

Introduction to modern Fortran (and Makefiles)

1 Introduction to programming

1.1 What is a programming language?

Computers speak binary. Each chipset (e.g. Intel x86 or the ARM A5 processor in your smartphone) has it's own set of low-level instructions known as "assembly language" that translates to the raw binary instructions. Assembly language is very low level: "push onto stack, move to this memory address, pop number off the stack, etc.".

A programming language is a human-readable language that you can use to specify your human intent (i.e. "solve this equation", "add these numbers and print the answer"). The compiler translates this into raw assembler code and ultimately into an executable binary file. Thus all programming languages roughly do the same thing - the difference is just how easy it is to specify what you want to do in a given language.

In Unix-speak a program is just a "text file" that you create with any standard text editor such as vi, emacs or gedit. The compiler converts your text into a binary file that it can execute. You then run the program. Scripting languages such as Python and Perl differ from more traditional compiled languages because these two steps are combined into one, that is the program is compiled 'on-the-fly' when you run it.

1.2 Hello world

With any language, you should always start with the simplest possible program, or:

"Make it work before you make it complicated"

To type and run a basic program in Fortran, create a text file with the extension .f90. For example type gedit hello.f90 & and enter contents as follows:

program hello
 print*,'hello world'
end program hello

We will use the *Gnu Fortran compiler*, gfortran (since it is free), so the relevant command is as follows (having made sure you've saved the file to disk):

\$ gfortran -o bob hello.f90

Assuming this did not throw up any errors, this creates an executable file with the name bob, that you can run from the unix shell:

\$./bob hello world

It is instructive to see what the assembler code actually looks like. You can do this by typing:

```
gfortran -S hello.f90
```

This creates a text file called hello.s containing the assembler code. The contents of this file shows how unreadable assembler language really is and hence why programming languages exist:

```
$ more hello.s
         .cstring
LCO:
         .ascii "hello.f90\0"
         .const
LC1:
         .ascii " hello world"
         .text
_MAIN__:
LFB0:
                 %rbp
        pushq
LCFIO:
        movq
                 %rsp, %rbp
LCFI1:
                 $480, %rsp
        subq
                 LCO(%rip), %rax
        leaq
                 %rax, -472(%rbp)
        movq
                 $3, -464(%rbp)
        movl
                 $128, -480(%rbp)
        movl
                 $6, -476(%rbp)
        movl
                 -480(\%rbp), \%rax
        leaq
```

2 Introduction to modern Fortran

Fortran (For-mula Trans-lation is a language written to solve mathematical equations. The modern language (Fortran), defined via standards set in 1990, 2003 and 2008, is

backwards-compatible with the old language (referred to as FORTRAN, standardised in 1977) meaning that it can be used to run old codes as well, but certain features have been declared obsolete. Most criticisms of Fortran refer to problems in FORTRAN that were fixed more than 25 years ago.

2.1 Why use modern Fortran instead of F77?

Here is why: Type the following into a file called test.f. Note that in the old FOR-TRAN language each instruction needed to start in the 6th column and cannot exceed 72 characters in length — this was to fit on the punchcards they used in the 1960s:

```
program badfort

do 30 i=1.20
    print*,i
30 continue

end
```

Compile and run this:

```
$ gfortran -o test test.f
```

First, notice that the compiler compiled this with no warnings or errors. But what is the output? Does the program do what you expect?

The above example has an obvious bug in it. But the compiler was not able to detect the bug because the program is written sloppily. Most of your time programming will be spent debugging. So anything in a language that can help you find problems more easily will save you a lot of time and pain. The big one in Fortran is the *implicit none* statement. Type this program in modern syntax, but with the same bug, into a file called test.f90 (no special line spacing is required if the file extension is .f90 instead of .f):

```
program badfort
  implicit none

do i=1.20
    print*,i
  enddo

end program badfort
```

Compiling this, the compiler will immediately tell you what is wrong with it:

This means you can fix the problems *before* you go launching your satellite. The line and column numbers are given in the error messages so you can pinpoint the problematic lines of code.

A longer explanation here — F77 allowed any variable to be assumed to be real or integer depending on the first letter. Specifically variables starting with letters a-h or o-z are assumed to be real (floating point) while variables starting with l-n are assumed integers. Thus in our example above the compiler assumed we had set a new variable called "do 30 i" and set it's value to 1.2. We then print i, which is a different variable that has not been set and so assumes a random value from memory, which is what was printed. Thus the famous joke:

"God is real unless declared integer"

Hence, *always* declare implicit none at the top of your code. With gfortran you can also enforce this even if you forgot to type it by adding the flag -fimplicit-none, i.e.:

```
$ gfortran -fimplicit-none -o test test.f90
```

2.2 A simple program

Let's perform a simple mathematical calculation in Fortran (e.g. in a file called maths.f90):

```
program maths
  implicit none
```

```
real :: r, area
real, parameter :: pi = 4.*atan(1.)

r = 2.0
area = pi*r**2
print*,' pi is ',pi
print*,' the area of a circle of radius ',r,' is ',area
end program maths
```

Compile and run this and confirm the output is as follows:

```
$ gfortran -o maths maths.f90
$ ./maths
pi is    3.14159274
the area of a circle of radius    2.00000000 is    12.5663710
```

2.3 Organising your code

Before we go into detail about how to do specific things, we must learn how to organise a code. The guiding principle is the following:

Never repeat code.

You will be tempted. What's a little cut-and-paste between friends? Cut-and-paste equals death to good programming. Instead, you need to organise your code so that it is broken up into small, re-usable pieces.

Let's take the example above, and put the area calculation into a function. The modern way to do this is to put it in a module. To do this, create a new file called area.f90 where the area function is defined. Let's also add some *comments* to the code:

```
! A module containing functions to compute the area of a circle
! Written by Daniel Price, 2015
module geometry
implicit none
real, parameter :: pi = 4.*atan(1.)
public :: area, pi
private

contains
!
```

```
! A function to calculate the area of a circle of given radius
!
real function area(r)
real, intent(in) :: r
area = pi*r**2
end function area
end module geometry
```

The main purpose of comments is for your future self. It is very easy to forget the thought process you had when you wrote the program, so use comments to record this information. At bare minimum, state the purpose of the module, subroutine, function or program in the comments. Making this a habit will save you pain later.

Let's re-write the main program (in maths.f90) to use the module:

```
program maths
  use geometry, only:area,pi
  implicit none
  real :: r

r = 2.0
  print*,' pi is ',pi
  print*,' the area of a circle of radius ',r,' is ',area(r)
end program maths
```

To compile this we need to tell the compiler about the different files and also compile them in the order that they are needed

```
$ gfortran -o maths area.f90 maths.f90
$ ./maths
```

Does your program give the same output as before? By putting functionality into modules in this way we can construct a software project as large as we like, and maximise the reuse of code. It also enables us to debug and test the functions in the module separately from the main code (in software parlance this is known as *unit testing*) which is a huge help in keeping the program bug-free.

You can see some other important things in the programs we just typed:

• We declared implicit none in both the main program and at the top of the module (it is unnecessary in the function, the global declaration at the top of the module already ensures this is the case)

- In the module we explicitly denoted which parts are public (that is, available for use in other parts of the program) and which are private (available only to functions inside the module).
- In the function we declared the radius variable as intent(in) which tells the compiler that this variable is expected to have a value on input and should not be changed by the function.
- In the program, we explicitly stated which bits of the module we wanted using only:area,pi rather than importing everything in the module.

Use these for same reason as the "do 30 i" example above — the more information you can give the compiler about what you are trying to do, the easier it is for the compiler to report an error and the less time you will waste debugging your code.

2.4 Compiler flags

Another time saver is to turn on *compiler warnings*. The syntax is different with different compilers, but with gfortran this is achieved by adding the -Wall flag:

\$ gfortran -Wall -o maths area.f90 maths.f90

To use the maximum possible level of warnings and the strictest adherence to the latest standards, we would use:

\$ gfortran -Wall -Wextra -pedantic -std=f2008 -o maths area.f90 maths.f90

Finally, when writing simulation code we want it to run fast. Achieve this by turning on optimisation flags:

\$ gfortran -03 -o maths area.f90 maths.f90

Note that the flag is the letter O followed by a 3 (for optimisation level 3), not zero.

3 Using Makefiles to compile your code

We only used two files in the example above, but it quickly becomes laborious to remember the various files that go together to make a program and the combination of compiler flags you last used. Also with larger codes, is is inefficient to recompile the whole code even though only one file has changed. To do this we must compile each file separately into an *object* file (.o) and then *link* the object files together to create the binary executable. We do this using the -c flag as follows:

```
$ gfortran -o area.o -c area.f90
$ gfortran -o maths.o -c maths.f90
$ gfortran -o maths maths.o area.o
```

Doing this by hand is laborious, but we can automate it using a Makefile. The make program is actually a programming language in itself — one designed to make compiling codes easy.

3.1 Hello world in Gnu Make

To use make, simply create a file in the working directory with the name Makefile (that is, make a makefile), with contents as follows:

VAR=hello

```
default:
<tab> echo ${VAR}
```

Here we have just set a string variable called VAR and set its value to 'hello'. <tab> means press the tab key. Tab is treated very different to spaces in Makefiles, so this is important. You should already be familiar with the unix echo command and what it does. Typing make on the command line will run this:

```
$ make
echo 'hello'
hello
```

The key elements of Makefiles are targets and dependencies.

3.2 Targets

To understand targets edit your Makefile to have two different targets, 'merry' and 'fun':

```
VAR=hello
VAR2=world

merry:
<tab> @echo ${VAR}

fun:
<tab> @echo ${VAR2}
```

(The @ stops the Makefile from printing the shell command to the screen before it executes it). You can now execute each of these targets separately:

```
$ make
hello
$ make merry
hello
$ make fun
world
```

Hopefully this makes targets clear — by convention the first target in the file is also the default (i.e. the one executed when you just type make).

3.3 Dependencies

Dependencies are specified on the right hand side of the colon. Let's modify the above example so that the 'fun' target depends on the 'merry' target by amending the appropriate line to read:

```
fun: merry
<tab> @echo ${VAR2}
```

Now type 'make fun' and 'make merry' as previously. Do you understand the output? In particular the 'merry' target should give:

```
$ make fun
hello
world
```

3.4 Variables in Makefiles

Variables are fairly straightforward in the examples above. You can override variable setting(s) on the command line:

```
$ make VAR=goodbye fun
goodbye
world
```

There are also some special predefined variables — \$@ refers to the name of the target (the text to the left of the colon), while \$< refers to the dependencies (everything to the right of the colon). Try the following example:

```
fun: merry
<tab>    @echo ${VAR2} is $@ and $<

What is the output?

We can also define new variables from existing variables, with various string replacement options, e.g.

VAR=hello
VAR2=world
VAR3=${VAR2:world=planet}}

hay: merry
<tab>    @echo ${VAR3}

and you should find:

$ make hay
hello
planet
```

3.5 Using a Makefile to compile your code

From the above you should be able to see how to use a Makefile to compile your code. Since this is what make is designed for there are standard names for the appropriate variables to use. A very simple Makefile for compiling our Fortran program would be:

```
FC=gfortran
FFLAGS=-03 -Wall -Wextra
SRC=area.f90 maths.f90
OBJ=${SRC:.f90=.o}
%.o: %.f90
<tab> $(FC) $(FFLAGS) -o $@ -c $<
maths: $(OBJ)
<tab> $(FC) $(FFLAGS) -o $@ $(OBJ)

clean:
<tab> rm *.o *.mod maths
```

A couple of things:

- The first target is a special 'wildcard' rule telling Make how to compile *any* file ending in .o from the corresponding .f90 file. This specifies that a file called fred.o would depend on the corresponding fred.f90 and if the .o does not already exist should be created using the rule specified on the line below.
- We added a clean target which cleans up the 'dog poo' generated by the intermediate step of creating the .o files. So we can just type make clean to clean up these temporary files.
- Notice the use of the special variables \$0 and \$<.

Now compiling your code should be as simple as typing 'make':

```
$ make
gfortran -03 -Wall -Wextra -o area.o -c area.f90
gfortran -03 -Wall -Wextra -o maths.o -c maths.f90
gfortran -03 -Wall -Wextra -o maths area.o maths.o
$ ./maths
```

Even better, if nothing has changed then typing make again does nothing:

```
$ make
make: 'maths' is up to date.
```

but if we make a change to just one file then make knows to only recompile that one file and redo the link step:

```
$ touch maths.f90
$ make
gfortran -03 -Wall -Wextra -o maths.o -c maths.f90
gfortran -03 -Wall -Wextra -o maths area.o maths.o
```

You should see, however, that this is dangerous if we apply it to the area.f90 module. That is, if area.f90 changes then we should also recompile maths.f90 as well, since our program calls functions in that module. How should we ensure this happens? By specifying the dependency in our Makefile!

```
maths.o: geometry.mod
```

Specifying all of the dependencies is hard to do for bigger codes, and only relevant if the interface to the routine changes, I often don't bother and instead just make sure the files are complied in the correct order (i.e. SRC=area.f90 maths.f90 not SRC=maths.f90 area.f90). Then I just type make clean if I know that I have changed the way a routine is called.

4 Key Fortran concepts

4.1 Floating point precision in Fortran

The basic types of variables in Fortran are integer, logical, real and character. Declaration of these takes an optional kind parameter. So for example to specify an 8-byte real instead of a 4-byte real one would use:

```
real(kind=8) :: x
or just
real(8) :: x
```

Technically, using the kind parameter to specify the number of bytes is compiler dependent, but it is very widespread. The correct way is to let Fortran define the kind according to the desired precision of the calculation using selected_real_kind. To do this, let's create another file called precision.f90 with contents as follows:

```
! module to define precision of real variables
module prec
 implicit none
 integer, parameter :: dp = selected_real_kind(P=10,R=30)
 integer, parameter :: sp = selected_real_kind(P=5,R=15)
 integer, parameter :: dp_alt = kind(0.d0)
public :: dp,sp,print_kind_info
private
contains
 ! routine to print information about the kinds being used
 subroutine print_kind_info()
 real(sp) :: pi_single
 real(dp) :: pi_double
 print*,' double precision is kind=',dp
 print*,' single precision is kind=',sp
 print*,' kind of a double precision number is ',dp_alt
 pi_single = 4.0_sp*atan(1.0_sp)
 pi_double = 4.0_dp*atan(1.0_dp)
 print*,' pi in single precision is ',pi_single
```

```
print*,' pi in double precision is ',pi_double

! see what happens if we accidentally mix precisions
pi_double = pi_single
print*,' pi converted from single is = ',pi_double

end subroutine print_kind_info
end module prec
```

Then add this file to the line in the Makefile:

```
SRC=precision.f90 area.f90 maths.f90
```

and edit the main program to call this subroutine and also print the kind of a default real. That is:

```
program maths
  use geometry, only:area,pi
  use prec,     only:print_kind_info
  implicit none
  real :: r

call print_kind_info()
  print*,' radius is of kind ',kind(r)
  ...
end program maths
```

Compiling and running this, you should find:

\$./maths

```
double precision is kind= 8
single precision is kind= 4
pi in single precision is 3.14159274
pi in double precision is 3.1415926535897931
pi converted from single is = 3.1415927410125732
radius is of kind 4
```

Notice the precision loss in converting from double to single precision.

Exercise 4.1.1: Rewrite the area function so that it works in double precision instead of single precision. To do this, redefine all the real variables to be real(dp), with dp imported from our prec module as above.

4.2 Arrays and array operations

4.2.1 One dimensional arrays and array operations

Arrays are very powerful in Fortran, and this is one of the key advantages over other languages such as C and C++. Defining an array just means giving one or more *dimensions* to a variable. For example a vector of 3 numbers can be defined using:

```
real :: x(3)
```

Let's create an example module for array operations, in a file called arrays.f90:

```
module arrays
 implicit none
contains
 subroutine array_examples()
  use prec, only:sp
  real(sp) :: x(3),y(3)
  ! set each element individually
  x(1) = 1.0
  x(2) = 2.0
  x(3) = 3.0
  print*, 'x = ',x
  ! set whole array equal to 1
  x = 1.
  print*,' x = ',x
  ! set array parts in one line
  x = (/1.0, 2.0, 3.0/)
  print*, 'x = ',x
  ! set y=x
  y = x
  print*,' y = ',y
  ! array operations
  print*,'sum=',sum(y)
  print*,'maxval=',maxval(y)
  print*,'minval=',minval(y)
  print*,'maxloc=',maxloc(y)
```

```
print*,'minloc=',minloc(y)

! magnitude
print*,' |x| = ',sqrt(dot_product(x,x))
print*,' |x| = ',norm2(x)

end subroutine array_examples

end module arrays

As previously, use your module from the main code:

program maths
use arrays, only:array_examples
implicit none

call array_examples()

end program maths
```

4.2.2 Multi dimensional arrays

Multi-dimensional arrays are similar. For a 3×3 matrix just define:

```
real :: x(3,3)
```

Again, you can add, subtract and multiply arrays just as if they were scalars.

5 Logic and loops: if/then/else and select case

Here is a simple example of how to construct and use logical statements:

```
program ifanimal
  implicit none
  logical :: isacow,isadog,hastwohorns
  integer, parameter :: nhorns = 2

  isacow = .true.
  isadog = .false.
  if (isacow) then ! check if our animal is a cow
      write(*,''(a)'',advance='no') my animal is a cow...'
```

```
if (nhorns==2) write(*,*) ' ...with two horns'
elseif (isadog) then
    print*,' my animal is a dog. Woof.'
else
    print ''(a)'','my animal is not a cow'
    hastwohorns = (nhorns==2)
    if (hastwohorns) print ''(a)'',' but it has two horns'
endif
end program ifanimal
```

6 Writing to and reading from files

Writing to/reading from files is just a matter of using the open,close,write and read statements.

```
program io
 implicit none
 character(len=20) :: filename
 real :: x, y
 filename='results.out'
 x = 1.
 y = 2.
 ! write to ascii file
 open(unit=1,file=filename,status='replace',form='formatted')
 write(1,*) x, y
 close(1)
 ! reset vars
 x = 0.
 y = 0.
 ! read from ascii file
 open(unit=2,file=filename,status='old')
 read(2,*) x,y
 close(2)
 ! print vars
 print*,' x = ',x,' y = ',y
end program io
```

7 Exercises

The actual mechanics of programming in Fortran is just a matter of learning the syntax. The best way is to try to implement practical examples. Here are a few simple exercises.

1. Implement a subroutine to return the n real solutions to a quadratic equation, i.e.

$$ax^2 + bx + c = 0, (1)$$

using the exact solution, i.e.

$$x = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a}.\tag{2}$$

Check that the solution returned by your routine is correct by evaluating (1) in your program.

2. Write a subroutine that sets up two one dimensional arrays for position (x) and velocity (v_x) . Set the position in equally spaced increments between 0 and 1, and velocity array equal to:

$$v_x = \sin(2\pi x) \tag{3}$$

Write the position and velocity arrays to a file and use gnuplot or similar to plot the arrays.

3. Write a simple program that asks for your name and birth year and prints back your name and age. Google for the date_and_time intrinsic routine.

To familiarise yourself further with the basic syntax of Fortran, take yourself through the 'Fortran primer'.

8 Advanced Fortran

While not designed to be comprehensive, in the remainder of this session we will cover some very useful "advanced" aspects of Fortran that are hard to discover without being told about them. Try to use git as you go along to save versions of the code that you write.

8.1 Making functions operate on whole arrays

In the previous session we wrote an 'area' function that returns the area for a circle. But the power of modern Fortran is in array operations. For example try performing the following operation in a simple program:

```
program array
implicit none
real :: x(10)

do i=1,size(x)
    x(i) = real(i)
enddo

print*,' sqrt(x) = ',sqrt(x)

end program array
```

The sqrt operation can be performed on the whole array at once, and return an array as the output. It is easy to make your own functions do this. The key is that they must have the elemental property. Elemental functions have certain restrictions. For example they must not print anything to the screen. From the above we see that it would be weird for the sqrt function to print anything. A related keyword is pure. Pure subroutines act entirely on their input, do not change global (shared) variables and do not perform any input or output. These kinds of functions can be easily parallelised.

Exercise

1. Modify your area function from the earlier lab session so that it can operate on a whole array. To do this, just add the elemental keyword to the function definition:

```
module blah
  implicit none

contains

real, elemental function area(r)
  real, intent(in) :: r
```

2. Fill an array with a bunch of different radii, and return the area calculated for each element of the array by calling the area function with an array as the input argument.

8.2 Generic interfaces

A second useful 'modern Fortran' concept is the use of generic interfaces. The idea is that you can use the same function name to refer to different routines with different kinds of arguments. For example, the intrinsic function sqrt() can take either a real or a double precision variable as an argument, but in both cases the same function name (sqrt) is used (in old FORTRAN these had separate names: sqrt and dsqrt). You will see an example of this in the prompting.f90 module in the examples folder.

To make your own 'generic interface', you must define the two different functions in your module with *different names* and then use an **interface** statement to give the generic name. For example:

```
module geometry
  implicit none
  interface area
    module procedure area_real, area_double
  end interface area
  public :: area
  private :: area_real, area_double
contains
 real, elemental function area_real(r)
  real, intent(in) :: r
  area_real = pi*r**2
 end function area_real
 real(kind=8), elemental function area_double(r)
  real(kind=8), intent(in) :: r
 end function area_double
end module geometry
```

In the above we made the generic name public but the specific names private, so that an external routine can *only* refer to the routine by its generic name.

Exercise

- 1. Modify your module so that it has both real and double precision version of the area function, and write a program to test this, by calling area with a real and double precision argument (for example area(1.0) and area(1.d0)). Make sure that you use enough precision in π for both cases!
- 2. Add a subroutine to your module that computes the area of a rectangle. The interface to this routine should look something like:

```
real function area_rectangle(width,height)
  real, intent(in) :: width, height
  ...
end function area_rectangle
```

3. Add this routine to the generic interface definition, and show that calling area with *two* arguments e.g. area(2.0,3.0) returns the area of a rectangle instead of a circle.

8.3 Optional arguments

Subroutines that are defined properly in modules can also have *optional* arguments. Whether or not these have been passed can be checked by the present() function. For example, let's add an optional argument factor, to the area_rectangle routine:

```
real function area_rectangle(width,height,factor)
real, intent(in) :: width, height
real, intent(in), optional :: factor

area_rectangle = width*height
if (present(factor)) then
    area_rectangle = area_rectangle*factor
endif

end function area_rectangle
```

Exercise

1. With the optional argument implemented as above, try calling the area function from the main program with the following argument lists:

```
area(1.0,2.0)
area(1.0,2.0,10.)
area(1.0,2.0,factor=1.)
area(1.0,2.0,factor=10.)
```

8.4 Calling C from Fortran

A key part of Fortran 2003 is the ability to interoperate with the C language. This is important because a lot of software libraries are written in C and it is useful to be able to call them directly. Let's give an example of how to do this.

First, paste the following code into a file called fastsqrt.c and save it to your current working directory:

```
/*
  implementation of a fast inverse sqrt function
  as described in Chris Lomont, "Fast Inverse Square Root"
  www.math.purdue.edu/~clomont
 precise implementation here by Daniel Price
*/
float finvsqrt(float *y)
      float x = *y;
      float xhalf = 0.5f*x;
      int i = *(int*)&x;
                              // get bits for floating value
      i = 0x5f375a86- (i>>1); // gives initial guess y0
                              // converts bits back to float
      x = *(float*)&i;
      x = x*(1.5f-xhalf*x*x); // Newton step, repeating increases accuracy
      x = x*(1.5f-xhalf*x*x); // Newton step, repeating increases accuracy
      x = x*(1.5f-xhalf*x*x); // Newton step, repeating increases accuracy
      return x;
      }
```

Then, open a file called fastmath.f90 that we will use to create the interface to this routine. Let's then define the interface to the C routine in this module:

```
module fastmath
  use iso_c_binding, only:c_float, c_double
  implicit none

private
  public :: fast_invsqrt
```

```
interface fast_invsqrt
pure function finvsqrt(x) bind(C)
  import
  implicit none
  real(kind=c_float), intent(in) :: x
  real(kind=c_float) :: finvsqrt
  end function finvsqrt
end interface fast_invsqrt
```

end module fastmath

Notice that we used an *intrinsic* module called <code>iso_c_binding</code> to be able to define a real variable that is the same as a <code>float</code> or <code>double</code> in the C language. The <code>bind(C)</code> attribute tells the compiler that this is actually a function call to C. There is no <code>contains</code> statement in the module above, since it does not contain any subroutines. We can then call the function with the 'Fortran' name we have given (<code>fast_invsqrt</code>) rather than it's name in C. So in your program add the following lines:

```
program testmath
 use fastmath, only:fast_invsqrt
 implicit none
 integer, parameter :: nmax = 10000000
 real :: x(nmax),y(nmax),t1,t2,t3
 integer :: i
 do i=1,nmax
     x(i) = real(i)
 enddo
 call cpu_time(t1)
 do i=1,nmax
    y(i) = 1./sqrt(x(i))
 enddo
 call cpu_time(t2)
 print*,'completed in ',t2-t1,' s'
 do i=1,nmax
    y(i) = fast_invsqrt(x(i))
 enddo
 call cpu_time(t3)
 print*,' fast version completed in ',t3-t2,'s'
end program testmath
```

To compile the code, we just have to compile the C routine using the C compiler, and the rest with gfortran:

```
$ gcc -o fastsqrt.o -c fastsqrt.c
$ gfortran -o fastmath.o -c astmath.f90
$ gfortran -o test.o -c test.f90
$ gfortran -o test test.o fastmath.o fastsqrt.o
$ ./test
```

Exercise

- 1. Which computation of the fast inverse square root is faster?
- 2. Add the appropriate interface definition for the double precision version of the fast sqrt routine (dinvsqrt) in fastsqrt.c as follows:

Include this in the definition of the generic interface fast_invsqrt and show that you can successfully call fast_invsqrt with either a real or double precision argument.

8.5 OpenMP parallelisation

A final aspect of Fortran and C is that both languages support parallelisation across multiple CPUs using OpenMP directives. OpenMP is a way to run a code in parallel in the case where all cpus share the same memory (i.e. are located on the same computer). Parallelisation across different machines connected across a network (known as 'distributed memory parallelisation') is significantly more complicated and is usually achieved using the Message Passing Interface (MPI). MPI (confusingly including the openMPI implementation) and openMP are *very* different things.

A simple openMP parallelisation of a loop would proceed as follows (in a file called par.f90):

```
program par
  implicit none
  integer :: i

!$omp parallel do default(none) private(i)
  do i=1,10
      print*,i
  enddo
  !$omp end parallel do

end program par
```

If the code is not compiled in parallel then the !\$omp is just treated as a comment. To compile in parallel, use the -fopenmp flag (the exact flag is specific to the compiler):

```
$ gfortran -fopenmp -o testpar par.f90
```

Exercise

- 1. Compare the output of the code compiled with and without the **-fopenmp** flag. Do you understand what is happening here?
- 2. Create a loop that evaluates a square root 1 million times. Run this in parallel and in serial and compare timings (using the cpu_time intrinsic function)

There is obviously a lot more to learn about writing parallel code, but this is merely to show you that such a possibility exists — you can proceed further by taking yourself through some of the openMP tutorials or documentation on the web.

8.6 Co-array Fortran

The Fortran 2008 standard defined a language standard for distributed memory parallelism within Fortran itself. This is known as 'co-array Fortran' and is now implemented in a few compilers though not yet widely used. Again, there are excellent tutorials on the web.