Scientific Fortran for Beginners

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Introduction

Welcome to Fortran!

If you've not downloaded gfortran or another compiler, please see https://fortran-lang.org/en/learn/os_setup/install_gfortran/

Fortran Source Files.

The standard modern Fortran file extension is .f90 . You might sometimes see .f95 or .f03 for specific Fortran standards.

If you happen to see .f or .for file extensions, you're dealing with old FORTRAN, which looks quite different.

Hello World!

program main

A Fortran file is made up of a list of statements, each taking place on its own line. There's no need for curly braces or indentation, but you'll probably want to use indentation.

```
implicit none
   print *, "Hello, World!"
end program main

Fortran is also case insensitive, so PROGRAM MAIN is equivalent
to program main or pRoGrAm MaIn
```

Compilation

Once you have written a Fortran file, it needs to be compiled. If you have installed gfortran this is done in the following way.

\$ gfortran myprogram.f90 -o myprogram.out You can then run the binary like so.

\$./myprogram.out

Variables

Fortran is a statically typed language, variables must be defined with a type and can only be assigned to values of that type. Here are a few examples:

```
integer :: i, j, ij, jsq
logical :: i_less_than_j

i = 12
j = 17
i_less_than_j = i < j
ij = i * j
jsq = j ** 2</pre>
```

Constants are defined using the keyword parameter .

Floating point numbers come in single and double precision forms. It's standard practice to define the **kinds** of these using the following syntax.

```
integer, parameter :: dp = kind(1.0d0), &
    & sp = kind(1.0e0)
real(sp) :: f1
real(dp) :: f2

f1 = 1.0_sp
f2 = 1.0_dp
```

Arrays

Arrays are assigned using parentheses () . To make an array of three dimensions you would:

```
real(dp) :: array_3d(3)

! Set all elements to zero.
    array_3d = 0._dp

! Set the first element to 1.
    array_3d(1) = 1._dp

! Set multiple elements.
    array_3d = (/1._dp, 0._dp, 0._dp/)
```

Flow Control

If statements use the following syntax.

```
if (some_logical) then
    ! Your first branch in here.
else if (another_logical) then
    ! Your second branch in here.
end if
```

Basic for loops, or do loops as they are known in Fortran use the following syntax:

```
do x = 1, n
     ! Your repeating code in here.
end do

While loops:
    do while (some_logical)
     ! Your repeating code in here.
end do
```

Exercise 1

Write a program which calculates the dot product of two vectors, and prints it. Use the following two vectors:

$$(1, 30, 1.2, -12) \cdot (100.05, 0.00034, 6, 67)$$

Subprograms

Fortran supports both functions (those which return a value) and subroutines (those which don't return a value). We'll start with function syntax as its what many of you may be most familiar with.

```
program main
    implicit none
    integer x
    x = 12
    print *, x
    x = add_two(x)
    print *, x
contains
    integer function add two(x)
        integer, intent(in) :: x
        add two = x + 2
    end function add two
end program main
```

And now subroutines.

```
program main
    implicit none
    integer x
    x = 12
    print *, x
    call add two(x)
    print *, x
contains
    subroutine add two(x)
        integer, intent(in out) :: x
        x = x + 2
    end subroutine add_two
end program main
```

A Small Project: Coulomb's Law

The second part of this workshop is about creating a little graph of two charged particles acting under Coulomb's law.

A quick refresher: the electrostatic force felt by a charged particle due to another charged particle is

$$\mathbf{F}_1 = \frac{q_1 q_2}{4\pi\varepsilon_0 |\mathbf{r}_{12}|^2} \hat{\mathbf{r}_{12}},$$

where ${f r}_{12}$ is a vector in the direction of particle 1 from particle 2.

Writing to a file

Fortran doesn't have any plotting tools that are as easy to use as matplotlib (in my opinion), for example, so its common practice to write out the data to a file and then to write scripts that analyse the data.

This is also good practice if you wish to save on time spent generating data. The syntax for opening a file, writing to it line by line, and closing it is as follows.

```
open(unit = 1, file = "myfile.data", status = "new")
write(1,*) "3.14"
close(1)
```

Exercise 2: Coulomb's Law

Back to the project, your code should:

- Create a file named coulomb_repulsion.data .
- Define two electrons, with positions and charges.
- Calculate the Coulomb force felt by the moving electron at 1,000 points in space as ${\bf r}_{12}$ is varied between (0.1,0.1,0.1) mm and (1.1,1.1,1.1) mm.
- Write each force on a new line in coulomb_repulsion.data

In your simulation please use the following constants:

$$\pi = 3.14, \;\; \varepsilon_0 = 8.85 \times 10^{-12} \mathrm{F/m}, \;\; e = 1.60 \times 10^{-19} \mathrm{C}$$

Here's the coulomb repulsion equation again:

$$\mathbf{F}_1 = \frac{q_1 q_2}{4\pi\varepsilon_0 {|\mathbf{r}_{12}|}^2} \hat{\mathbf{r}_{12}},$$