# Lecture 4: Functions and modular code

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## Aim

This lecture will introduce how to simplify your code by writing functions and how to reuse code easily in many different places.

#### 1 Functions

Occasionally, there is a particular section of your code that you would like to reuse over-and-over, without having to write the code over and over (as previously mentioned, programmers are lazy). For this, we make use of *functions*, the use of which will be familiar. For example, we have used the print() function already in previous weeks, and the NumPy library contains a wide variety of functions, some of which were introduced last week. However, this week we shall see how it is possible to write our own functions in Python.

The general syntax for defining a function in Python is as follows,

```
# Defining a function

def my_function(argument_1, argument_2):
    """

Adds together two arguments.
    """

result = argument_1 + argument_2
    return result
```

Above, we defined a function named my\_function which took two arguments, added them together to produce a result, which was returned. Once defined it is possible to use this function in our code as follows,

```
# Using our function
a = 1
b = 2
c = my_function(a, b)
print(c)
```

It can be seen when looking at the above example, that the object that is returned from the function is then assigned to the variable c. The text immediately following the function definition line is the *docstring*, this is important to help others understand the purpose of the function without having to read the code (basically it is a description of the function's action).

The above example of a function is relatively simple, just adding together two numbers. However, a function can contain a large amount of code abstracted to a single line. Furthermore, a well named function can increase the readibility of code significantly (consider the atomic distance code from previous weeks, the abstraction of the distance calculation to an appropriately named function would make this more understandable).

The my\_function example above contained two required arguments. These are the objects that are passed of the function when it is called (and typically these are operated on in some fashion) and are necessary for the function to run. In addition to these required arguments, other arugments may be included in a given function; such as default arguments and variable-length arguments. These are showing in the functions below,

```
# A function with defaults
def my_function(arg1, arg2, arg3=0):
   Adds together two arguments,
   and an optional third.
   result = arg1 + arg2 + arg3
   return result
print(my_function(1, 2))
print(my_function(1, 2, arg3=3))
# A variable length argument
def my_function(arg1, arg2, *arg3):
   Definitely adds together two arguments,
   but can optionally add together any
   number!
   result = arg1 + arg2
   for i in arg3:
       result += i
   return result
print(my_function(1, 2))
print(my_function(1, 2, 3, 4, 5))
```

Note that in default argument example, if no value was given it would default to 0.

You should be aware that a Python function does not need to have any

arguments. For example, you could write a function that returns the value of the Planck constant (ideally using the SciPy constants library),

```
# A function with no arguments

from scipy import constants

def h():
    """
    Planck constant.
    """
    return constants.h

print('Planck constant = {:.3e}'.format(h()))
```

## 2 Modules

In addition to writing your own functions, it is possible to *clean up* your code further by creating your own *modules*. These modules are similar to the libraries (like NumPy and SciPy) that we discussed last week, however, they are created by you. This is achieved by creating a file in the folder where you would like to use the module, and then *importing* the module in the code where your would like to use it. For example, say you have a file called atom\_helper.py in which you will put many functions related to atomistic calculations. Lets start it off with our distance calculation,

```
# atom_helper.py
import numpy as np

def distance(a, b):
    """
    Determines the distance between two points
    in any number of dimensions.
    """
    return np.sqrt(np.sum(np.square(a - b)))
```

Having created this module, we can then import it into a Jupyter Notebook that is in the same folder, and access the distances function as shown below,

```
# Using the atom_helper module
import numpy as np
import atom_helper
atom1 = np.array([1, 4, 2])
atom2 = np.array([5, 2, 6])
dist = atom_helper.distance(atom1, atom2)
```

#### print(dist)

It is clear from the above example, that the *user-defined* module can be imported in the same way as a library. Then any function defined in the module may be called from the Jupyter Notebook. This means that you can create modules containing a wide variety of functions and use them in many different Notebooks.

## 3 Problem

It is possible to estimate the interaction energy,  $E_{LJ}$  between two atoms using the Lennard-Jones expression,

$$E_{LJ} = \epsilon \left[ \left( \frac{r_m}{r} \right)^{12} - 2 \left( \frac{r_m}{r} \right)^6 \right], \tag{1}$$

where, r is the distances between the two atoms,  $\epsilon$  and  $r_m$  are constants describing the shape of the interaction potential surface. Add a function that can calculate the interaction energy, using the above relationship with constants of  $\epsilon = -2.2\,\mathrm{kJ\,mol^{-1}}$  and  $r_m = 3.7\,\mathrm{\mathring{A}}$ , to the atom\_helper.py module. Remember to plan the algorithm first!

Use the new function to calculate the interactions energy for each of the pairs of atoms used in Week 2.