

CH40208: TOPICS IN COMPUTATIONAL CHEMISTRY

FUNCTIONS & MODULAR CODE

FUNCTIONS

- ▶ Functions save programmers time and makes code simpler
- ▶ Essentially it is a way stop you having to write the same thing time and time again
- ▶ We have already encountered the use of functions
 - ▶ `print()`
 - ▶ `np.sum()`

FUNCTIONS

- ▶ We define a function, can pass it arguments and possibly receive information returned
- ▶ Not all functions need arguments or necessary return an object
- ▶ It is considered good form to add a *docstring* to all of your functions

FUNCTIONS



DEMO

FUNCTIONS

- ▶ Not all arguments are the same
- ▶ Those we have already seen included *required* arguments
- ▶ There is also *default* arguments, where the argument will default to a given value if not defined
- ▶ And *variable-length* arguments, essentially a list that is passed that can be any length

FUNCTIONS



DEMO

MODULES

- ▶ Another way to simplify your code is to store frequently used functions in *modules*
- ▶ Modules are files (with the file format `.py`) where we can store a lot of functions that we need to use
- ▶ Like libraries (such as NumPy) it is necessary to `import` any modules that we want to use in our code

MODULES



DEMO

PROBLEM

- ▶ The interaction energy between two atoms may be estimated using the Lennard-Jones potential model

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

- ▶ Where ϵ and r_m are constants describing the shape of the interaction energy surface

PROBLEM

- ▶ Add a function to calculate the interaction energy between two atoms to your `atom_helper.py` module

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

$$\epsilon = -2.2 \text{ kJ mol}^{-1}$$

$$r_m = 3.7 \text{ \AA}$$

PROBLEM

- ▶ Use this function to calculate the interaction energy between each of the pairs of atoms from Week 2

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

PROBLEM

- ▶ A common aim in computational chemistry is to fit the energetically optimised structure of a given chemical system
- ▶ To find this, we must find the energy minimum for all of the interactions
- ▶ This is achieved using an optimisation algorithm

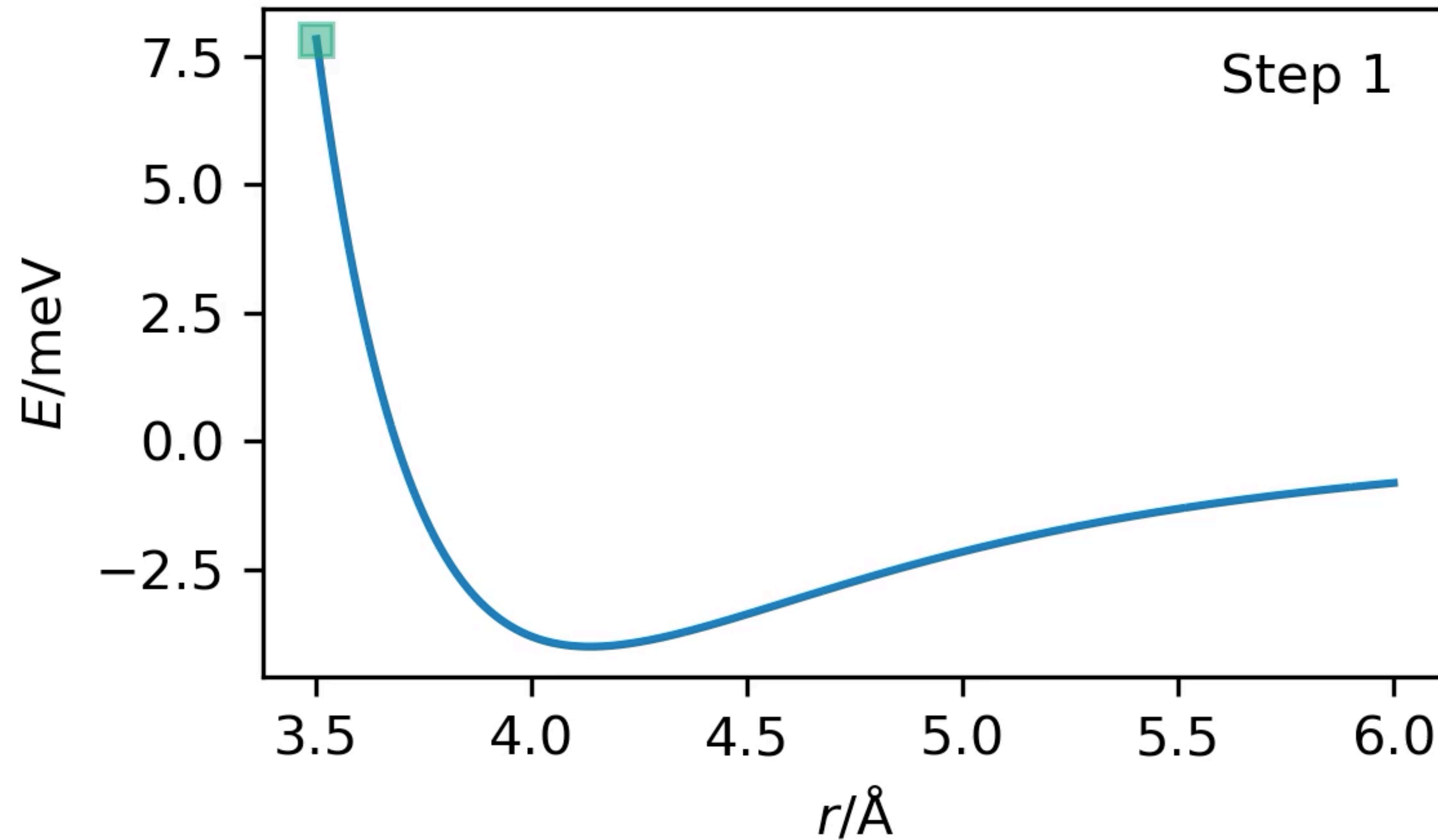
PROBLEM

- ▶ A popular, and simple, optimisation algorithm is the Newton-Raphson Method

$$r_1 = r_0 - \frac{E'}{E''}$$

- ▶ The distance between the two atoms is updated iteratively based on the first and second derivatives of the energy

PROBLEM



PROBLEM

- ▶ A popular, and simple, optimisation algorithm is the Newton-Raphson Method

$$r_1 = r_0 - \frac{E'}{E''}$$

- ▶ The distance between the two atoms is updated iteratively based on the first and second derivatives of the energy (given in the handout)

PROBLEM

- ▶ Write an implementation of the Newton-Raphson Method that will perform 10 iterations
 - ▶ Hint: it is not straightforward to implement with this to take advantage of NumPy optimisation
- ▶ Calculate the energy-minimum distance for the two sets of A and B

SCIPY.OPTIMIZE.MINIMIZE



DEMO