for system programmers. It is very widespread for developments of non-numerical software
- The C++ syntax has inspired lots of popular languages, such as Perl, Python and Java.
- It is an extremely portable language, all Linux and Unix operated machines have a C++ compiler.
- In the last years there has been an enormous effort towards developing numerical libraries for C++. Numerous tools (numerical libraries such as MPI[10? ?]) are written in C++ and interfacing them requires knowledge of C++. Most C++ and Fortran compilers compare fairly well when it comes to speed and numerical efficiency. Although Fortran 77 and C are regarded as slightly faster than C++ or Fortran, compiler improvements during the last few years have diminished such differences. The Java numerics project has lost some of its steam recently, and Java is therefore normally slower than C++ or Fortran.
- Complex variables, one of Fortran's strongholds, can also be defined in the new ANSI C++ standard.
- C++ is a language which catches most of the errors as early as possible, typically at compilation time. Fortran has some of these features if one omits implicit variable declarations.
- C++ is also an object-oriented language, to be contrasted with C and Fortran. This means that it supports three fundamental ideas, namely objects, class hierarchies and polymorphism. Fortran has, through the `MODULE` declaration the capability of defining classes, but lacks inheritance, although polymorphism is possible. Fortran is then considered as an object-based programming language, to be contrasted with C++ which has the capability of relating classes to each other in a hierarchical way.

An important aspect of C++ is its richness with more than 60 keywords allowing for a good balance between object orientation and numerical efficiency. Furthermore, careful programming can result in an efficiency close to Fortran 77. The language is well-suited for large projects and has presently good standard libraries suitable for computational science projects, although many of these still lag behind the large body of libraries for numerics available to Fortran programmers. However, it is not difficult to interface libraries written in Fortran with C++ codes, if care is exercised. Other weak sides are the fact that it can be easy to write inefficient code and that there are many ways of writing the same things, adding to the confusion for beginners and professionals as well. The language is also under continuous development, which often causes portability problems.

C++ is also a difficult language to learn. Grasping the basics is rather straightforward, but takes time to master. A specific problem which often causes unwanted or odd errors is dynamic memory management.

The efficiency of C++ codes are close to those provided by Fortran. This means often that a code written in Fortran 77 can be faster, however for large numerical projects C++ and Fortran are to be preferred. If speed is an issue, one could port critical parts of the code to Fortran 77.

Future plans

Since our undergraduate curriculum has changed considerably from the beginning of the fall semester of 2007, with the introduction of Python as programming language, the content of this course will change accordingly from the fall semester 2009. C++ and Fortran will then coexist with Python and students can choose between these three programming languages. The emphasis in the text will be on C++ programming, but how to interface C++ or Fortran programs with Python codes will also be discussed. Tools like Cython (or SWIG) are highly recommended, see for example the Cython link at <http://cython.org>.

1.2 Designing programs

Before we proceed with a discussion of numerical methods, we would like to remind you of some aspects of program writing.

In writing a program for a specific algorithm (a set of rules for doing mathematics or a precise description of how to solve a problem), it is obvious that different programmers will apply different styles, ranging from barely readable ² (even for the programmer) to well documented codes which can be used and extended upon

²As an example, a bad habit is to use variables with no specific meaning, like x1, x2 etc, or names for subprograms which go like routine1, routine2 etc.

by others in e.g., a project. The lack of readability of a program leads in many cases to credibility problems, difficulty in letting others extend the codes or remembering oneself what a certain statement means, problems in spotting errors, not always easy to implement on other machines, and so forth. Although you should feel free to follow your own rules, we would like to focus certain suggestions which may improve a program. What follows here is a list of our recommendations (or biases/prejudices).

First about designing a program.

- Before writing a single line, have the algorithm clarified and understood. It is crucial to have a logical structure of e.g., the flow and organization of data before one starts writing.
- Always try to choose the simplest algorithm. Computational speed can be improved upon later.
- Try to write a as clear program as possible. Such programs are easier to debug, and although it may take more time, in the long run it may save you time. If you collaborate with other people, it reduces spending time on debugging and trying to understand what the codes do. A clear program will also allow you to remember better what the program really does!
- Implement a working code with emphasis on design for extensions, maintenance etc. Focus on the design of your code in the beginning and don't think too much about efficiency before you have a thoroughly debugged and verified program. A rule of thumb is the so-called 80 – 20 rule, 80 % of the CPU time is spent in 20 % of the code and you will experience that typically only a small part of your code is responsible for most of the CPU expenditure. Therefore, spend most of your time in devising a good algorithm.
- The planning of the program should be from top down to bottom, trying to keep the flow as linear as possible. Avoid jumping back and forth in the program. First you need to arrange the major tasks to be achieved. Then try to break the major tasks into subtasks. These can be represented by functions or subprograms. They should accomplish limited tasks and as far as possible be independent of each other. That will allow you to use them in other programs as well.
- Try always to find some cases where an analytical solution exists or where simple test cases can be applied. If possible, devise different algorithms for solving the same problem. If you get the same answers, you may have coded things correctly or made the same error twice.
- When you have a working code, you should start thinking of the efficiency. Analyze the efficiency with a tool (profiler) to predict the CPU-intensive parts.

Attack then the CPU-intensive parts after the program reproduces benchmark results.

However, although we stress that you should post-pone a discussion of the efficiency of your code to the stage when you are sure that it runs correctly, there are some simple guidelines to follow when you design the algorithm.

- Avoid lists, sets etc., when arrays can be used without too much waste of memory. Avoid also calls to functions in the innermost loop since that produces an overhead in the call.
- Heavy computation with small objects might be inefficient, e.g., vector of class complex objects
- Avoid small virtual functions (unless they end up in more than (say) 5 multiplications)
- Save object-oriented constructs for the top level of your code.
- Use tailored library functions for various operations, if possible.
- Reduce pointer-to-pointer-to....-pointer links inside loops.
- Avoid implicit type conversion, use rather the explicit keyword when declaring constructors in C++.
- Never return (copy) of an object from a function, since this normally implies a hidden allocation.

Finally, here are some of our favorite approaches to code writing.

- Use always the standard ANSI version of the programming language. Avoid local dialects if you wish to port your code to other machines.
- Add always comments to describe what a program or subprogram does. Comment lines help you remember what you did e.g., one month ago.
- Declare all variables. Avoid totally the `IMPLICIT` statement in Fortran. The program will be more readable and help you find errors when compiling.
- Do not use `GOTO` structures in Fortran. Although all varieties of spaghetti are great culinary temptations, spaghetti-like Fortran with many `GOTO` statements is to be avoided. Extensive amounts of time may be wasted on decoding other authors' programs.

- When you name variables, use easily understandable names. Avoid `v1` when you can use `speed_of_light`. Associative names make it easier to understand what a specific subprogram does.
- Use compiler options to test program details and if possible also different compilers. They make errors too.
- Writing codes in C++ and Fortran may often lead to segmentation faults. This means in most cases that we are trying to access elements of an array which are not available. When developing a code it is then useful to compile with debugging options. The use of debuggers and profiling tools is something we highly recommend during the development of a program.

For more detailed texts on C++ programming in engineering and science are the books by Flowers [4] and Barton and Nackman [1]. The classic text on C++ programming is the book of Bjarne Stroustrup [2]. The Fortran 95 standard is well documented in Refs. [9, 10] while the new details of Fortran 2003 and 2008 can be found in Ref. [11]. The reader should note that this is not a text on C++ or Fortran. It is therefore important that one tries to find additional literature on these programming languages. Good Python texts on scientific computing are [12, 13].

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Chapter 2

Introduction to C++ and Fortran

This chapter aims at catching two birds with a stone; to introduce to you essential features of the programming languages C++ and Fortran with a brief reminder on Python specific topics, and to stress problems like overflow, underflow, round off errors and eventually loss of precision due to the finite amount of numbers a computer can represent. The programs we discuss are tailored to these aims. You will also learn to

2.1 Getting Started

In programming languages we encounter data entities such as constants, variables, results of evaluations of functions etc. Common to these objects is that they can be represented through the type concept. There are intrinsic types and derived types. Intrinsic types are provided by the programming language whereas derived types are provided by the programmer. If one specifies the type to be for example `INTEGER (KIND=2)` for Fortran¹ or `short int/int` in C++, the programmer selects a particular data type with 2 bytes (16 bits) for every item of the class `INTEGER (KIND=2)` or `int`. Intrinsic types come in two classes, numerical (like integer, real or complex) and non-numeric (as logical and character). The general form for declaring variables is `data type name of variable` and Table 2.1 lists the standard variable declarations of C++ and Fortran (note well that there be may compiler and machine differences from the table below). An important aspect when declaring variables is their region of validity. Inside a function we define a variable through the expression `int var` or `INTEGER :: var`. The question is whether this variable is available in other functions as well, moreover where is `var` initialized and

¹Our favoured display mode for Fortran statements will be capital letters for language statements and low key letters for user-defined statements. Note that Fortran does not distinguish between capital and low key letters while C++ does.

finally, if we call the function where it is declared, is the value conserved from one call to the other?

Table 2.1: Examples of variable declarations for C++ and Fortran . We reserve capital letters for Fortran declaration statements throughout this text, although Fortran is not sensitive to upper or lowercase letters. Note that there are machines which allow for more than 64 bits for doubles. The ranges listed here may therefore vary.

type in C++ and Fortran	bits	range
int/INTEGER (2)	16	−32768 to 32767
unsigned int	16	0 to 65535
signed int	16	−32768 to 32767
short int	16	−32768 to 32767
unsigned short int	16	0 to 65535
signed short int	16	−32768 to 32767
int/long int/INTEGER(4)	32	−2147483648 to 2147483647
signed long int	32	−2147483648 to 2147483647
float/REAL(4)	32	10^{-44} to 10^{+38}
double/REAL(8)	64	10^{-322} to $10e^{+308}$

Both C++ and Fortran operate with several types of variables and the answers to these questions depend on how we have defined for example an integer via the statement `int var`. Python on the other hand does not use variable or function types (they are not explicitly written), allowing thereby for a better potential for reuse of the code.

The following list may help in clarifying the above points:

type of variable	validity
local variables	defined within a function, only available within the scope of the function.
formal parameter	If it is defined within a function it is only available within that specific function.
global variables	Defined outside a given function, available for all functions from the point where it is defined.

In Table 2.1 we show a list of some of the most used language statements in Fortran and C++.

Fortran	C++
Program structure	
PROGRAM something	main ()
FUNCTION something(input)	double (int) something(input)
SUBROUTINE something(inout)	
Data type declarations	
REAL (4) x, y	float x, y;
REAL(8) :: x, y	double x, y;
INTEGER :: x, y	int x,y;
CHARACTER :: name	char name;
REAL(8), DIMENSION(dim1,dim2) :: x	double x[dim1][dim2];
INTEGER, DIMENSION(dim1,dim2) :: x	int x[dim1][dim2];
LOGICAL :: x	
TYPE name	struct name {
declarations	declarations;
END TYPE name	}
POINTER :: a	double (int) *a;
ALLOCATE	new;
DEALLOCATE	delete;
Logical statements and control structure	
IF (a == b) THEN	if (a == b)
b=0	{ b=0;
ENDIF	}
DO WHILE (logical statement)	while (logical statement)
do something	{do something
ENDDO	}
IF (a >= b) THEN	if (a >= b)
b=0	{ b=0;
ELSE	else
a=0	a=0; }
ENDIF	
SELECT CASE (variable)	switch(variable)
CASE (variable=value1)	{
do something	case 1:
CASE (...)	variable=value1;
...	do something;
	break;
END SELECT	case 2:
	do something; break; ...
	}
DO i=0, end, 1	for(i=0; i<= end; i++)
do something	{ do something ;
ENDDO	}

Table 2.2: Elements of programming syntax.

In addition, both C++ and Fortran allow for complex variables. In Fortran we would declare a complex variable as `COMPLEX (KIND=16):: x, y` which refers to a double with word length of 16 bytes. In C++ we would need to include a complex library through the statements

```
#include <complex>
complex<double> x, y;
```

We will discuss the above declaration `complex<double> x,y;` in more detail in chapter ??.

2.1.1 Scientific hello world

Our first programming encounter is the 'classical' one, found in almost every text-book on computer languages, the 'hello world' code, here in a scientific disguise. We present first the C version.

[Click here to view code](#)

```
/* comments in C begin like this and end with */
#include <stdlib.h> /* atof function */
#include <math.h> /* sine function */
#include <stdio.h> /* printf function */

int main (int argc, char* argv[])
{
    double r, s; /* declare variables */
    r = atof(argv[1]); /* convert the text argv[1] to double */
    s = sin(r);
    printf("Hello, World! sin(%g)=%g\n", r, s);
    return 0; /* success execution of the program */
}
```

The compiler must see a declaration of a function before you can call it (the compiler checks the argument and return types). The declaration of library functions appears in so-called header files that must be included in the program, for example `#include <stdlib.h>`.

We call three functions `atof`, `sin`, `printf` and these are declared in three different header files. The main program is a function called `main` with a return value set to an integer, returning 0 if success. The operating system stores the return value, and other programs/utilities can check whether the execution was successful or not. The command-line arguments are transferred to the main function through the statement

```
int main (int argc, char* argv[])
```

The integer `argc` stands for the number of command-line arguments, set to one in our case, while `argv` is a vector of strings containing the command-line arguments with `argv[0]` containing the name of the program and `argv[1]`, `argv[2]`, ... are the command-line args, i.e., the number of lines of input to the program.

This means that we would run the programs as `mhjensen@compphys: ./myprogram.exe 0.3`. The name of the program enters `argv[0]` while the text string `0.2` enters `argv[1]`. Here we define a floating point variable, see also below, through the keywords `float` for single precision real numbers and `double` for double precision. The function `atof` transforms a text (`argv[1]`) to a float. The sine function is declared in `math.h`, a library which is not automatically included and needs to be linked when computing an executable file.

With the command `printf` we obtain a formatted printout. The `printf` syntax is used for formatting output in many C-inspired languages (Perl, Python, awk, partly C++).

In C++ this program can be written as

[Click here to view code](#)

```
// A comment line begins like this in C++ programs
using namespace std;
#include <iostream>
#include <cstdlib>
#include <cmath>
int main (int argc, char* argv[])
{
    // convert the text argv[1] to double using atof:
    double r = atof(argv[1]);
    double s = sin(r);
    cout << "Hello, World! sin(" << r << ")=" << s << endl;
    // success
    return 0;
}
```

We have replaced the call to `printf` with the standard C++ function `cout`. The header file `iostream` is then needed. In addition, we don't need to declare variables like `r` and `s` at the beginning of the program. I personally prefer however to declare all variables at the beginning of a function, as this gives me a feeling of greater readability. Note that we have used the declaration using `namespace std`;. Namespace is a way to collect all functions defined in C++ libraries. If we omit this declaration on top of the program we would have to add the declaration `std` in front of `cout` or `cin`. Our program would then read

[Click here to view code](#)

```
// Hello world code without using namespace std
#include <iostream>
```

```
#include <cstdlib>
#include <cmath>
int main (int argc, char* argv[])
{
    // convert the text argv[1] to double using atof:
    double r = atof(argv[1]);
    double s = sin(r);
    std::cout << "Hello, World! sin(" << r << ")=" << s << endl;
    // success
    return 0;
}
```

Another feature which is worth noting is that we have skipped exception handling here. Later in this chapter we discuss examples that test our input from the command line. But it is easy to add such a feature, as shown in our modified hello world program

[Click here to view code](#)

```
// Hello world code with exception handling
using namespace std;
#include <cstdlib>
#include <cmath>
#include <iostream>
int main (int argc, char* argv[])
{
    // Read in output file, abort if there are too few command-line arguments
    if( argc <= 1 ){
        cout << "Bad Usage: " << argv[0] <<
            " read also a number on the same line, e.g., prog.exe 0.2" << endl;
        exit(1); // here the program stops.
    }
    // convert the text argv[1] to double using atof:
    double r = atof(argv[1]);
    double s = sin(r);
    cout << "Hello, World! sin(" << r << ")=" << s << endl;
    // success
    return 0;
}
```

Here we test that we have more than one argument. If not, the program stops and writes to screen an error message. Observe also that we have included the mathematics library via the `#include <cmath>` declaration.

To run these programs, you need first to compile and link them in order to obtain an executable file under operating systems like e.g., UNIX or Linux. Before we proceed we give therefore examples on how to obtain an executable file under Linux/Unix.

In order to obtain an executable file for a C++ program, the following instructions under Linux/Unix can be used @@

```
c++ -c -Wall myprogram.c
c++ -o myprogram myprogram.o
```

where the compiler is called through the command c++. The compiler option -Wall means that a warning is issued in case of non-standard language. The executable file is in this case myprogram. The option -c is for compilation only, where the program is translated into machine code, while the -o option links the produced object file myprogram.o and produces the executable myprogram.

The corresponding Fortran code is

[Click here to view code](#)

```
PROGRAM shw
  IMPLICIT NONE
  REAL (KIND=8) :: r  ! Input number
  REAL (KIND=8) :: s  ! Result

  ! Get a number from user
  WRITE(*,*) 'Input a number: '
  READ(*,*) r
  ! Calculate the sine of the number
  s = SIN(r)
  ! Write result to screen
  WRITE(*,*) 'Hello World! SINE of ', r, ' =', s
END PROGRAM shw
```

The first statement must be a program statement; the last statement must have a corresponding end program statement. Integer numerical variables and floating point numerical variables are distinguished. The names of all variables must be between 1 and 31 alphanumeric characters of which the first must be a letter and the last must not be an underscore. Comments begin with a ! and can be included anywhere in the program. Statements are written on lines which may contain up to 132 characters. The asterisks (*,*) following WRITE represent the default format for output, i.e., the output is e.g., written on the screen. Similarly, the READ(*,*) statement means that the program is expecting a line input. Note also the IMPLICIT NONE statement which we strongly recommend the use of. In many Fortran 77 programs one can find statements like IMPLICIT REAL*8(a-h,o-z), meaning that all variables beginning with any of the above letters are by default floating numbers. However, such a usage makes it hard to spot eventual errors due to misspelling of variable names. With IMPLICIT NONE you have to declare all variables and therefore detect possible errors already while compiling. I recommend strongly that you declare all variables when using Fortran.

We call the Fortran compiler (using free format) through @@

```
gfortran -c -free myprogram.f90
gfortran -o myprogram.x myprogram.o
```

Under Linux/Unix it is often convenient to create a so-called makefile, which is a script which includes possible compiling commands, in order to avoid retyping the above lines every once and then we have made modifications to our program. A typical makefile for the above *cc* compiling options is listed below @@

```
# General makefile for c - choose PROG =   name of given program
```

```
# Here we define compiler option, libraries and the target
CC= c++ -Wall
PROG= myprogram
```

```
# Here we make the executable file
${PROG} :          ${PROG}.o
                  ${CC} ${PROG}.o -o ${PROG}
```

```
# whereas here we create the object file
```

```
${PROG}.o :        ${PROG}.cpp
                  ${CC} -c ${PROG}.cpp
```

If you name your file for 'makefile', simply type the command **make** and Linux/Unix executes all of the statements in the above makefile. Note that C++ files have the extension .cpp

For Fortran, a similar makefile is @@

```
# General makefile for F90 - choose PROG =   name of given program
```

```
# Here we define compiler options, libraries and the target
F90= gfortran
PROG= myprogram
```

```
# Here we make the executable file
${PROG} :          ${PROG}.o
                  ${F90} ${PROG}.o -o ${PROG}
```

```
# whereas here we create the object file
```

```

${PROG}.o :      ${PROG}.f90
                  ${F90} -c ${PROG}.f

```

Finally, for the sake of completeness, we list the corresponding Python code

[Click here to view code](#)

```

#!/usr/bin/env python

import sys, math
# Read in a string a convert it to a float
r = float(sys.argv[1])
s = math.sin(r)
print "Hello, World! sin(%g)=%12.6e" % (r,s)

```

where we have used a formatted printout with scientific notation. In Python we do not need to declare variables. Mathematical functions like the `sin` function are imported from the `math` module. For further references to Python and its syntax, we recommend the text of Hans Petter Langtangen [?]. The corresponding codes in Python are available at the webpage of the course. All programs are listed as a directory tree beginning with `programs/chapterxx`. Each chapter has in turn three directories, one for C++, one for Fortran and finally one for Python codes. The Fortran codes in this chapter can be found in the directory `programs/chapter02/Fortran`.

2.2 Representation of Integer Numbers

In Fortran a keyword for declaration of an integer is `INTEGER (KIND=n)`, $n = 2$ reserves 2 bytes (16 bits) of memory to store the integer variable whereas $n = 4$ reserves 4 bytes (32 bits). In Fortran, although it may be compiler dependent, just declaring a variable as `INTEGER`, reserves 4 bytes in memory as default.

In C++ keywords are `short int`, `int`, `long int`, `long long int`. The byte-length is compiler dependent within some limits. The GNU C++-compilers (called by `gcc` or `g++`) assign 4 bytes (32 bits) to variables declared by `int` and `long int`. Typical byte-lengths are 2, 4, 4 and 8 bytes, for the types given above. To see how many bytes are reserved for a specific variable, C++ has a library function called `sizeof(type)` which returns the number of bytes for `type`.

An example of a program declaration is

```

Fortran:      INTEGER (KIND=2) :: age_of_participant
C++:          short int          age_of_participant;

```

Note that the `(KIND=2)` can be written as `(2)`. Normally however, we will for Fortran programs just use the 4 bytes default assignment `INTEGER`.

In the above examples one bit is used to store the sign of the variable `age_of_participant` and the other 15 bits are used to store the number, which then may range from zero to $2^{15} - 1 = 32767$. This should definitely suffice for human lifespans. On the other hand, if we were to classify known fossils by age we may need

```
Fortran:      INTEGER (4) :: age_of_fossile
C++:         int          age_of_fossile;
```

Again one bit is used to store the sign of the variable `age_of_fossile` and the other 31 bits are used to store the number which then may range from zero to $2^{31} - 1 = 2.147.483.647$. In order to give you a feeling how integer numbers are represented in the computer, think first of the decimal representation of the number 417

$$417 = 4 \times 10^2 + 1 \times 10^1 + 7 \times 10^0,$$

which in binary representation becomes

$$417 = a_n 2^n + a_{n-1} 2^{n-1} + a_{n-2} 2^{n-2} + \dots + a_0 2^0,$$

where the coefficients a_k with $k = 0, \dots, n$ are zero or one. They can be calculated through successive division by 2 and using the remainder in each division to determine the numbers a_n to a_0 . A given integer in binary notation is then written as

$$a_n 2^n + a_{n-1} 2^{n-1} + a_{n-2} 2^{n-2} + \dots + a_0 2^0.$$

In binary notation we have thus

$$(417)_{10} = (110100001)_2,$$

since we have

$$(110100001)_2 = 1 \times 2^8 + 1 \times 2^7 + 0 \times 2^6 + 1 \times 2^5 + 0 \times 2^4 + 0 \times 2^3 + 0 \times 2^2 + 0 \times 2^1 + 1 \times 2^0.$$

To see this, we have performed the following divisions by 2

417/2=208	remainder 1	coefficient of 2^0 is 1
208/2=104	remainder 0	coefficient of 2^1 is 0
104/2=52	remainder 0	coefficient of 2^2 is 0
52/2=26	remainder 0	coefficient of 2^3 is 0
26/2=13	remainder 0	coefficient of 2^4 is 0
13/2= 6	remainder 1	coefficient of 2^5 is 1
6/2= 3	remainder 0	coefficient of 2^6 is 0
3/2= 1	remainder 1	coefficient of 2^7 is 1
1/2= 0	remainder 1	coefficient of 2^8 is 1

We see that nine bits are sufficient to represent 417. Normally we end up using 32 bits as default for integers, meaning that our number reads

[illegible]

A simple program which performs these operations is listed below. Here we employ the modulus operation (with division by 2), which in C++ is given by the `a%2` operator. In Fortran we would call the function `MOD(a,2)` in order to obtain the remainder of a division by 2.

[Click here to view code](#)

```
using namespace std;
#include <iostream>

int main (int argc, char* argv[])
{
    int i;
    int terms[32]; // storage of a0, a1, etc, up to 32 bits
    int number = atoi(argv[1]);
    // initialise the term a0, a1 etc
    for (i=0; i < 32 ; i++){ terms[i] = 0;}
    for (i=0; i < 32 ; i++){
        terms[i] = number%2;
        number /= 2;
    }
    // write out results
    cout << `` Number of bytes used= `` << sizeof(number) << endl;
    for (i=0; i < 32 ; i++){
        cout << `` Term nr: `` << i << ``Value= `` << terms[i];
        cout << endl;
    }
    return 0;
}
```

The C++ function `sizeof` yields the number of bytes reserved for a specific variable. Note also the `for` construct. We have reserved a fixed array which contains the values of a_i being 0 or 1, the remainder of a division by two. We have enforced the integer to be represented by 32 bits, or four bytes, which is the default integer representation.

Note that for 417 we need 9 bits in order to represent it in a binary notation, while a number like the number 3 is given in an 32 bits word as

$$(3)_{10} = (0000000000000000000000000000000011)_2.$$

For this number 2 significant bits would be enough.

With these prerequisites in mind, it is rather obvious that if a given integer variable is beyond the range assigned by the declaration statement we may encounter problems.

If we multiply two large integers $n_1 \times n_2$ and the product is too large for the bit size allocated for that specific integer assignment, we run into an overflow problem. The most significant bits are lost and the least significant kept. Using 4 bytes for integer variables the result becomes

$$2^{20} \times 2^{20} = 0.$$

However, there are compilers or compiler options that preprocess the program in such a way that an error message like 'integer overflow' is produced when running the program. Here is a small program which may cause overflow problems when running (try to test your own compiler in order to be sure how such problems need to be handled).

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/cpp/program3.cpp>

```
// Program to calculate 2**n
using namespace std;
#include <iostream>

int main()
{
    int int1, int2, int3;
    // print to screen
    cout << "Read in the exponential N for 2^N =\n";
    // read from screen
    cin >> int2;
    int1 = (int) pow(2., (double) int2);
    cout << " 2^N * 2^N = " << int1*int1 << "\n";
    int3 = int1 - 1;
    cout << " 2^N*(2^N - 1) = " << int1 * int3 << "\n";
    cout << " 2^N- 1 = " << int3 << "\n";
    return 0;
}
// End: program main()
```

If we run this code with an exponent $N = 32$, we obtain the following output @@

```
2^N * 2^N = 0
2^N*(2^N - 1) = -2147483648
2^N- 1 = 2147483647
```

We notice that 2^{64} exceeds the limit for integer numbers with 32 bits. The program returns 0. This can be dangerous, since the results from the operation $2^N(2^N - 1)$ is obviously wrong. One possibility to avoid such cases is to add compilation options which flag if an overflow or underflow is reached.

2.2.1 Fortran codes

The corresponding Fortran code is

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/Fortran/program2.f90>

```
PROGRAM binary_integer
IMPLICIT NONE
  INTEGER i, number, terms(0:31) ! storage of a0, a1, etc, up to 32 bits,
  ! note array length running from 0:31. Fortran allows negative indexes as well.

  WRITE(*,*) 'Give a number to transform to binary notation'
  READ(*,*) number
  ! Initialise the terms a0, a1 etc
  terms = 0
  ! Fortran takes only integer loop variables
  DO i=0, 31
    terms(i) = MOD(number,2) ! Modulus function in Fortran
    number = number/2
  ENDDO
  ! write out results
  WRITE(*,*) 'Binary representation '
  DO i=0, 31
    WRITE(*,*) ' Term nr and value', i, terms(i)
  ENDDO
END PROGRAM binary_integer
```

and

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/Fortran/program3.f90>

```
PROGRAM integer_exp
IMPLICIT NONE
INTEGER :: int1, int2, int3
  ! This is the begin of a comment line in Fortran 90
  ! Now we read from screen the variable int2
  WRITE(*,*) 'Read in the number to be exponentiated'
  READ(*,*) int2
  int1=2**int2
  WRITE(*,*) '2^N*2^N', int1*int1
  int3=int1-1
  WRITE(*,*) '2^N*(2^N-1)', int1*int3
  WRITE(*,*) '2^N-1', int3
END PROGRAM integer_exp
```

In Fortran the modulus division is performed by the intrinsic function `MOD(number, 2)` in case of a division by 2. The exponentiation of a number is given by for example `2**N` instead of the call to the `pow` function in C++.

2.3 Real Numbers and Numerical Precision

An important aspect of computational physics is the numerical precision involved. To design a good algorithm, one needs to have a basic understanding of propagation of inaccuracies and errors involved in calculations. There is no magic recipe for dealing with underflow, overflow, accumulation of errors and loss of precision, and only a careful analysis of the functions involved can save one from serious problems.

Since we are interested in the precision of the numerical calculus, we need to understand how computers represent real and integer numbers. Most computers deal with real numbers in the binary, octal and/or hexadecimal systems, in contrast to the decimal system that we humans prefer to use. The binary system uses 2 as the base, in much the same way that the decimal system uses 10. Since the typical computer communicates with us in the decimal system, but works internally in e.g., the binary system, conversion procedures must be executed by the computer, and these conversions involve hopefully only small roundoff errors

Computers are also not able to operate using real numbers expressed with more than a fixed number of digits, and the set of values possible is only a subset of the mathematical integers or real numbers. The so-called word length we reserve for a given number places a restriction on the precision with which a given number is represented. This means in turn, that for example floating numbers are always rounded to a machine dependent precision, typically with 6-15 leading digits to the right of the decimal point. Furthermore, each such set of values has a processor-dependent smallest negative and a largest positive value.

Why do we at all care about rounding and machine precision? The best way is to consider a simple example first. In the following example we assume that we can represent a floating number with a precision of 5 digits only to the right of the decimal point. This is nothing but a mere choice of ours, but mimicks the way numbers are represented in the machine.

Suppose we wish to evaluate the function

$$f(x) = \frac{1 - \cos(x)}{\sin(x)},$$

for small values of x . If we multiply the denominator and numerator with $1 + \cos(x)$ we obtain the equivalent expression

$$f(x) = \frac{\sin(x)}{1 + \cos(x)}.$$

If we now choose $x = 0.006$ (in radians) our choice of precision results in

$$\sin(0.007) \approx 0.59999 \times 10^{-2},$$

and

$$\cos(0.007) \approx 0.99998.$$

The first expression for $f(x)$ results in

$$f(x) = \frac{1 - 0.99998}{0.59999 \times 10^{-2}} = \frac{0.2 \times 10^{-4}}{0.59999 \times 10^{-2}} = 0.33334 \times 10^{-2},$$

while the second expression results in

$$f(x) = \frac{0.59999 \times 10^{-2}}{1 + 0.99998} = \frac{0.59999 \times 10^{-2}}{1.99998} = 0.30000 \times 10^{-2},$$

which is also the exact result. In the first expression, due to our choice of precision, we have only one relevant digit in the numerator, after the subtraction. This leads to a loss of precision and a wrong result due to a cancellation of two nearly equal numbers. If we had chosen a precision of six leading digits, both expressions yield the same answer. If we were to evaluate $x \sim \pi$, then the second expression for $f(x)$ can lead to potential losses of precision due to cancellations of nearly equal numbers.

This simple example demonstrates the loss of numerical precision due to roundoff errors, where the number of leading digits is lost in a subtraction of two near equal numbers. The lesson to be drawn is that we cannot blindly compute a function. We will always need to carefully analyze our algorithm in the search for potential pitfalls. There is no magic recipe however, the only guideline is an understanding of the fact that a machine cannot represent correctly **all** numbers.

2.3.1 Representation of real numbers

Real numbers are stored with a decimal precision (or mantissa) and the decimal exponent range. The mantissa contains the significant figures of the number (and thereby the precision of the number). A number like $(9.90625)_{10}$ in the decimal representation is given in a binary representation by

$$(1001.11101)_2 = 1 \times 2^3 + 0 \times 2^2 + 0 \times 2^1 + 1 \times 2^0 + 1 \times 2^{-1} + 1 \times 2^{-2} + 1 \times 2^{-3} + 0 \times 2^{-4} + 1 \times 2^{-5},$$

and it has an exact machine number representation since we need a finite number of bits to represent this number. This representation is however not very practical. Rather, we prefer to use a scientific notation. In the decimal system we would write a number like 9.90625 in what is called the normalized scientific notation. This means simply that the decimal point is shifted and appropriate powers of 10 are supplied. Our number could then be written as

$$9.90625 = 0.990625 \times 10^1,$$

and a real non-zero number could be generalized as

$$x = \pm r \times 10^n,$$

with a r a number in the range $1/10 \leq r < 1$. In a similar way we can represent a binary number in scientific notation as

$$x = \pm q \times 2^m,$$

with a q a number in the range $1/2 \leq q < 1$. This means that the mantissa of a binary number would be represented by the general formula

$$(0.a_{-1}a_{-2}\dots a_{-n})_2 = a_{-1} \times 2^{-1} + a_{-2} \times 2^{-2} + \dots + a_{-n} \times 2^{-n}.$$

In a typical computer, floating-point numbers are represented in the way described above, but with certain restrictions on q and m imposed by the available word length. In the machine, our number x is represented as

$$x = (-1)^s \times \text{mantissa} \times 2^{\text{exponent}},$$

where s is the sign bit, and the exponent gives the available range. With a single-precision word, 32 bits, 8 bits would typically be reserved for the exponent, 1 bit for the sign and 23 for the mantissa. This means that if we define a variable as

```
Fortran:      REAL (4) :: size_of_fossile
C++:         float      size_of_fossile;
```

we are reserving 4 bytes in memory, with 8 bits for the exponent, 1 for the sign and 23 bits for the mantissa, implying a numerical precision to the sixth or seventh digit, since the least significant digit is given by $1/2^{23} \approx 10^{-7}$. The range of the exponent goes from $2^{-128} = 2.9 \times 10^{-39}$ to $2^{127} = 3.4 \times 10^{38}$, where 128 stems from the fact that 8 bits are reserved for the exponent.

A modification of the scientific notation for binary numbers is to require that the leading binary digit 1 appears to the left of the binary point. In this case the representation of the mantissa q would be $(1.f)_2$ and $1 \leq q < 2$. This form is rather useful when storing binary numbers in a computer word, since we can always assume that the leading bit 1 is there. One bit of space can then be saved meaning that a 23 bits mantissa has actually 24 bits. This means explicitly that a binary number with 23 bits for the mantissa reads

$$(1.a_{-1}a_{-2}\dots a_{-23})_2 = 1 \times 2^0 + a_{-1} \times 2^{-1} + a_{-2} \times 2^{-2} + \dots + a_{-23} \times 2^{-23}.$$

As an example, consider the 32 bits binary number

$$(10111110111101000000000000000000)_2,$$

where the first bit is reserved for the sign, 1 in this case yielding a negative sign. The exponent m is given by the next 8 binary numbers 01111101 resulting in 125 in the decimal system. However, since the exponent has eight bits, this means it has $2^8 - 1 = 255$ possible numbers in the interval $-128 \leq m \leq 127$, our final exponent is $125 - 127 = -2$ resulting in 2^{-2} . Inserting the sign and the mantissa yields the final number in the decimal representation as

$$-2^{-2} \left(1 \times 2^0 + 1 \times 2^{-1} + 1 \times 2^{-2} + 1 \times 2^{-3} + 0 \times 2^{-4} + 1 \times 2^{-5} \right) = (-0.4765625)_{10}.$$

In this case we have an exact machine representation with 32 bits (actually, we need less than 23 bits for the mantissa).

If our number x can be exactly represented in the machine, we call x a machine number. Unfortunately, most numbers cannot and are thereby only approximated in the machine. When such a number occurs as the result of reading some input data or of a computation, an inevitable error will arise in representing it as accurately as possible by a machine number.

A floating number x , labelled $fl(x)$ will therefore always be represented as

$$fl(x) = x(1 \pm \epsilon_x), \quad (2.1)$$

with x the exact number and the error $|\epsilon_x| \leq |\epsilon_M|$, where ϵ_M is the precision assigned. A number like $1/10$ has no exact binary representation with single or double precision. Since the mantissa

$$1.(a_{-1}a_{-2}\dots a_{-n})_2$$

is always truncated at some stage n due to its limited number of bits, there is only a limited number of real binary numbers. The spacing between every real binary number is given by the chosen machine precision. For a 32 bit words this number is approximately $\epsilon_M \sim 10^{-7}$ and for double precision (64 bits) we have $\epsilon_M \sim 10^{-16}$, or in terms of a binary base as 2^{-23} and 2^{-52} for single and double precision, respectively.

2.3.2 Machine numbers

To understand that a given floating point number can be written as in Eq. (2.1), we assume for the sake of simplicity that we work with real numbers with words of length 32 bits, or four bytes. Then a given number x in the binary representation can be represented as

$$x = (1.a_{-1}a_{-2}\dots a_{-23}a_{-24}a_{-25}\dots)_2 \times 2^n,$$

or in a more compact form

$$x = r \times 2^n,$$

with $1 \leq r < 2$ and $-126 \leq n \leq 127$ since our exponent is defined by eight bits.

In most cases there will not be an exact machine representation of the number x . Our number will be placed between two exact 32 bits machine numbers x_- and x_+ . Following the discussion of Kincaid and Cheney [1] these numbers are given by

$$x_- = (1.a_{-1}a_{-2}\dots a_{-23})_2 \times 2^n,$$

and

$$x_+ = ((1.a_{-1}a_{-2}\dots a_{-23}))_2 + 2^{-23} \times 2^n.$$

If we assume that our number x is closer to x_- we have that the absolute error is constrained by the relation

$$|x - x_-| \leq \frac{1}{2}|x_+ - x_-| = \frac{1}{2} \times 2^{n-23} = 2^{n-24}.$$

A similar expression can be obtained if x is closer to x_+ . The absolute error conveys one type of information. However, we may have cases where two equal absolute errors arise from rather different numbers. Consider for example the decimal numbers $a = 2$ and $\bar{a} = 2.001$. The absolute error between these two numbers is 0.001. In a similar way, the two decimal numbers $b = 2000$ and $\bar{b} = 2000.001$ give exactly the same absolute error. We note here that $\bar{b} = 2000.001$ has more leading digits than b .

If we compare the relative errors

$$\frac{|a - \bar{a}|}{|a|} = 1.0 \times 10^{-3}, \quad \frac{|b - \bar{b}|}{|b|} = 1.0 \times 10^{-6},$$

we see that the relative error in b is much smaller than the relative error in a . We will see below that the relative error is intimately connected with the number of leading digits in the way we approximate a real number. The relative error is therefore the quantity of interest in scientific work. Information about the absolute error is normally of little use in the absence of the magnitude of the quantity being measured.

We define then the relative error for x as

$$\frac{|x - x_-|}{|x|} \leq \frac{2^{n-24}}{r \times 2^n} = \frac{1}{q} \times 2^{-24} \leq 2^{-24}.$$

Instead of using x_- and x_+ as the machine numbers closest to x , we introduce the relative error

$$\frac{|x - \bar{x}|}{|x|} \leq 2^{n-24},$$

with \bar{x} being the machine number closest to x . Defining

$$\epsilon_x = \frac{\bar{x} - x}{x},$$

we can write the previous inequality

$$fl(x) = x(1 + \epsilon_x)$$

where $|\epsilon_x| \leq \epsilon_M = 2^{-24}$ for variables of length 32 bits. The notation $fl(x)$ stands for the machine approximation of the number x . The number ϵ_M is given by the specified machine precision, approximately 10^{-7} for single and 10^{-16} for double precision, respectively.

There are several mathematical operations where an eventual loss of precision may appear. A subtraction, especially important in the definition of numerical derivatives discussed in chapter ?? is one important operation. In the computation of derivatives we end up subtracting two nearly equal quantities. In case of such a subtraction $a = b - c$, we have

$$fl(a) = fl(b) - fl(c) = a(1 + \epsilon_a),$$

or

$$fl(a) = b(1 + \epsilon_b) - c(1 + \epsilon_c),$$

meaning that

$$fl(a)/a = 1 + \epsilon_b \frac{b}{a} - \epsilon_c \frac{c}{a},$$

and if $b \approx c$ we see that there is a potential for an increased error in the machine representation of $fl(a)$. This is because we are subtracting two numbers of equal size and what remains is only the least significant part of these numbers. This part is prone to roundoff errors and if a is small we see that (with $b \approx c$)

$$\epsilon_a \approx \frac{b}{a}(\epsilon_b - \epsilon_c),$$

can become very large. The latter equation represents the relative error of this calculation. To see this, we define first the absolute error as

$$|fl(a) - a|,$$

whereas the relative error is

$$\frac{|fl(a) - a|}{a} \leq \epsilon_a.$$

The above subtraction is thus

$$\frac{|fl(a) - a|}{a} = \frac{|fl(b) - fl(c) - (b - c)|}{a},$$

yielding

$$\frac{|fl(a) - a|}{a} = \frac{|b\epsilon_b - c\epsilon_c|}{a}.$$

An interesting question is then how many significant binary bits are lost in a subtraction $a = b - c$ when we have $b \approx c$. The loss of precision theorem for a subtraction

$a = b - c$ states that [1]: if b and c are positive normalized floating-point binary machine numbers with $b > c$ and

$$2^{-r} \leq 1 - \frac{c}{b} \leq 2^{-s}, \quad (2.2)$$

then at most r and at least s significant binary bits are lost in the subtraction $b - c$. For a proof of this statement, see for example Ref. [1].

But even additions can be troublesome, in particular if the numbers are very different in magnitude. Consider for example the seemingly trivial addition $1 + 10^{-8}$ with 32 bits used to represent the various variables. In this case, the information contained in 10^{-8} is simply lost in the addition. When we perform the addition, the computer equates first the exponents of the two numbers to be added. For 10^{-8} this has however catastrophic consequences since in order to obtain an exponent equal to 10^0 , bits in the mantissa are shifted to the right. At the end, all bits in the mantissa are zeros.

This means in turn that for calculations involving real numbers (if we omit the discussion on overflow and underflow) we need to carefully understand the behavior of our algorithm, and test all possible cases where round-off errors and loss of precision can arise. Other cases which may cause serious problems are singularities of the type $0/0$ which may arise from functions like $\sin(x)/x$ as $x \rightarrow 0$. Such problems may also need the restructuring of the algorithm.

2.4 Programming Examples on Loss of Precision and Round-off Errors

2.4.1 Algorithms for e^{-x}

In order to illustrate the above problems, we discuss here some famous and perhaps less famous problems, including a discussion on specific programming features as well.

We start by considering three possible algorithms for computing e^{-x} :

1. by simply coding

$$e^{-x} = \sum_{n=0}^{\infty} (-1)^n \frac{x^n}{n!}$$

2. or to employ a recursion relation for

$$e^{-x} = \sum_{n=0}^{\infty} s_n = \sum_{n=0}^{\infty} (-1)^n \frac{x^n}{n!}$$

using

$$s_n = -s_{n-1} \frac{x}{n},$$

3. or to first calculate

$$\exp x = \sum_{n=0}^{\infty} s_n$$

and thereafter taking the inverse

$$e^{-x} = \frac{1}{\exp x}$$

Below we have included a small program which calculates

$$e^{-x} = \sum_{n=0}^{\infty} (-1)^n \frac{x^n}{n!},$$

for x -values ranging from 0 to 100 in steps of 10. When doing the summation, we can always define a desired precision, given below by the fixed value for the variable TRUNCATION = $1.0E-10$, so that for a certain value of $x > 0$, there is always a value of $n = N$ for which the loss of precision in terminating the series at $n = N$ is always smaller than the next term in the series $\frac{x^N}{N!}$. The latter is implemented through the while{...} statement.

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/cpp/program4.cpp>

```
// Program to calculate function exp(-x)
// using straightforward summation with differing precision
using namespace std;
#include <iostream>
// type float: 32 bits precision
// type double: 64 bits precision
#define TYPE double
#define PHASE(a) (1 - 2 * (abs(a) % 2))
#define TRUNCATION 1.0E-10
// function declaration
TYPE factorial(int);

int main()
{
    int n;
    TYPE x, term, sum;
    for(x = 0.0; x < 100.0; x += 10.0) {
        sum = 0.0;           //initialization
        n = 0;
        term = 1;
        while(fabs(term) > TRUNCATION) {
            term = PHASE(n) * (TYPE) pow((TYPE) x, (TYPE) n) / factorial(n);
            sum += term;
            n++;
        } // end of while() loop
        cout << `` x = ' ' << x << `` exp = `` << exp(-x) << `` series = `` << sum;
```

```

    cout << `` number of terms = " << n << endl;
} // end of for() loop
return 0;
} // End: function main()

// The function factorial()
// calculates and returns n!

TYPE factorial(int n)
{
    int loop;
    TYPE fac;
    for(loop = 1, fac = 1.0; loop <= n; loop++) {
        fac *= loop;
    }
    return fac;
} // End: function factorial()

```

There are several features to be noted². First, for low values of x , the agreement is good, however for larger x values, we see a significant loss of precision. Secondly, for $x = 70$ we have an overflow problem, represented (from this specific compiler) by NaN (not a number). The latter is easy to understand, since the calculation of a factorial of the size $171!$ is beyond the limit set for the double precision variable factorial. The message NaN appears since the computer sets the factorial of 171 equal to zero and we end up having a division by zero in our expression for e^{-x} .

The overflow problem can be dealt with via a recurrence formula³ for the terms in the sum, so that we avoid calculating factorials. A simple recurrence formula for our equation

$$\exp(-x) = \sum_{n=0}^{\infty} s_n = \sum_{n=0}^{\infty} (-1)^n \frac{x^n}{n!},$$

is to note that

$$s_n = -s_{n-1} \frac{x}{n},$$

so that instead of computing factorials, we need only to compute products. This is exemplified through the next program.

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/cpp/program5.cpp>

```

// program to compute exp(-x) without factorials
using namespace std;

```

²Note that different compilers may give different messages and deal with overflow problems in different ways.

³Recurrence formulae, in various disguises, either as ways to represent series or continued fractions, are among the most commonly used forms for function approximation. Examples are Bessel functions, Hermite and Laguerre polynomials, discussed for example in chapter ??.

x	$\exp(-x)$	Series	Number of terms in series
0.0	0.100000E+01	0.100000E+01	1
10.0	0.453999E-04	0.453999E-04	44
20.0	0.206115E-08	0.487460E-08	72
30.0	0.935762E-13	-0.342134E-04	100
40.0	0.424835E-17	-0.221033E+01	127
50.0	0.192875E-21	-0.833851E+05	155
60.0	0.875651E-26	-0.850381E+09	171
70.0	0.397545E-30	NaN	171
80.0	0.180485E-34	NaN	171
90.0	0.819401E-39	NaN	171
100.0	0.372008E-43	NaN	171

Table 2.3: Result from the brute force algorithm for $\exp(-x)$.

```

#include <iostream>
#define TRUNCATION 1.0E-10

int main()
{
    int    loop, n;
    double x, term, sum;

    for(loop = 0; loop <= 100; loop += 10){
        x  = (double) loop;    // initialization
        sum = 1.0;
        term = 1;
        n  = 1;
        while(fabs(term) > TRUNCATION){
            term *= -x/((double) n);
            sum += term;
            n++;
        } // end while loop
        cout << "x =" << x << "exp = " << exp(-x) << "series = " << sum;
        cout << "number of terms = " << n << endl;
    } // end of for loop
} // End: function main()

```

In this case, we do not get the overflow problem, as can be seen from the large number of terms. Our results do however not make much sense for larger values of x . Decreasing the truncation test will not help! (try it). This is a much more serious problem.

In order better to understand this problem, let us consider the case of $x = 20$,

x	$\exp(-x)$	Series	Number of terms in series
0.000000	0.10000000E+01	0.10000000E+01	1
10.000000	0.45399900E-04	0.45399900E-04	44
20.000000	0.20611536E-08	0.56385075E-08	72
30.000000	0.93576230E-13	-0.30668111E-04	100
40.000000	0.42483543E-17	-0.31657319E+01	127
50.000000	0.19287498E-21	0.11072933E+05	155
60.000000	0.87565108E-26	-0.33516811E+09	182
70.000000	0.39754497E-30	-0.32979605E+14	209
80.000000	0.18048514E-34	0.91805682E+17	237
90.000000	0.81940126E-39	-0.50516254E+22	264
100.000000	0.37200760E-43	-0.29137556E+26	291

Table 2.4: Result from the improved algorithm for $\exp(-x)$.

which already differs largely from the exact result. Writing out each term in the summation, we obtain the largest term in the sum appears at $n = 19$, with a value that equals -43099804 . However, for $n = 20$ we have almost the same value, but with an interchanged sign. It means that we have an error relative to the largest term in the summation of the order of $43099804 \times 10^{-10} \approx 4 \times 10^{-2}$. This is much larger than the exact value of 0.21×10^{-8} . The large contributions which may appear at a given order in the sum, lead to strong roundoff errors, which in turn is reflected in the loss of precision. We can rephrase the above in the following way: Since $\exp(-20)$ is a very small number and each term in the series can be rather large (of the order of 10^8 , it is clear that other terms as large as 10^8 , but negative, must cancel the figures in front of the decimal point and some behind as well. Since a computer can only hold a fixed number of significant figures, all those in front of the decimal point are not only useless, they are crowding out needed figures at the right end of the number. Unless we are very careful we will find ourselves adding up series that finally consists entirely of roundoff errors! An analysis of the contribution to the sum from various terms shows that the relative error made can be huge. This results in an unstable computation, since small errors made at one stage are magnified in subsequent stages.

To this specific case there is a simple cure. Noting that $\exp(x)$ is the reciprocal of $\exp(-x)$, we may use the series for $\exp(x)$ in dealing with the problem of alternating signs, and simply take the inverse. One has however to beware of the fact that $\exp(x)$ may quickly exceed the range of a double variable.

2.4.2 Fortran codes

The Fortran programs are rather similar in structure to the C++ program.

In Fortran Real numbers are written as 2.0 rather than 2 and declared as REAL (KIND=8) or REAL (KIND=4) for double or single precision, respectively. In general we discourage the use of single precision in scientific computing, the achieved precision is in general not good enough. Fortran uses a do construct to have the computer execute the same statements more than once. Note also that Fortran does not allow floating numbers as loop variables. In the example below we use both a do construct for the loop over x and a DO WHILE construction for the truncation test, as in the C++ program. One could alternatively use the EXIT statement inside a do loop. Fortran has also if statements as in C++. The IF construct allows the execution of a sequence of statements (a block) to depend on a condition. The if construct is a compound statement and begins with IF ... THEN and ends with ENDIF. Examples of more general IF constructs using ELSE and ELSEIF statements are given in other program examples. Another feature to observe is the CYCLE command, which allows a loop variable to start at a new value.

Subprograms are called from the main program or other subprograms. In the C++ codes we declared a function `TYPE factorial(int);`. Subprograms are always called functions in C++. If we declare it with `void` it has the same meaning as subroutines in Fortran. Subroutines are used if we have more than one return value. In the example below we compute the factorials using the function `factorial`. This function receives a dummy argument n . `INTENT(IN)` means that the dummy argument cannot be changed within the subprogram. `INTENT(OUT)` means that the dummy argument cannot be used within the subprogram until it is given a value with the intent of passing a value back to the calling program. The statement `INTENT(INOUT)` means that the dummy argument has an initial value which is changed and passed back to the calling program. We recommend that you use these options when calling subprograms. This allows better control when transferring variables from one function to another. In chapter ?? we discuss call by value and by reference in C++. Call by value does not allow a called function to change the value of a given variable in the calling function. This is important in order to avoid unintentional changes of variables when transferring data from one function to another. The `INTENT` construct in Fortran allows such a control. Furthermore, it increases the readability of the program.

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/Fortran/program4.f90>

```
! In this module you can define for example global constants
MODULE constants
  ! definition of variables for double precisions and complex variables
  INTEGER, PARAMETER :: dp = KIND(1.0D0)
  INTEGER, PARAMETER :: dpc = KIND((1.0D0,1.0D0))
```

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```

! Global Truncation parameter
REAL(DP), PARAMETER, PUBLIC :: truncation=1.0E-10
END MODULE constants

! Here you can include specific functions which can be used by
! many subroutines or functions

MODULE functions
CONTAINS
REAL(DP) FUNCTION factorial(n)
  USE CONSTANTS
  INTEGER, INTENT(IN) :: n
  INTEGER :: loop

  factorial = 1.0_dp
  IF ( n > 1 ) THEN
    DO loop = 2, n
      factorial=factorial*loop
    ENDDO
  ENDIF
END FUNCTION factorial
END MODULE functions

! Main program starts here
PROGRAM exp_prog
  USE constants
  USE functions
  IMPLICIT NONE
  REAL (DP) :: x, term, final_sum
  INTEGER :: n, loop_over_x

  ! loop over x-values
  DO loop_over_x=0, 100, 10
    x=loop_over_x
    ! initialize the EXP sum
    final_sum= 0.0_dp; term = 1.0_dp; n = 0
    DO WHILE ( ABS(term) > truncation)
      term = ((-1.0_dp)**n)*(x**n)/ factorial(n)
      final_sum=final_sum+term
      n=n+1
    ENDDO
    ! write the argument x, the exact value, the computed value and n
    WRITE(*,*) x ,EXP(-x), final_sum, n
  ENDDO
END PROGRAM exp_prog

```

The **MODULE** declaration in Fortran allows one to place functions like the one which

calculates the factorials. Note also the usage of the module **constants** where we define double and complex variables. If one wishes to switch to another precision, one just needs to change the declaration in one part of the program only. This hinders possible errors which arise if one has to change variable declarations in every function and subroutine. In addition we have defined a global variable **truncation** which is accessible to all functions which have the `USE constants` declaration. These declarations have to come before any variable declarations and `IMPLICIT NONE` statement.

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/Fortran/program5.f90>

```
! In this module you can define for example global constants
MODULE constants
  ! definition of variables for double precisions and complex variables
  INTEGER, PARAMETER :: dp = KIND(1.0D0)
  INTEGER, PARAMETER :: dpc = KIND((1.0D0,1.0D0))
  ! Global Truncation parameter
  REAL(DP), PARAMETER, PUBLIC :: truncation=1.0E-10
END MODULE constants

PROGRAM improved_exp
  USE constants
  IMPLICIT NONE
  REAL (dp) :: x, term, final_sum
  INTEGER :: n, loop_over_x

  ! loop over x-values, no floats as loop variables
  DO loop_over_x=0, 100, 10
    x=loop_over_x
    ! initialize the EXP sum
    final_sum=1.0 ; term=1.0 ; n = 1
    DO WHILE ( ABS(term) > truncation)
      term = -term*x/FLOAT(n)
      final_sum=final_sum+term
      n=n+1
    ENDDO
    ! write the argument x, the exact value, the computed value and n
    WRITE(*,*) x ,EXP(-x), final_sum, n
  ENDDO
END PROGRAM improved_exp
```


2.4.3 Further examples

Summing $1/n$

Let us look at another roundoff example which may surprise you more. Consider the series

$$s_1 = \sum_{n=1}^N \frac{1}{n},$$

which is finite when N is finite. Then consider the alternative way of writing this sum

$$s_2 = \sum_{n=N}^1 \frac{1}{n},$$

which when summed analytically should give $s_2 = s_1$. Because of roundoff errors, numerically we will get $s_2 \neq s_1$! Computing these sums with single precision for $N = 1.000.000$ results in $s_1 = 14.35736$ while $s_2 = 14.39265$! Note that these numbers are machine and compiler dependent. With double precision, the results agree exactly, however, for larger values of N , differences may appear even for double precision. If we choose $N = 10^8$ and employ double precision, we get $s_1 = 18.9978964829915355$ while $s_2 = 18.9978964794618506$, and one notes a difference even with double precision.

This example demonstrates two important topics. First we notice that the chosen precision is important, and we will always recommend that you employ double precision in all calculations with real numbers. Secondly, the choice of an appropriate algorithm, as also seen for e^{-x} , can be of paramount importance for the outcome.

The standard algorithm for the standard deviation

Yet another example is the calculation of the standard deviation σ when σ is small compared to the average value \bar{x} . Below we illustrate how one of the most frequently used algorithms can go wrong when single precision is employed.

However, before we proceed, let us define σ and \bar{x} . Suppose we have a set of N data points, represented by the one-dimensional array $x(i)$, for $i = 1, N$. The average value is then

$$\bar{x} = \frac{\sum_{i=1}^N x(i)}{N},$$

while

$$\sigma = \sqrt{\frac{\sum_i x(i)^2 - \bar{x} \sum_i x(i)}{N - 1}}.$$

Let us now assume that

$$x(i) = i + 10^5,$$

and that $N = 127$, just as a mere example which illustrates the kind of problems which can arise when the standard deviation is small compared with the mean value \bar{x} .

The standard algorithm computes the two contributions to σ separately, that is we sum $\sum_i x(i)^2$ and subtract thereafter $\bar{x} \sum_i x(i)$. Since these two numbers can become nearly equal and large, we may end up in a situation with potential loss of precision as an outcome.

The second algorithm on the other hand computes first $x(i) - \bar{x}$ and then squares it when summing up. With this recipe we may avoid having nearly equal numbers which cancel.

Using single precision results in a standard deviation of $\sigma = 40.05720139$ for the first and most used algorithm, while the exact answer is $\sigma = 36.80579758$, a number which also results from the above second algorithm. With double precision, the two algorithms result in the same answer.

The reason for such a difference resides in the fact that the first algorithm includes the subtraction of two large numbers which are squared. Since the average value for this example is $\bar{x} = 100063.00$, it is easy to see that computing $\sum_i x(i)^2 - \bar{x} \sum_i x(i)$ can give rise to very large numbers with possible loss of precision when we perform the subtraction. To see this, consider the case where $i = 64$. Then we have

$$x_{64}^2 - \bar{x}x_{64} = 100352,$$

while the exact answer is

$$x_{64}^2 - \bar{x}x_{64} = 100064!$$

You can even check this by calculating it by hand.

The second algorithm computes first the difference between $x(i)$ and the average value. The difference gets thereafter squared. For the second algorithm we have for $i = 64$

$$x_{64} - \bar{x} = 1,$$

and we have no potential for loss of precision.

The standard text book algorithm is expressed through the following program, where we have also added the second algorithm

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/cpp/program6.cpp>

```
// program to calculate the mean and standard deviation of
// a user created data set stored in array x[]
using namespace std;
#include <iostream>
int main()
{
    int    i;
    float  sum, sumsq2, xbar, sigma1, sigma2;
    // array declaration with fixed dimension
```

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```

float x[127];
// initialise the data set
for ( i=0; i < 127 ; i++){
    x[i] = i + 100000.;
}
// The variable sum is just the sum over all elements
// The variable sumsq2 is the sum over x^2
sum=0.;
sumsq2=0.;
// Now we use the text book algorithm
for ( i=0; i < 127; i++){
    sum += x[i];
    sumsq2 += pow((double) x[i],2.);
}
// calculate the average and sigma
xbar=sum/127.;
sigma1=sqrt((sumsq2-sum*xbar)/126.);
/*
** Here comes the second algorithm where we evaluate
** separately first the average and thereafter the
** sum which defines the standard deviation. The average
** has already been evaluated through xbar
*/
sumsq2=0.;
for ( i=0; i < 127; i++){
    sumsq2 += pow( (double) (x[i]-xbar),2.);
}
sigma2=sqrt(sumsq2/126.);
cout << "xbar = " << xbar << "sigma1 = " << sigma1 << "sigma2 = " << sigma2;
cout << endl;
return 0;
} // End: function main()

```

The corresponding Fortran program is given below.

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/Fortran/program6.f90>

```

PROGRAM standard_deviation
IMPLICIT NONE
REAL (KIND = 4) :: sum, sumsq2, xbar
REAL (KIND = 4) :: sigma1, sigma2
REAL (KIND = 4), DIMENSION (127) :: x
INTEGER :: i

x=0;
DO i=1, 127
    x(i) = i + 100000.
ENDDO
sum=0.; sumsq2=0.
!    standard deviation calculated with the first algorithm

```

```

DO i=1, 127
    sum = sum +x(i)

    sumsq2 = sumsq2+x(i)**2
ENDDO
!    average
xbar=sum/127.
sigma1=SQRT((sumsq2-sum*xbar)/126.)
!    second algorithm to evaluate the standard deviation
sumsq2=0.
DO i=1, 127
    sumsq2=sumsq2+(x(i)-xbar)**2
ENDDO
sigma2=SQRT(sumsq2/126.)
WRITE(*,*) xbar, sigma1, sigma2

END PROGRAM standard_deviation

```

2.5 Additional Features of C++ and Fortran

2.5.1 Operators in C++

In the previous program examples we have seen several types of operators. In the tables below we summarize the most important ones. Note that the modulus in C++ is represented by the operator % whereas in Fortran we employ the intrinsic function MOD. Note also that the increment operator ++ and the decrement operator -- is not available in Fortran. In C++ these operators have the following meaning

++x; or x++; has the same meaning as x = x + 1;
 --x; or x--; has the same meaning as x = x - 1;

Table 2.5 lists several relational and arithmetic operators. Logical operators in C++ and Fortran are listed in 2.6. while Table 2.7 shows bitwise operations.

C++ offers also interesting possibilities for combined operators. These are collected in Table 2.8.

Finally, we show some special operators pertinent to C++ only. The first one is the ? operator. Its action can be described through the following example

A = expression1 ? expression2 : expression3;

Here expression1 is computed first. If this is "true" ($\neq 0$), then expression2 is computed and assigned A. If expression1 is "false", then expression3 is computed and assigned A.

arithmetic operators		relation operators	
operator	effect	operator	effect
−	Subtraction	>	Greater than
+	Addition	>=	Greater or equal
*	Multiplication	<	Less than
/	Division	<=	Less or equal
% or MOD	Modulus division	==	Equal
−−	Decrement	!=	Not equal
++	Increment		

Table 2.5: Relational and arithmetic operators. The relation operators act between two operands. Note that the increment and decrement operators ++ and −− are not available in Fortran .

Logical operators		
C++	Effect	Fortran
0	False value	.FALSE.
1	True value	.TRUE.
!x	Logical negation	.NOT.x
x&& y	Logical AND	x.AND.y
x y	Logical inclusive OR	x.OR.y

Table 2.6: List of logical operators in C++ and Fortran .

Bitwise operations		
C++	Effect	Fortran
~i	Bitwise complement	NOT(j)
i&j	Bitwise and	IAND(i,j)
i^j	Bitwise exclusive or	IEOR(i,j)
i j	Bitwise inclusive or	IOR(i,j)
i<<j	Bitwise shift left	ISHFT(i,j)
i>>n	Bitwise shift right	ISHFT(i,-j)

Table 2.7: List of bitwise operations.

Expression	meaning	expression	meaning
a += b;	a = a + b;	a -= b;	a = a - b;
a *= b;	a = a * b;	a /= b;	a = a / b;
a %= b;	a = a % b;	a <= b;	a = a < b;
a >= b;	a = a > b;	a &= b;	a = a & b;
a = b;	a = a b;	a ^= b;	a = a ^ b;

Table 2.8: C++ specific expressions.

2.5.2 Pointers and arrays in C++.

In addition to constants and variables C++ contain important types such as pointers and arrays (vectors and matrices). These are widely used in most C++ program. C++ allows also for pointer algebra, a feature not included in Fortran . Pointers and arrays are important elements in C++. To shed light on these types, consider the following setup

<code>int name</code>	defines an integer variable called name. It is given an address in memory where we can store an integer number.
<code>&name</code>	is the address of a specific place in memory where the integer name is stored. Placing the operator & in front of a variable yields its address in memory.
<code>int *pointer</code>	defines an integer pointer and reserves a location in memory for this specific variable The content of this location is viewed as the address of another place in memory where we have stored an integer.

Note that in C++ it is common to write `int* pointer` while in C one usually writes `int *pointer`. Here are some examples of legal C++ expressions.

```
name = 0x56;           /* name gets the hexadecimal value hex 56. */
pointer = &name;       /* pointer points to name. */
printf("Address of name = %p",pointer); /* writes out the address of name. */
printf("Value of name= %d",*pointer);  /* writes out the value of name. */
```

Here's a program which illustrates some of these topics.

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/cpp/program7.cpp>

```
1 using namespace std;
2 main()
3 {
```

```

4   int var;
5   int *pointer;
6
7   pointer = &var;
8   var = 421;
9   printf("Address of the integer variable var : %p\n",&var);
10  printf("Value of var : %d\n", var);
11  printf("Value of the integer pointer variable: %p\n",pointer);
12  printf("Value which pointer is pointing at : %d\n",*pointer);
13  printf("Address of the pointer variable : %p\n",&pointer);
14 }

```

Line	Comments
4	• Defines an integer variable var.
5	• Define an integer pointer – reserves space in memory.
7	• The content of the address of pointer is the address of var.
8	• The value of var is 421.
9	• Writes the address of var in hexadecimal notation for pointers %p.
10	• Writes the value of var in decimal notation %d.

The output of this program, compiled with g++, reads @@

Address of the integer variable var : 0xbfffeb74

Value of var: 421

Value of integer pointer variable : 0xbfffeb74

The value which pointer is pointing at : 421

Address of the pointer variable : 0xbfffeb70

In the next example we consider the link between arrays and pointers.

```

int matr[2]    defines a matrix with two integer members – matr[0] og
               matr[1].
matr           is a pointer to matr[0].
(matr + 1)     is a pointer to matr[1].

```

<http://folk.uio.no/mhjensen/compphys/programs/chapter02/cpp/program8.cpp>

```

1  using namespace std;
2  #included <iostream>
3  int main()
4  {
5      int matr[2];
6      int *pointer;
7      pointer = &matr[0];

```

```

8     matr[0] = 321;
9     matr[1] = 322;
10    printf("\nAddress of the matrix element matr[1]: %p",&matr[0]);
11    printf("\nValue of the matrix element matr[1]; %d",matr[0]);
12    printf("\nAddress of the matrix element matr[2]: %p",&matr[1]);
13    printf("\nValue of the matrix element matr[2]: %d\n", matr[1]);
14    printf("\nValue of the pointer : %p",pointer);
15    printf("\nValue which pointer points at : %d",*pointer);
16    printf("\nValue which (pointer+1) points at: %d\n",*(pointer+1));
17    printf("\nAddress of the pointer variable: %p\n",&pointer);
18 }

```

You should especially pay attention to the following

Line	
5	• Declaration of an integer array matr with two elements
6	• Declaration of an integer pointer
7	• The pointer is initialized to point at the first element of the array matr.
8-9	• Values are assigned to the array matr.

The output of this example, compiled again with g++, is @@

```

Address of the matrix element matr[1]: 0xbffef70
Value of the matrix element matr[1]; 321
Address of the matrix element matr[2]: 0xbffef74
Value of the matrix element matr[2]: 322
Value of the pointer: 0xbffef70
The value pointer points at: 321
The value that (pointer+1) points at: 322
Address of the pointer variable : 0xbffef6c

```

2.5.3 Macros in C++

In C we can define macros, typically global constants or functions through the define statements shown in the simple C-example below for

```

1. #define ONE 1
2. #define TWO ONE + ONE
3. #define THREE ONE + TWO
4.
5. main()
6. {
7.     printf("ONE=%d, TWO=%d, THREE=%d",ONE,TWO,THREE);
8. }

```


In C++ the usage of macros is discouraged and you should rather use the declaration for constant variables. You would then replace a statement like `#define ONE 1` with `const int ONE = 1;`. There is typically much less use of macros in C++ than in C. C++ allows also the definition of our own types based on other existing data types. We can do this using the keyword `typedef`, whose format is: `typedef existing_type new_type_name ;`, where `existing_type` is a C++ fundamental or compound type and `new_type_name` is the name for the new type we are defining. For example:

```
typedef char new_name;
typedef unsigned int word ;
typedef char * test;
typedef char field [50];
```

In this case we have defined four data types: `new_name`, `word`, `test` and `field` as `char`, `unsigned int`, `char*` and `char[50]` respectively, that we could perfectly use in declarations later as any other valid type

```
new_name mychar, anotherchar, *ptc1;
word myword;
test ptc2;
field name;
```

The use of `typedef` does not create different types. It only creates synonyms of existing types. That means that the type of `myword` can be considered to be either `word` or `unsigned int`, since both are in fact the same type. Using `typedef` allows to define an alias for a type that is frequently used within a program. It is also useful to define types when it is possible that we will need to change the type in later versions of our program, or if a type you want to use has a name that is too long or confusing.

In C we could define macros for functions as well, as seen below.

```
1. #define MIN(a,b) ( ((a) < (b)) ? (a) : (b) )
2. #define MAX(a,b) ( ((a) > (b)) ? (a) : (b) )
3. #define ABS(a)   ( ((a) < 0) ? -(a) : (a) )
4. #define EVEN(a)  ( (a) %2 == 0 ? 1 : 0 )
5. #define TOASCII(a) ( (a) & 0x7f )
```

In C++ we would replace such function definition by employing so-called `inline` functions. The above functions could then read

```
inline double MIN(double a, double b) {return ((a)<(b)) ? (a):(b);}
inline double MAX(double a, double b){return ((a)>(b)) ? (a):(b);}
inline double ABS(double a) {return ((a)<0) ? -(a):(a);}
```

where we have defined the transferred variables to be of type `double`. The functions also return a `double` type. These functions could easily be generalized through the use of classes and templates, see chapter ??, to return whatever types of real, complex or integer variables.

Inline functions are very useful, especially if the overhead for calling a function implies a significant fraction of the total function call cost. When such function call overhead is significant, a function definition can be preceded by the keyword `inline`. When this function is called, we expect the compiler to generate inline code without function call overhead. However, although inline functions eliminate function call overhead, they can introduce other overheads. When a function is inlined, its code is duplicated for each call. Excessive use of `inline` may thus generate large programs. Large programs can cause excessive paging in virtual memory systems. Too many inline functions can also lengthen compile and link times, on the other hand not inlining small functions like the above that do small computations, can make programs bigger and slower. However, most modern compilers know better than programmer which functions to inline or not. When doing this, you should also test various compiler options. With the compiler option `-O3` inlining is done automatically by basically all modern compilers.

A good strategy, recommended in many C++ textbooks, is to write a code without inline functions first. As we also suggested in the introductory chapter, you should first write a as simple and clear as possible program, without a strong emphasis on computational speed. Thereafter, when profiling the program one can spot small functions which are called many times. These functions can then be candidates for inlining. If the overall time consumption is reduced due to inlining specific functions, we can proceed to other sections of the program which could be speeded up.

Another problem with inlined functions is that on some systems debugging an inline function is difficult because the function does not exist at runtime.

2.5.4 Structures in C++ and TYPE in Fortran

A very important part of a program is the way we organize our data and the flow of data when running the code. This is often a neglected aspect especially during the development of an algorithm. A clear understanding of how data are represented makes the program more readable and easier to maintain and extend upon by other users. Till now we have studied elementary variable declarations through keywords like `int` or `INTEGER`, `double` or `REAL(KIND(8))` and `char` or its Fortran equivalent `CHARACTER`. These declarations could also be extended to general multi-dimensional arrays.

However, C++ and Fortran offer other ways as well by which we can organize our data in a more transparent and reusable way. One of these options is through the `struct` declaration of C++, or the correspondingly similar `TYPE` in Fortran. The latter data type will also be discussed in chapter ??.

The following example illustrates how we could make a general variable which can be reused in defining other variables as well.

Suppose you would like to make a general program which treats quantum mechanical problems from both atomic physics and nuclear physics. In atomic and nuclear physics the single-particle degrees are represented by quantum numbers such orbital angular momentum, total angular momentum, spin and energy. An independent particle model is often assumed as the starting point for building up more complicated many-body correlations in systems with many interacting particles. In atomic physics the effective degrees of freedom are often reduced to electrons interacting with each other, while in nuclear physics the system is described by neutrons and protons. The structure `single_particle_descript` contains a list over different quantum numbers through various pointers which are initialized by a calling function.

```
struct single_particle_descript{
    int total_states;
    int* n;
    int* lorb;
    int* m_l;
    int* jang;
    int* spin;
    double* energy;
    char* orbit_status
};
```

To describe an atom like Neon we would need three single-particle orbits to describe the ground state wave function if we use a single-particle picture, i.e., the $1s$, $2s$ and $2p$ single-particle orbits. These orbits have a degeneracy of $2(2l + 1)$, where the first number stems from the possible spin projections and the second from the possible projections of the orbital momentum. Note that we reserve the naming orbit for the generic labelling $1s$, $2s$ and $2p$ while we use the naming states when we include all possible quantum numbers. In total there are 10 possible single-particle states when we account for spin and orbital momentum projections. In this case we would thus need to allocate memory for arrays containing 10 elements.

The above structure is written in a generic way and it can be used to define other variables as well. For electrons we could write `struct single_particle_descript electrons;` and is a new variable with the name `electrons` containing all the elements of this structure.

The following program segment illustrates how we access these elements To access these elements we could for example read from a given device the various quantum numbers:

```
for ( int i = 0; i < electrons.total_states; i++){
    cout << `` Read in the quantum numbers for electron i: `` << i << endl;
    cin >> electrons.n[i];
    cin > electrons.lorb[i];
    cin >> electrons.m_l[i];
```

```

    cin >> electrons.jang[i];
    cin >> electrons.spin[i];
}

```

The structure `single_particle_descript` can also be used for defining quantum numbers of other particles as well, such as neutrons and protons through the new variables `struct single_particle_descript protons` and `struct single_particle_descript`.

The corresponding declaration in Fortran is given by the `TYPE` construct, seen in the following example.

```

TYPE, PUBLIC :: single_particle_descript
  INTEGER :: total_states
  INTEGER, DIMENSION(:), POINTER :: n, lorb, jang, spin, m_l
  CHARACTER (LEN=10), DIMENSION(:), POINTER :: orbit_status
  REAL(8), DIMENSION(:), POINTER :: energy
END TYPE single_particle_descript

```

This structure can again be used to define variables like `electrons`, `protons` and `neutrons` through the statement `TYPE (single_particle_descript) :: electrons, protons`. More detailed examples on the use of these variable declarations, classes and templates will be given in subsequent chapters.

2.6 Reading and writing to file

Furthermore, we will use this section to introduce three important C++-programming features, namely reading and writing to a file, call by reference and call by value, and dynamic memory allocation. We are also going to split the tasks performed by the program into subtasks. We define one function which reads in the input data, one which calculates the second derivative and a final function which writes the results to file.

Let us look at a simple case first, the use of `printf` and `scanf`. If we wish to print a variable defined as `double speed_of_sound`; we could for example write

```

double speed_of_sound;
.....
printf("`speed_of_sound = %lf\n'", speed_of_sound);

```

In this case we say that we transfer the value of this specific variable to the function `printf`. The function `printf` *can however not change the value of this variable* (there is no need to do so in this case). Such a call of a specific function is called *call by value*. The crucial aspect to keep in mind is that the value of this specific variable does not change in the called function.

When do we use call by value? And why care at all? We do actually care, because if a called function has the possibility to change the value of a variable when this

is not desired, calling another function with this variable may lead to totally wrong results. In the worst cases you may even not be able to spot where the program goes wrong.

We do however use call by value when a called function simply receives the value of the given variable without changing it.

If we however wish to update the value of say an array in a called function, we refer to this call as **call by reference**. What is transferred then is the address of the first element of the array, and the called function has now access to where that specific variable 'lives' and can thereafter change its value.

The function `scanf` is then an example of a function which receives the address of a variable and is allowed to modify it. Afterall, when calling `scanf` we are expecting a new value for a variable. A typical call could be `scanf("%lf\n", &speed_of_sound);`.

Consider now the following program

```
1 using namespace std;
2 # include <iostream>
3 // begin main function
4 int main(int argc, char argv[])
5 {
6     int a;
7     int *b;
8     a = 10;
9     b = new int[10];
10    for( int i = 0; i < 10; i++){
11        b[i] = i;
12    }
13    func(a,b);
14    return 0;
15 } // end of main function
16 // definition of the function func
17 void func(int x, int *y)
18 {
19     x += 7;
20     *y += 10;
21     y[6] += 10;
22     return;
23 } // end function func
```

There are several features to be noted.

- Lines 5 and 6: Declaration of two variables `a` and `b`. The compiler reserves two locations in memory. The size of the location depends on the type of variable. Two properties are important for these locations – the address in memory and the content in the
- Line 7: The value of `a` is now 10.

- Line 8: Memory to store 10 integers is reserved. The address to the first location is stored in b. The address of element number 6 is given by the expression (b + 6).
- Line 10: All 10 elements of b are given values: b[0] = 0, b[1] = 1,, b[9] = 9;
- Line 12: The main() function calls the function func() and the program counter transfers to the first statement in func(). With respect to data the following happens. The content of a (= 10) and the content of b (a memory address) are copied to a stack (new memory location) associated with the function func()
- Line 16: The variable x and y are local variables in func(). They have the values – x = 10, y = address of the first element in b in the main() program.
- Line 18: The local variable x stored in the stack memory is changed to 17. Nothing happens with the value a in main().
- Line 19: The value of y is an address and the symbol *y stands for the position in memory which has this address. The value in this location is now increased by 10. This means that the value of b[0] in the main program is equal to 10. Thus func() has modified a value in main().
- Line 20: This statement has the same effect as line 9 except that it modifies element b[6] in main() by adding a value of 10 to what was there originally, namely 6.
- Line 21: The program counter returns to main(), the next expression after *func(a,b)*;. All data on the stack associated with func() are destroyed.
- The value of a is transferred to func() and stored in a new memory location called x. Any modification of x in func() does not affect in any way the value of a in main(). This is called **transfer of data by value**. On the other hand the next argument in func() is an address which is transferred to func(). This address can be used to modify the corresponding value in main(). In the programming language C it is expressed as a modification of the value which y points to, namely the first element of b. This is called **transfer of data by reference** and is a method to transfer data back to the calling function, in this case main().

C++ allows however the programmer to use solely call by reference (note that call by reference is implemented as pointers). To see the difference between C and C++, consider the following simple examples. In C we would write

```
int n; n = 8;
func(&n); /* &n is a pointer to n */
....
```

```
void func(int *i)
{
    *i = 10; /* n is changed to 10 */
    ....
}
```

whereas in C++ we would write

```
int n; n =8;
func(n); // just transfer n itself
....
void func(int& i)
{
    i = 10; // n is changed to 10
    ....
}
```

Note well that the way we have defined the input to the function `func(int& i)` or `func(int *i)` decides how we transfer variables to a specific function. The reason why we emphasize the difference between call by value and call by reference is that it allows the programmer to avoid pitfalls like unwanted changes of variables. However, many people feel that this reduces the readability of the code. It is more or less common in C++ to use call by reference, since it gives a much cleaner code. Recall also that behind the curtain references are usually implemented as pointers. When we transfer large objects such a matrices and vectors one should always use call by reference. Copying such objects to a called function slows down considerably the execution. If you need to keep the value of a call by reference object, you should use the `const` declaration.

In programming languages like Fortran one uses only call by reference, but you can flag whether a called function or subroutine is allowed or not to change the value by declaring for example an integer value as `INTEGER, INTENT(IN) :: i`. The local function cannot change the value of *i*. Declaring a transferred values as `INTEGER, INTENT(OUT) :: i`. allows the local function to change the variable *i*.

Initializations and main program

In every program we have to define the functions employed. The style chosen here is to declare these functions at the beginning, followed thereafter by the main program and the detailed tasks performed by each function. Another possibility is to include these functions and their statements before the main program, meaning that the main program appears at the very end. I find this programming style less readable however since I prefer to read a code from top to bottom. A further option, specially in connection with larger projects, is to include these function definitions in a user defined header file. The following program shows also (although it is rather unnec-

essary in this case due to few tasks) how one can split different tasks into specialized functions. Such a division is very useful for larger projects and programs.

In the first version of this program we use a more C-like style for writing and reading to file. At the end of this section we include also the corresponding C++ and Fortran files.

<http://folk.uio.no/mhjensen/compphys/programs/chapter03/cpp/program1.cpp>

```

/*
**  Program to compute the second derivative of exp(x).
**  Three calling functions are included
**  in this version. In one function we read in the data from screen,
**  the next function computes the second derivative
**  while the last function prints out data to screen.
*/
using namespace std;
# include <iostream>

void initialize (double *, double *, int *);
void second_derivative( int, double, double, double *, double *);
void output( double *, double *, double, int);

int main()
{
    // declarations of variables
    int number_of_steps;
    double x, initial_step;
    double *h_step, *computed_derivative;
    // read in input data from screen
    initialize (&initial_step, &x, &number_of_steps);
    // allocate space in memory for the one-dimensional arrays
    // h_step and computed_derivative
    h_step = new double[number_of_steps];
    computed_derivative = new double[number_of_steps];
    // compute the second derivative of exp(x)
    second_derivative( number_of_steps, x, initial_step, h_step,
                      computed_derivative);
    // Then we print the results to file
    output(h_step, computed_derivative, x, number_of_steps );
    // free memory
    delete [] h_step;
    delete [] computed_derivative;
    return 0;
} // end main program

```

We have defined three additional functions, one which reads in from screen the value of x , the initial step length h and the number of divisions by 2 of h . This function is called `initialize`. To calculate the second derivatives we define the function `second_derivative`. Finally, we have a function which writes our results together

with a comparison with the exact value to a given file. The results are stored in two arrays, one which contains the given step length h and another one which contains the computed derivative.

These arrays are defined as pointers through the statement

```
double *h_step, *computed_derivative;
```

A call in the main function to the function `second_derivative` looks then like this

```
second_derivative( number_of_steps, x, initial_step, h_step, computed_derivative);
```

while the called function is declared in the following way

```
void second_derivative(int number_of_steps, double x, double *h_step, double
    *computed_derivative);
```

indicating that `double *h_step, double *computed_derivative;` are pointers and that we transfer the address of the first elements. The other variables `int number_of_steps` are transferred by value and are not changed in the called function.

Another aspect to observe is the possibility of dynamical allocation of memory through the new function. In the included program we reserve space in memory for these three arrays in the following way

```
h_step = new double[number_of_steps];
computed_derivative = new double[number_of_steps];
```

When we no longer need the space occupied by these arrays, we free memory through the declarations

```
delete [] h_step;
delete [] computed_derivative;
```

The function initialize

```
// Read in from screen the initial step, the number of steps
// and the value of x

void initialize (double *initial_step, double *x, int *number_of_steps)
{
    printf("Read in from screen initial step, x and number of steps\n");
    scanf("%lf %lf %d", initial_step, x, number_of_steps);
    return;
} // end of function initialize
```

This function receives the addresses of the three variables

```
void initialize (double *initial_step, double *x, int *number_of_steps)
```

and returns updated values by reading from screen.

The function `second_derivative`

```
// This function computes the second derivative

void second_derivative( int number_of_steps, double x,
                       double initial_step, double *h_step,
                       double *computed_derivative)
{
    int counter;
    double h;
    // calculate the step size
    // initialize the derivative, y and x (in minutes)
    // and iteration counter
    h = initial_step;
    // start computing for different step sizes
    for (counter=0; counter < number_of_steps; counter++ )
    {
        // setup arrays with derivatives and step sizes
        h_step[counter] = h;
        computed_derivative[counter] =
            (exp(x+h)-2.*exp(x)+exp(x-h))/(h*h);
        h = h*0.5;
    } // end of do loop
    return;
} // end of function second_derivative
```

The loop over the number of steps serves to compute the second derivative for different values of h . In this function the step is halved for every iteration (you could obviously change this to larger or smaller step variations). The step values and the derivatives are stored in the arrays `h_step` and `double computed_derivative`.

The output function

This function computes the relative error and writes the results to a chosen file.

The last function here illustrates how to open a file, write and read possible data and then close it. In this case we have fixed the name of the file. Another possibility is obviously to read the name of this file together with other input parameters. The way the program is presented here is slightly unpractical since we need to recompile the program if we wish to change the name of the output file.

An alternative is represented by the following C++ program. This program reads from screen the names of the input and output files.

<http://folk.uio.no/mhjensen/compphys/programs/chapter03/cpp/program2.cpp>

```
1 #include <stdio.h>
2 #include <stdlib.h>
3 int col:
```

```

4
5 int main(int argc, char *argv[])
6 {
7     FILE *inn, *out;
8     int c;
9     if( argc < 3) {
10    printf("You have to read in :\n");
11    printf("in_file and out_file \n");
12    exit(1);
13    inn = fopen( argv[1], "r"); // returns pointer to the in_file
14    if( inn == NULL ) { // can't find in_file
15        printf("Can't find the input file %s\n", argv[1]);
16        exit(1);
17    }
18    out = fopen( argv[2], "w"); // returns a pointer to the out_file
19    if( out == NULL ) { // can't find out_file
20        printf("Can't find the output file %s\n", argv[2]);
21        exit(1);
22    }
23    ... program statements
24
25    fclose(inn);
26    fclose(out);
27    return 0;
28 }

```

This program has several interesting features.

Line	Program comments
5	<ul style="list-style-type: none"> • The function <code>main()</code> takes three arguments, given by <code>argc</code>. The variable <code>argv</code> points to the following: the name of the program, the first and second arguments, in this case the file names to be read from screen.
7	<ul style="list-style-type: none"> • C++ has a data type called <code>FILE</code>. The pointers <code>inn</code> and <code>out</code> point to specific files. They must be of the type <code>FILE</code>.
10	<ul style="list-style-type: none"> • The command line has to contain 2 filenames as parameters.
13-17	<ul style="list-style-type: none"> • The input file has to exist, else the pointer returns <code>NULL</code>. It has only read permission.
18-22	<ul style="list-style-type: none"> • This applies for the output file as well, but now with write permission only.
23-24	<ul style="list-style-type: none"> • Both files are closed before the main program ends.

The main part of the code includes now an object declaration `ofstream ofile` which is included in C++ and allows the programmer to open and declare files. This is done via the statement `ofile.open(outfilename);`. We close the file at the

end of the main program by writing `ofile.close();`. There is a corresponding object for reading inputfiles. In this case we declare prior to the main function, or in an eventual header file, `ifstream ifile` and use the corresponding statements `ifile.open(infilename);` and `ifile.close();` for opening and closing an input file. Note that we have declared two character variables `char* outfilename;` and `char* infilename;`. In order to use these options we need to include a corresponding library of functions using `# include <fstream>`.

One of the problems with C++ is that formatted output is not as easy to use as the `printf` and `scanf` functions in C. The output function using the C++ style is included below.

```
// function to write out the final results
void output(double *h_step, double *computed_derivative, double x,
            int number_of_steps )
{
    int i;
    ofile << "RESULTS:" << endl;
    ofile << setiosflags(ios::showpoint | ios::uppercase);
    for( i=0; i < number_of_steps; i++)
    {
        ofile << setw(15) << setprecision(8) << log10(h_step[i]);
        ofile << setw(15) << setprecision(8) <<
            log10(fabs(computed_derivative[i]-exp(x))/exp(x))) << endl;
    }
} // end of function output
```

The function `setw(15)` reserves an output of 15 spaces for a given variable while `setprecision(8)` yields eight leading digits. To use these options you have to use the declaration `# include <iomanip>`.

Before we discuss the results of our calculations we list here the corresponding Fortran program. The corresponding Fortran example is

<http://folk.uio.no/mhjensen/compphys/programs/chapter03/Fortran/program1.f90>

```
! Program to compute the second derivative of exp(x).
! Only one calling function is included.
! It computes the second derivative and is included in the
! MODULE functions as a separate method
! The variable h is the step size. We also fix the total number
! of divisions by 2 of h. The total number of steps is read from
! screen
MODULE constants
    ! definition of variables for double precisions and complex variables
    INTEGER, PARAMETER :: dp = KIND(1.0D0)
    INTEGER, PARAMETER :: dpc = KIND((1.0D0,1.0D0))
END MODULE constants

! Here you can include specific functions which can be used by
```

```

! many subroutines or functions

MODULE functions
USE constants
IMPLICIT NONE
CONTAINS
  SUBROUTINE derivative(number_of_steps, x, initial_step, h_step, &
    computed_derivative)
    USE constants
    INTEGER, INTENT(IN) :: number_of_steps
    INTEGER :: loop
    REAL(DP), DIMENSION(number_of_steps), INTENT(INOUT) :: &
      computed_derivative, h_step
    REAL(DP), INTENT(IN) :: initial_step, x
    REAL(DP) :: h
    ! calculate the step size
    ! initialize the derivative, y and x (in minutes)
    ! and iteration counter
    h = initial_step
    ! start computing for different step sizes
    DO loop=1, number_of_steps
      ! setup arrays with derivatives and step sizes
      h_step(loop) = h
      computed_derivative(loop) = (EXP(x+h)-2.*EXP(x)+EXP(x-h))/(h*h)
      h = h*0.5
    ENDDO
  END SUBROUTINE derivative

END MODULE functions

PROGRAM second_derivative
  USE constants
  USE functions
  IMPLICIT NONE
  ! declarations of variables
  INTEGER :: number_of_steps, loop
  REAL(DP) :: x, initial_step
  REAL(DP), ALLOCATABLE, DIMENSION(:) :: h_step, computed_derivative
  ! read in input data from screen
  WRITE(*,*) 'Read in initial step, x value and number of steps'
  READ(*,*) initial_step, x, number_of_steps
  ! open file to write results on
  OPEN(UNIT=7,FILE='out.dat')
  ! allocate space in memory for the one-dimensional arrays
  ! h_step and computed_derivative
  ALLOCATE(h_step(number_of_steps),computed_derivative(number_of_steps))
  ! compute the second derivative of exp(x)
  ! initialize the arrays
  h_step = 0.0_dp; computed_derivative = 0.0_dp

```

```

CALL derivative(number_of_steps,x,initial_step,h_step,computed_derivative)

! Then we print the results to file
DO loop=1, number_of_steps
  WRITE(7, '(E16.10,2X,E16.10)') LOG10(h_step(loop)),&
    LOG10 ( ABS ( (computed_derivative(loop)-EXP(x))/EXP(x)))
ENDDO
! free memory
DEALLOCATE( h_step, computed_derivative)
! close the output file
CLOSE(7)

END PROGRAM second_derivative

```

The `MODULE` declaration in Fortran allows one to place functions like the one which calculates second derivatives in a module. Since this is a general method, one could extend its functionality by simply transferring the name of the function to differentiate. In our case we use explicitly the exponential function, but there is nothing which hinders us from defining other functions. Note the usage of the module **constants** where we define double and complex variables. If one wishes to switch to another precision, one needs to change the declaration in one part of the program only. This hinders possible errors which arise if one has to change variable declarations in every function and subroutine. Finally, dynamic memory allocation and deallocation is in Fortran done with the keywords `ALLOCATE(array(size))` and `DEALLOCATE(array)`. Although most compilers deallocate and thereby free space in memory when leaving a function, you should always deallocate an array when it is no longer needed. In case your arrays are very large, this may block unnecessarily large fractions of the memory. Furthermore, you should always initialize arrays. In the example above, we note that Fortran allows us to simply write `h_step = 0.0_dp; computed_derivative = 0.0_dp`, which means that all elements of these two arrays are set to zero. Coding arrays in this manner brings us much closer to the way we deal with mathematics. In Fortran it is irrelevant whether this is a one-dimensional or multi-dimensional array. In chapter ??, where we deal with allocation of matrices, we will introduce the numerical libraries Armadillo and Blitz++ which allow for similar treatments of arrays in C++. By default however, these features are not included in the ANSI C++ standard.

2.7 Exercises

Set up an algorithm which converts a floating number given in the decimal representation to the binary representation. You may or may not use a scientific representation. Write thereafter a program which implements this algorithm.

Make a program which sums

1.

$$s_{\text{up}} = \sum_{n=1}^N \frac{1}{n},$$

and

$$s_{\text{down}} = \sum_{n=N}^{n=1} \frac{1}{n}.$$

The program should read N from screen and write the final output to screen.

2. Compare s_{up} og s_{down} for different N using both single and double precision for N up to $N = 10^{10}$. Which of the above formula is the most reliable one? Try to give an explanation of possible differences. One possibility for guiding the eye is for example to make a log-log plot of the relative difference as a function of N in steps of 10^n with $n = 1, 2, \dots, 10$. This means you need to compute $\log_{10}(|(s_{\text{up}}(N) - s_{\text{down}}(N))/s_{\text{down}}(N)|)$ as function of $\log_{10}(N)$.

Write a program which computes

$$f(x) = x - \sin x,$$

for a wide range of values of x . Make a careful analysis of this function for values of x near zero. For $x \approx 0$ you may consider to write out the series expansions of $\sin x$

$$\sin x = x - \frac{x^3}{3!} + \frac{x^5}{5!} - \frac{x^7}{7!} + \dots$$

Use the loss of precision theorem of Eq. (2.2) to show that the loss of bits can be limited to at most one bit by restricting x so that

$$1 - \frac{\sin x}{x} \geq \frac{1}{2}.$$

One finds then that x must at least be 1.9, implying that for $|x| < 1.9$ we need to carefully consider the series expansion. For $|x| \geq 1.9$ we can use directly the expression $x - \sin x$.

For $|x| < 1.9$ you should device a recurrence relation for the terms in the series expansion in order to avoid having to compute very large factorials.

Assume that you do not have access to the intrinsic function for $\exp x$. Write your own algorithm for $\exp(-x)$ for all possible values of x , with special care on how to avoid the loss of precision problems discussed in the text. Write thereafter a program which implements this algorithm.

The classical quadratic equation $ax^2 + bx + c =$ with solution

$$x = \left(-b \pm \sqrt{b^2 - 4ac} \right) / 2a,$$

needs particular attention when $4ac$ is small relative to b^2 . Find an algorithm which yields stable results for all possible values of a , b and c . Write thereafter a program and test the results of your computations.

Write a Fortran program which reads a real number x and computes the precision in bits (using the function `DIGIT(x)`) for single and double precision, the smallest positive number (using `TINY(x)`), the largest positive number (using the function `HUGE(x)`) and the number of leading digits (using the function `PRECISION(x)`). Try thereafter to find similar functionalities in C++ and Python.

Write an algorithm and program which reads in a real number x and finds the two nearest machine numbers x_- and x_+ , the corresponding relative errors and absolute errors.

Recurrence relations are extremely useful in representing functions, and form expedient ways of representing important classes of functions used in the Sciences. We will see two such examples in the discussion below. One example of recurrence relations appears in studies of Fourier series, which enter studies of wave mechanics, be it either in classical systems or quantum mechanical ones. We may need to calculate in an efficient way sums like

$$F(x) = \sum_{n=0}^N a_n \cos(nx), \quad (2.3)$$

where the coefficients a_n are known numbers and x is the argument of the function $F()$. If we want to solve this problem right on, we could write a simple repetitive loop that multiplies each of the cosines with its respective coefficient a_n like

```
for ( n=0; n < N; n++) {
    f += an*cos(n*x)
}
```

Even though this seems rather straightforward, it may actually yield a waste of computer time if N is large. The interesting point here is that through the three-term recurrence relation

$$\cos(n-1)x - 2\cos(x)\cos(nx) + \cos(n+1)x = 0, \quad (2.4)$$

we can express the entire finite Fourier series in terms of $\cos(x)$ and two constants. The essential device is to define a new sequence of coefficients b_n recursively by

$$b_n = (2\cos(x))b_{n-1} - b_{n+2} + a_n \quad n = 0, \dots, N-1, N, \quad (2.5)$$

defining $b_{N+1} = b_{N+2} = \dots = 0$ for all $n > N$, the upper limit. We can then determine all the b_n coefficients from a_n and one evaluation of $2\cos(x)$. If we replace a_n with b_n

in the sum for $F(x)$ in Eq. (2.3) we obtain

$$\begin{aligned} F(x) = & b_N [\cos(Nx) - 2\cos((N-1)x)\cos(x) + \cos((N-2)x)] + \\ & b_{N-1} [\cos((N-1)x) - 2\cos((N-2)x)\cos(x) + \cos((N-3)x)] + \dots \\ & b_2 [\cos(2x) - 2\cos^2(x) + 1] + b_1 [\cos(x) - 2\cos(x)] + b_0. \end{aligned} \quad (2.6)$$

Using Eq. (2.4) we obtain the final result

$$F(x) = b_0 - b_1 \cos(x), \quad (2.7)$$

and b_0 and b_1 are determined from Eq. (2.3). The latter relation is after Chensaw. This method of evaluating finite series of orthogonal functions that are connected by a linear recurrence is a technique generally available for all standard special functions in mathematical physics, like Legendre polynomials, Bessel functions etc. They all involve two or three terms in the recurrence relations. The general relation can then be written as

$$F_{n+1}(x) = \alpha_n(x)F_n(x) + \beta_n(x)F_{n-1}(x).$$

Evaluate the function $F(x) = \sum_{n=0}^N a_n \cos(nx)$ in two ways: first by computing the series of Eq. (2.3) and then using the equation given in Eq. (2.5). Assume that $a_n = (n+2)/(n+1)$, set e.g., $N = 1000$ and try with different x -values as input.

Often, especially when one encounters singular behaviors, one may need to rewrite the function to be evaluated in terms of a Taylor expansion. Another possibility is to use so-called continued fractions, which may be viewed as generalizations of a Taylor expansion. When dealing with continued fractions, one possible approach is that of successive substitutions. Let us illustrate this by a simple example, namely the solution of a second order equation $x^2 - 4x - 1 = 0$, which we rewrite as $x = \frac{1}{4+x}$, which in turn could be represented through an iterative substitution process

$$x_{n+1} = \frac{1}{4 + x_n},$$

with $x_0 = 0$. This means that we have

$$x_1 = \frac{1}{4},$$

$$x_2 = \frac{1}{4 + \frac{1}{4}},$$

$$x_3 = \frac{1}{4 + \frac{1}{4 + \frac{1}{4}}},$$

and so forth. This is often rewritten in a compact way as

$$x_n = x_0 + \frac{a_1}{x_1 + \frac{a_2}{x_2 + \frac{a_3}{x_3 + \frac{a_4}{\dots}}}},$$

or as

$$x_n = x_0 + \frac{a_1}{x_1 +} \frac{a_2}{x_2 +} \frac{a_3}{x_3 +} \dots$$

Write a program which implements this continued fraction algorithm and solve iteratively Eq. (2.7). The exact solution is $x = 0.23607$ while already after three iterations you should obtain $x_3 = 0.236111$.

Many physics problems have spherical harmonics as solutions, such as the angular part of the Schrödinger equation for the hydrogen atom or the angular part of the three-dimensional wave equation or Poisson's equation.

The spherical harmonics for a given orbital momentum L , its projection M for $-L \leq M \leq L$ and angles $\theta \in [0, \pi]$ and $\phi \in [0, 2\pi]$ are given by

$$Y_L^M(\theta, \phi) = \sqrt{\frac{(2L+1)(L-M)!}{4\pi(L+M)!}} P_L^M(\cos(\theta)) \exp(iM\phi),$$

The functions $P_L^M(\cos(\theta))$ are the so-called associated Legendre functions. They are normally determined via the usage of recurrence relations. Recurrence relations are unfortunately often unstable, but the following relation is stable (with $x = \cos(\theta)$)

$$(L-M)P_L^M(x) = x(2L-1)P_{L-1}^M(x) - (L+M-1)P_{L-2}^M(x),$$

and with the analytic (on closed form) expressions

$$P_M^M(x) = (-1)^M (2M-1)!! (1-x^2)^{M/2},$$

and

$$P_{M+1}^M(x) = x(2M+1)P_M^M(x),$$

we have the starting values and the equations necessary for generating the associated Legendre functions for a general value of L .

1. Make first a function which computes the associated Legendre functions for different values of L and M . Compare with the closed-form results listed in chapter ??.
2. Make thereafter a program which calculates the real part of the spherical harmonics

3. Make plots for various $L = M$ as functions of θ (set $\phi = 0$) and study the behavior as L is increased. Try to explain why the functions become more and more narrow as L increases. In order to make these plots you can use for example gnuplot, as discussed in appendix ??.
4. Study also the behavior of the spherical harmonics when θ is close to 0 and when it approaches 180 degrees. Try to extract a simple explanation for what you see.

Other well-known polynomials are the Laguerre and the Hermite polynomials, both being solutions to famous differential equations. The Laguerre polynomials arise from the solution of the differential equation

$$\left(\frac{d^2}{dx^2} - \frac{d}{dx} + \frac{\lambda}{x} - \frac{l(l+1)}{x^2} \right) \mathcal{L}(x) = 0,$$

where l is an integer $l \geq 0$ and λ a constant. This equation arises for example from the solution of the radial Schrödinger equation with a centrally symmetric potential such as the Coulomb potential. The first polynomials are

$$\mathcal{L}_0(x) = 1,$$

$$\mathcal{L}_1(x) = 1 - x,$$

$$\mathcal{L}_2(x) = 2 - 4x + x^2,$$

$$\mathcal{L}_3(x) = 6 - 18x + 9x^2 - x^3,$$

and

$$\mathcal{L}_4(x) = x^4 - 16x^3 + 72x^2 - 96x + 24.$$

They fulfil the orthogonality relation

$$\int_{-\infty}^{\infty} e^{-x} \mathcal{L}_n(x)^2 dx = 1,$$

and the recursion relation

$$(n+1)\mathcal{L}_{n+1}(x) = (2n+1-x)\mathcal{L}_n(x) - n\mathcal{L}_{n-1}(x).$$

Similalry, the Hermite polynomials are solutions of the differential equation

$$\frac{d^2 H(x)}{dx^2} - 2x \frac{dH(x)}{dx} + (\lambda - 1)H(x) = 0,$$

which arises for example by solving Schrödinger's equation for a particle confined to move in a harmonic oscillator potential. The first few polynomials are

$$H_0(x) = 1,$$

$$H_1(x) = 2x,$$

$$H_2(x) = 4x^2 - 2,$$

$$H_3(x) = 8x^3 - 12x,$$

and

$$H_4(x) = 16x^4 - 48x^2 + 12.$$

They fulfil the orthogonality relation

$$\int_{-\infty}^{\infty} e^{-x^2} H_n(x)^2 dx = 2^n n! \sqrt{\pi},$$

and the recursion relation

$$H_{n+1}(x) = 2xH_n(x) - 2nH_{n-1}(x).$$

Write a program which computes the above Laguerre and Hermite polynomials for different values of n using the pertinent recursion relations. Check your results against some selected closed-form expressions.

Bibliography

- [1] D. Kincaid and W. Cheney. *Numerical Analysis*. Brooks/Gole Publishing Company, 1996.