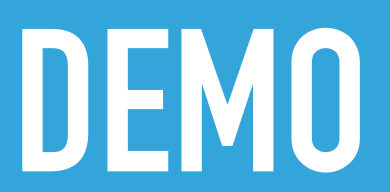
CH40208: TOPICS IN COMPUTATIONAL CHEMISTRY

FUNCTIONS & MODULAR CODE

- 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()

- 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



- 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



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



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

$$E_{LJ} = \frac{A}{r^{12}} - \frac{B}{r^6}$$

 \blacktriangleright Where A and B are constants describing the shape of the interaction energy surface

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

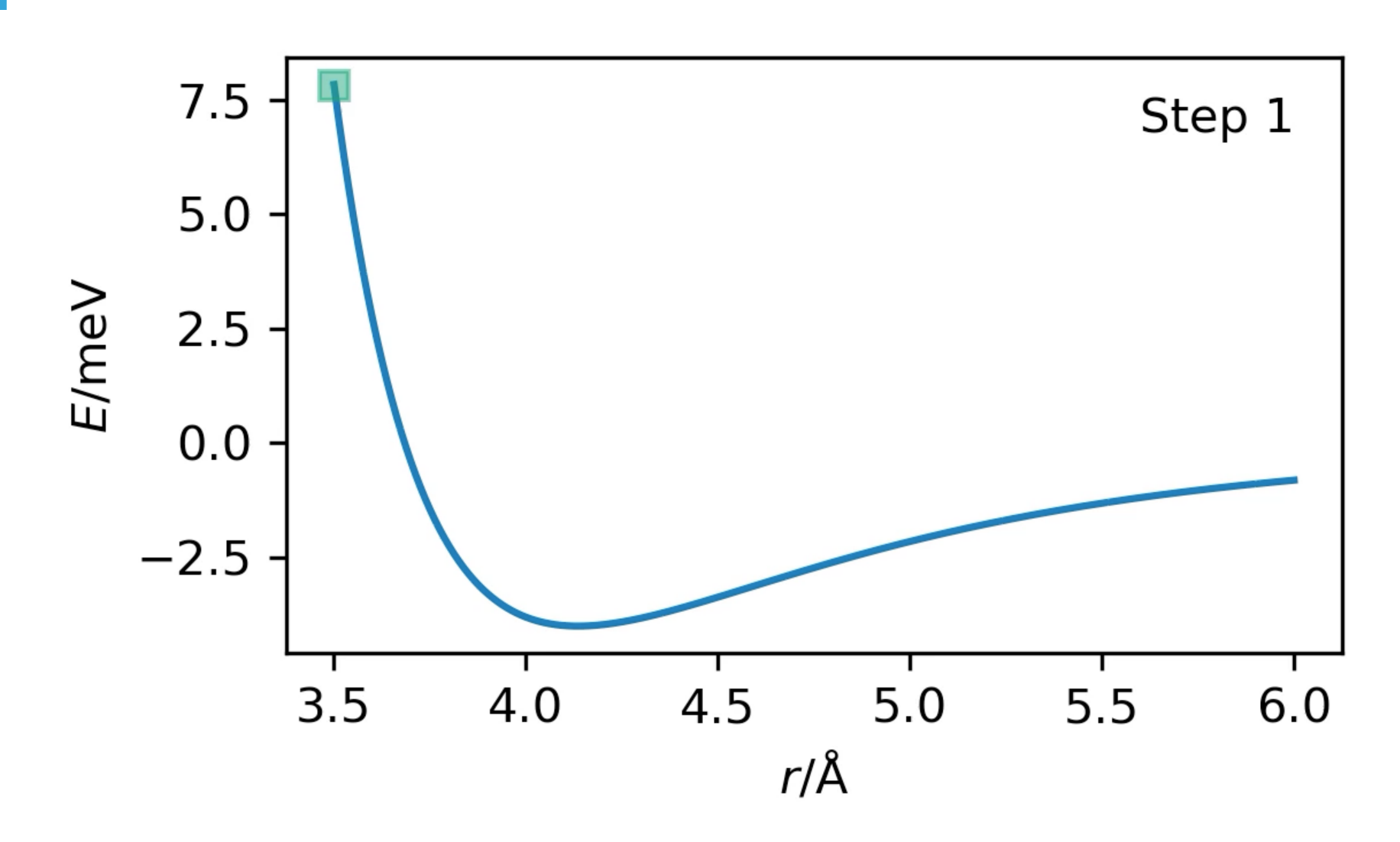
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- A common aim in computational chemistry is to fit the energetically optimised structure of a given chemical system
- To find this, we must find the energy minimum for all of the interactions
- This is achieved using an optimisation algorithm

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



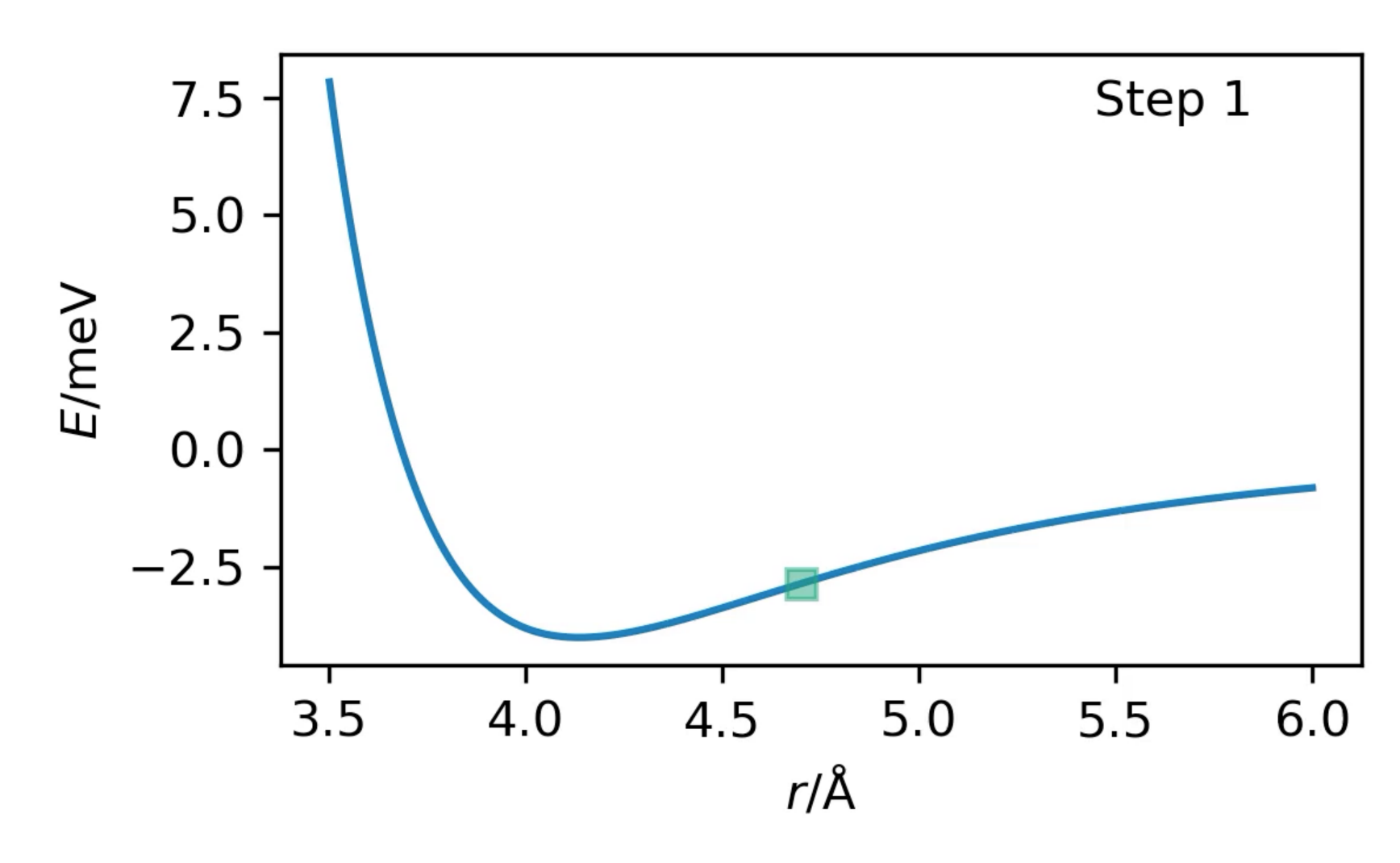
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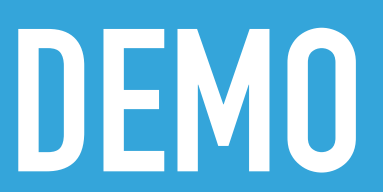
The distance between the two atoms is updated iteratively based on the first and second derivatives of the energy (given in the handout)

- 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

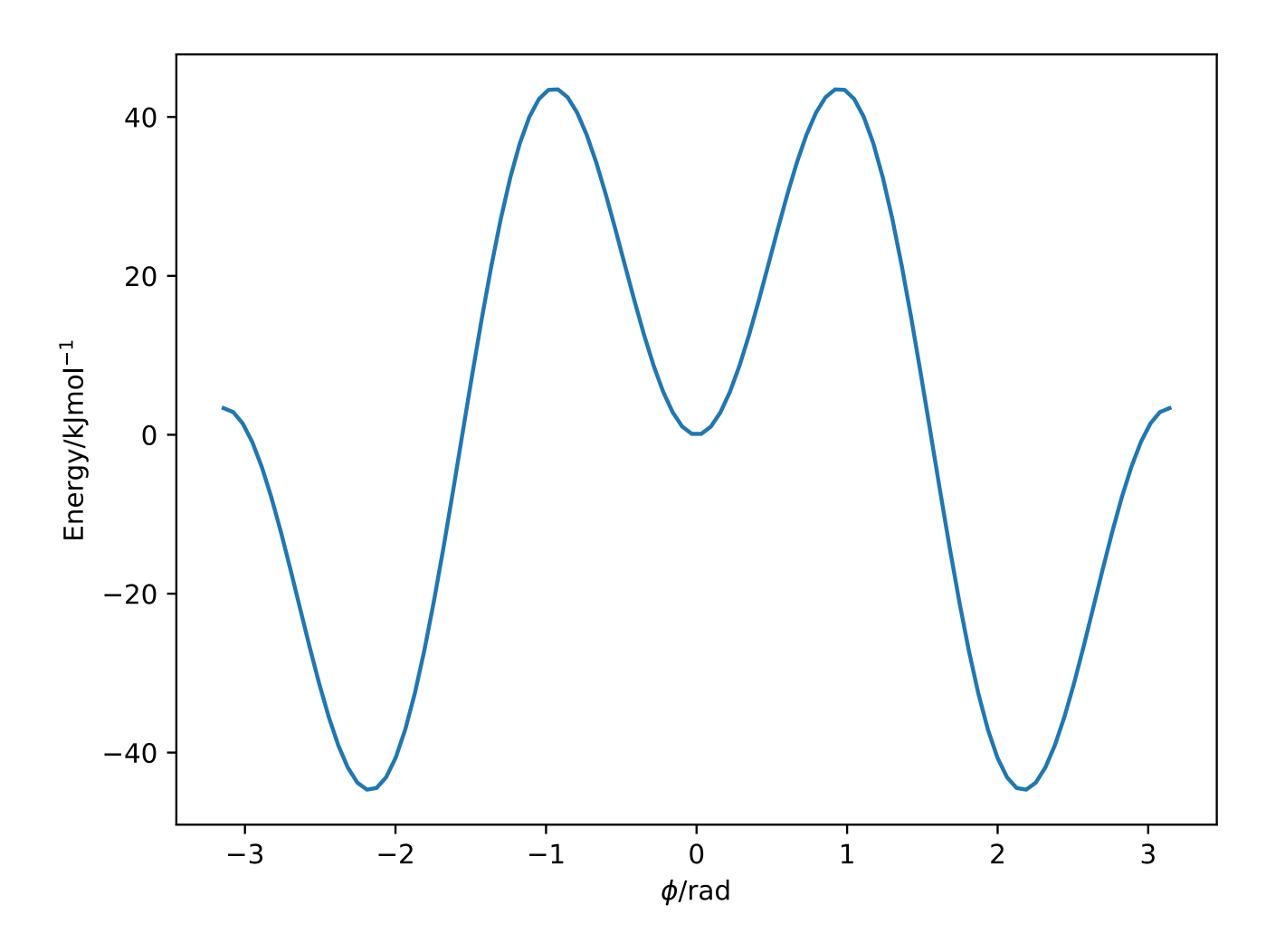
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LOCAL VS GLOBAL MINIMISATION



LOCAL VS GLOBAL OPTIMISATION

