
Introduction to Python for Data Analysis

How to manipulate and represent data with python

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Preamble

These notes contain the material for lecture of a [data scientist university degree](#) proposed at Université Clermont-Auvergne (UCA), which is mostly based on a [NumPy tutorial](#) that I gave at the [Laboratoire de Physique de Clermont](#) in April 2019. No prerequisite knowledge is assumed but being familiar with one programming language might be useful. It is better to know about some basic mathematics like simple vectorial operation or statistics. All the material of this lecture can be found in this [github repository](#).

This lecture is only a support to help you doing things yourself. As any other language, you must practice it if you want to progress. If you don't write and test code on your own, this lecture is close to be useless. I am available for any questions or general feedback on this lecture, so feel free to contact me: romain.madar@clermont.in2p3.fr.

General scope of the lecture

Python offers a rapidly evolving ecosystem to perform data analysis and it is both out of scope and hopeless to be extensive in this lecture. The main goal is therefore to make people familiar with the basic of python and data analysis tools in order to *make them able to extend their knowledge on themself*. Object oriented programming is not presented in this lecture. Practical exercises are also available to provide few working examples as a starting point.

What this lecture is? A *basic* and *practical* introduction to python together with some of the most important data analysis tools namely `numpy`, `matplotlib` and `pandas`.

What this lecture isn't? Neither a formal introduction to python, nor a extensive demonstration of all features available in the tools mentioned above.

Content of the lecture

There are a lot of information in this lecture. In order to help you to focus on important aspect, each chapter start with a list of expected skills that you should take away, ranked with three levels: *basic*, *medium*, *expert*.

1. Introduction to Python. This first section is dedicated to basic object type and operation in python. Fonctions will also be described but object oriented programming will not be covered – [online notebook](#)

2. Introduction to numpy. Differences between usual python objects and numpy objects will be introduced – [online notebook](#)

3. Three tools to know. This section gives a glimpse of `matplotlib`, `pandas` and `scipy` packages allowing powerful data analysis – [online notebook](#)

4. Multidimensional data manipulation. Non-trivial operation for multidimensional data using the full power of `numpy`. Most of these operation can be performed with existing tools but it is instructive to do it once with native `numpy` – [online notebook](#)

5. Introduction to image processing. Very first steps of image processing (definition, plotting, operation) including basic filters application (noising, sharpen, border detection) – [online notebook](#)

Other practical examples. Depending on the remaining time (and the people taste), we can go through different topics among the following ones. Some of them can be also used as a project performed by students.

- Fourier analysis
- Principal component analysis (PCA)
- Random Forest regression
- Gaussian processes

0.1 How to get prepared

1. Get familiar with python. I would recommend two links: [w3school tutorial](#) (both basic and complete) and <https://www.learnpython.org> (code can be ran directly within your web browser).

2. Install python with anaconda. In order to run python on your own machine, you should install it. I would recommend [anaconda](#) for this, which also includes jupyter-notebook.

3. Install git. This is a versioning software which can be installed following these [instructions](#). This whole repository can be *cloned* using `git clone https://github.com/radar/lecture-python` command.

4. Get familiar with notebooks. This represents a nice environment combining codes, notes and plots. This is very powerful to learn something and play with it. You can checkout [this video](#) or [this post](#).

Chapter 1

Practical Introduction to Python

Skills to take away

- *basic*: int/float/str, list/dictionnary, indexing/slicing, loops, functions, reading/writing files
- *medium*: docstring, comprehension, zip()/enumerate(), sorting dictionary
- *expert*: packing/unpacking, parsing file with correct casting, basic plotting

1.1 General information

Python can be installed using [anaconda](#). [Jupyter-notebook](#) (also coming with anaconda) is probably the easiest way to follow this lecture and make your own notes. The goal of this first chapter is to give a *very quick introduction basis*, but practice is mandatory to get comfortable with python objects and syntax. Practicing is possible with a web browser only using <https://www.learnpython.org>. A more complete tutorial (but not interactive) can be found in [w3school python tutorials](#). I recommend to follow the last tutorial up to *Arrays*.

In python, there is one instruction per line. Variable assignment is done with `=`, indentation is used to group instructions together under a loop or a condition block: there is no bracket (like in C++) or equivalent. Comments (*i.e.* uninterpreted text) start with `#`. Importation of external modules or function can be done with three different ways: `import module`, `import module as m` or `from module import this_function`.

In the following example, the result of the command will be printed so that people can check that the computer is doing what is expected. The instruction `print(x)` will print the content of `x`. If several variables are printed, is it convenient to use `print('x={} and y={}'.format(x, y))` syntax that will print `x` and `y` in bracket fields with one command - even if they have different types.

1.2 Object types

Numbers. There are three type of numbers: `int`, `float` and `complex`. The usual operations `(+, -, *, /)` are available. In addition, there is also `a**b` (which means a^b), `a // b` and `a % b` (which are the result of integer divisions - see example below).

```
# Basic numbers and operations
a = 2
b = 3.14
print(a+b)
print(a**b)
```

```
5.140000000000001
8.815240927012887
```

```
# Complex numbers and power
a = 1j
a**2
```

```
(-1+0j)
```

```
# Integer division example (// and % operators)
a, b = 10, 4
divisor, rest = a//b, a%b
print('{} = {}{}{}'.format(a, divisor, b, rest))
```

```
10 = 2x4 + 2
```

Strings. String allows to manipulate words, sentence or even text with specific methods. String are also python lists and list methods work as well (see below). The common and useful string manipulations can be counting the number of letters with `len(word)` or even manipulate a collection of words using `sentence.split('')`. Many methods exist, which can be looked at by typing `help(str)` in a python terminal or a jupyter notebook.

```
w1 = 'hello'
print(w1, len(w1), w1[3])
```

```
hello 5 l
```

```
# Summing two strings is possible (all other operators dont work)
blank, w2 = ' ', 'world'
sentence = w1 + blank + w2
print(sentence)
```

```
hello world
```

```
# Multiplying a string by an integer is also possible
repetition = sentence*3
print(repetition)
```

```
hello worldhello worldhello world
```

```
# Get a list of words from a sentence (cf. below for list objects)
s = 'It is rainy today'
list_words = s.split(' ')
print(list_words)
```

```
['It', 'is', 'rainy', 'today']
```

```
# Looping over the words and get the number of letters
for w in s.split(' '):
    print(w, len(w))
```

```
It 2
is 2
rainy 5
today 5
```

1.3 Object collections

There are four types of collection, which share several methods but differ from various aspects:

- list
- dictionnary
- tuple
- set

The most commonly used are the lists and dictionnary. The specificity of the set is that it is unordered, while the specificity of the tuple is that it cannot be modified. The common methods are

- number of items: `len(x)`
- loop over items with `for element in x:`
- check if an item is in the list: `element in x`

Lists. This is one of the most used collection object in python because it is the next-to-simplest level, after individual variables. A python list is a *list of objects* with possibly different types. One can search, loop, count with list. One can also add two lists or multiply a list by an integer, which makes a *concatenation* or a *duplication* (these points will be important for numpy arrays). The indexing of elements is also a nice way to access the information of interest: one can access the i^{th} element with `my_list[i]` or get a sub-list with `my_list[i:j]`. One can also take only one element every n with `my_list[i:j:n]` (more precisely this takes elements of index $i + p \times n$ until j , with $p = 0, 1, 2, \dots$). With this syntax, reverting the order of the list is easy: `reverted_list = my_list[::-1]`, where empty variable are default values (namely 0 and `len(my_list)`).

```
# Defining a list and access basic information
my_list1 = [1, 3, 4, 'banana']
print('Second element is {}'.format(my_list1[1]))
print('Number of elements: {}'.format(len(my_list1)))
print('Is \'banana\' in the list? {}'.format('banana' in my_list1))
```

```
Second element is 3
Number of elements: 4
Is 'banana' in the list? True
```

```
# Sum of two lists
my_list2 = ['string', 1+3j, [100, 1000]]
my_list = my_list1 + my_list2
print(my_list)
```

```
[1, 3, 4, 'banana', 'string', (1+3j), [100, 1000]]
```

```
# List multiplied by an integer
my_list = my_list*2
print(my_list)
```

```
[1, 3, 4, 'banana', 'string', (1+3j), [100, 1000], 1, 3, 4, 'banana', 'string', (1+3j), [100, 1000]]
```

```
# Looping over list element and print the type of seven first elements in the reversed order.
for element in my_list[6:0:-1]:
    print('{} is {}'.format(element, type(element)))
```

```
[100, 1000] is <class 'list'>
(1+3j) is <class 'complex'>
string is <class 'str'>
banana is <class 'str'>
4 is <class 'int'>
3 is <class 'int'>
```

sets and tuples. Tuples and sets are modified version of python list. Tuples are ordered but cannot be modified (no assignment, no addition, while sets are not ordered but can be modified). In this context, order means indexing (so `x[i:j:n]` syntax, among others). Search or loop over elements work in the same way as for list.

```
# Tuple
t = (1, 3, 7)
print(t)

# Access the third element
```

```

print(t[2])

# Try to modify the second element - using the 'try - except' synthax
try:
    t[1] = 'hello'
except TypeError:
    print('Impossible to change the value of a tuple')

```

```

(1, 3, 7)
7
Impossible to change the value of a tuple

```

Sets can modified with methods like `s.add(x)` or `s.update([x, y])`.

```

# Set
s = {'apple', 'banana', 'orange'}
print(s)

# Add one element
s.add('pineapple')
print(s)

# Add a list
s.update(['pear', 'prune'])
print(s)

```

```

{'orange', 'banana', 'apple'}
{'pineapple', 'orange', 'banana', 'apple'}
{'orange', 'pineapple', 'apple', 'pear', 'prune', 'banana'}

```

```

# Try to access the second element - using the 'try - except' synthax
try:
    print(s[1])
except TypeError:
    print('Impossible to access element via indexing')

```

```
Impossible to access element via indexing
```

Dictionnaries. Various object types are important to manipulate and organize data. The most common one is the dictionnary which work with a pair of (key, value). The key must be a non-modifiable object, in practice string or integer, but cannot be a list. This is a very powerful concept to store different types of information into the same object. One can easily loop, search, modify a given key value, or even add a new key quite easily.

```

# dictionary
person = {'name': 'Charles', 'age': 78, 'size': 173, 'gender': 'M'}
print(person)

```

```
{'name': 'Charles', 'age': 78, 'size': 173, 'gender': 'M'}
```

```
# Accessing value using the key
template = '{} ({} ) is {} years old and is {} cm'
print(template.format(person['name'], person['gender'], person['age'], person['size']))
```

```
Charles (M) is 78 years old and is 173 cm
```

```
# Adding a key and its value
person['eyes'] = 'blue'
print(person)
```

```
{'name': 'Charles', 'age': 78, 'size': 173, 'gender': 'M', 'eyes': 'blue'}
```

```
# Test if a key is present
print('name' in person)
print('brand' in person)
```

```
True
False
```

1.4 Loops

Loops are at the core of programming and especially for data analysis oriented tasks. There are two way of repeating a instruction several times: the `for` loop and the `while` loop. Several instructions are common to both loops, such as `continue` (skip instruction after and switch to the next element) or `break` (stop the loop), but the use case of these two ways are different.

For loops. For data analysis, I think these are the most used ones. But as we will see in the introduction to numpy, for loops must not be used for heavy computations in python. For loops are relevant for small (~1000) data samples (and computations). We'll come back to this point in the lecture. Below, few example are given.

```
# Compute sum(i^2) for i from 0 to 9
x = 0
for i in range(0, 10):
    x += i**2
print(x)
```

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```
# Loop over fruit via a set and print only ones with a 'p'
for fruit in s:
    if 'p' in fruit:
        print(fruit)
```

```
pineapple
apple
pear
prune
```

There are several ways to loop over dictionary depending on how we want to access the information. Indeed, you can access information by keys, values, or both. An example of each are given below.

```
# Loop over keys for a dictionnary and access the value of each
for properties in person:
    value = person[properties]
    print('{}: {}'.format(properties, value))
```

```
name: Charles
age: 78
size: 173
gender: M
eyes: blue
```

```
# Loop over dictionnary values only
for v in person.values():
    print(v)
```

```
Charles
78
173
M
blue
```

```
# Loop over both keys and values directly
for key, value in person.items():
    print('{}: {}'.format(key, value))
```

```
name: Charles
age: 78
size: 173
gender: M
eyes: blue
```

Tip: how to sort a dictionnary? It can be noted that dictionary are natively *not ordered*. This means that you cannot access a item with its index, since there is no index. However, it can be convenient to sort dictionary keys according to their values, using the general python function `sorted(collection, key=function)` and a type of object called `OrderedDict` from `collections` module, as explained below.

```
# Define a dictionnary
students_marks = {
```

```

'Jean': 12,
'Chloe': 17,
'Olivier': 8,
'Helene': 10
}

# Print the key, values items
for name, mark in students_marks.items():
    print(name, mark)

```

```

Jean 12
Chloe 17
Olivier 8
Helene 10

```

The `sorted()` function works on any type of collection than can be looped over (called *iterable*). It needs the collection and the function which return a key on which to sort for each object of the collection. This can be for e.g. a letter or a number. Let's try both by defining a function getting either the mark or the first letter of the name, from a dictionnary item (`name, mark`).

```

# Order it by increasing marks
def get_mark(item):
    return item[1]

# Or by the first letter of the name
def get_name(item):
    return item[0][0]

# Testing with an item
item_test = ('Jacques', 12)
print('{} has a mark of {} and {} as 1st letter'.format(item_test, get_mark(item_test),
    get_name(item_test)))

```

```
('Jacques', 12) has a mark of 12 and J as 1st letter
```

We can now apply the `sorted()` function to the collection of items of the initial directory. This will return a collection of sorted items that can be later converted into a dictionnary. This last step depends on python version (in version 2.5, one has to use `OrderDict` from `collections` module while it's not needed in python 3 - the version of python can be dynamically checked with `sys.version_info` from `sys` module).

```

# Get all items and sort them by increasing mark
all_items = students_marks.items()
items_sorted_by_marks = sorted(all_items, key=get_mark)
items_sorted_by_names = sorted(all_items, key=get_name)

```

```
# Check the version of python
import sys
version = sys.version_info[0]
isPython2 = version == 2

# Final conversion of collection of items into a dictionnary
if isPython2:
    from collections import OrderedDict
    marks_sorted_dict = OrderedDict(items_sorted_by_marks)
    names_sorted_dict = OrderedDict(items_sorted_by_names)

else:
    marks_sorted_dict = {k: v for k, v in items_sorted_by_marks}
    names_sorted_dict = {k: v for k, v in items_sorted_by_names}

# Check the mark sorted results:
for k, v in marks_sorted_dict.items():
    print(k, v)
```

Olivier 8
 Helene 10
 Jean 12
 Chloe 17

```
# Check the name sorted results:
for k, v in names_sorted_dict.items():
    print(k, v)
```

Chloe 17
 Helene 10
 Jean 12
 Olivier 8

While loops. They are bit less used in practice but they are quickly described for completeness. The idea is to repeat a given instruction until a condition is reached.

```
# Cast (ie change type) the set s into a list
my_list = list(s)

# Remove item one by one until there are no items left.
while len(my_list)>0:
    my_list.pop()
    print(my_list)
```

['orange', 'pineapple', 'apple', 'pear', 'prune']
 ['orange', 'pineapple', 'apple', 'pear']

```
['orange', 'pineapple', 'apple']
['orange', 'pineapple']
['orange']
[]
```

1.5 Few python synthax tips

Comprehension. This is the action of building a collection with one line of code. The comprehension syntax work for all collections, with conditions, or even nested loops (loops of loops). Few examples are given below.

```
# List
list_squares = [i**2 for i in range(1, 10)]
print(list_squares)

# Dictionnary
dict_squares = {i:i**2 for i in list_squares[0:5]}
print(dict_squares)
```

```
[1, 4, 9, 16, 25, 36, 49, 64, 81]
{1: 1, 4: 16, 9: 81, 16: 256, 25: 625}
```

```
# Comprehension list with a condition (e.g. keep only even numbers)
list_even = [i for i in range(0, 10) if i%2==0]
print(list_even)
```

```
[0, 2, 4, 6, 8]
```

```
# Comprehension with nested loops
sum_integers = [i*10+j for i in range(0,5) for j in range(0, 5)]
print(sum_integers)
```

```
[0, 1, 2, 3, 4, 10, 11, 12, 13, 14, 20, 21, 22, 23, 24, 30, 31, 32, 33, 34, 40, 41, 42, 43, 44]
```

Looping with enumerate and zip.

The keyword enumerate return directly a counter together with the element arising in the loop. This is useful if you need to count the number of iterations of the loop. This can be done without enumerate but you need to add two lines (initialisation of the counter, and incrementation).

```
# Position of each word in a sentence
sentence = 'I would like to analyse this sentence in term of word position'
words = sentence.split(' ')
for i, w in enumerate(words):
    print(w.ljust(10) + ': ' + str(i))
```

```
I      : 0
would : 1
like   : 2
to     : 3
analyse : 4
this   : 5
sentence : 6
in     : 7
term   : 8
of     : 9
word   : 10
position : 11
```

The `zip(list1, list2)` synthax allows to form pairs using elements of each list at *the same position*. This is quite convenient to associate some objects which are stored in different collections in a very quick and readable way. If `list1` and `list2` don't have the same size, the minimum of the two lenght is taken. Finally, the `zip()` command can take more than two collections and return then a group of element which has the size of the number of collection.

```
# Associate fruits and colors
fruits = ['banana', 'orange', 'pineapple', 'pear', 'prune']
colors = ['yellow', 'orange', 'brown', 'green', 'purple']
for f, c in zip(fruits, colors):
    print('{} is {}'.format(f, c))
```

```
banana is yellow
orange is orange
pineapple is brown
pear is green
prune is purple
```

```
# Using zip() with three lists
l1, l2, l3 = range(0, 10), range(0, 100, 10), range(0, 1000, 100)
for i, j, k in zip(l1, l2, l3):
    print(i, j, k, i+j+k)
```

```
0 0 0 0
1 10 100 111
2 20 200 222
3 30 300 333
4 40 400 444
5 50 500 555
6 60 600 666
7 70 700 777
8 80 800 888
9 90 900 999
```

1.6 Functions

Functions are defined as a set of instruction encapsulated into one object. This is particularly convenient when one has to the same list of instructions several times. A good guideline to know when to write a function could be

If you copy-paste the same piece of code more than two times, then make a function

Definition. A function takes some arguments, perform some instruction and return a result. The syntax to define and call a function is showed below. In python, function must be defined *before* being used (as opposed to C++ where it can be different, as soon as the function is declared). This makes the concept of *package* quite relevant to wrap-up several function into a python file which can be imported in the main code.

```
# Definition syntax
def function(argument):
    result = argument * 3
    return result

# Call syntax
function(2)
```

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The type of the argument is not fixed (since it is a general feature of python) so the same instruction will be interpreted differently depending on the type. The following example shows the different result of the function above for two argument types.

```
# Print the result for two types of arguments
print('function(10) = {}'.format(function(10)))
print('function(\\"ouh\\") = {}'.format(function('ouh')))

function(10) = 30
function('ouh') = ouhouhouh

def person_printer(p):
    template = '{} {} is {} years old and is {} cm'
    print(template.format(p['name'], p['gender'], p['age'], p['size']))
    return

def grow_old(p, n_years):

    # 1. copy the dictionary person (otherwise p *will* be modified)
    res = p.copy()

    # 2. Compute the new age and size
    new_age = res['age'] + n_years
```

```

new_size = res['size'] - n_years*0.13

# 3. Assign the new age/size to the result
res['age'] = new_age
res['size'] = new_size

# 4. Return the result
return res

```

```

# Print before growing old
person_printer(person)

# Growing old
old_guy = grow_old(person, 10)

# Print after growing old
person_printer(old_guy)

```

Charles (M) is 78 years old and is 173 cm
 Charles (M) is 88 years old and is 171.7 cm

Docstring.

This offers the possibility to document your code in a proper way, which is quite useful for others (and for you, when you will re-use a code after several years). This is then a good practice to do, even if it takes time. This can be accessed using the command `help(function)` or by using the keyboard shortcut Shift+Tab in jupyter notebook (when the cursor is after the opening parenthesis of the function). The syntax to add docstring is `'''My documentation'''` at the very begining of the function.

```

def grow_old(p, n_years):
    """
    Take a person dictionnary and update the age and the size to make the person older.

    Parameters
    -----
    p: dictionnary
        Person object as defined earlier in the code, with at least 'age' (year)
        and 'size' (cm) keys, to get old.
    n_years: integer
        Number of years to be added to the age of the person.

    Returns
    -----
    person: dictionnary
        Person object as defined earlier in the code with age and size updated as
        age -> age+n_years
        size -> size-n_years*0.13
    """

```

```
# 1. copy the dictionary person (otherwise p will be modified, which might problematic)
res = p.copy()

# 2. Compute the new age and size
new_age = res['age'] + n_years
new_size = res['size'] - n_years*0.13

# 3. Assign the new age/size to the result
res['age'] = new_age
res['size'] = new_size

# 4. Return the result
return res
```

```
help(grow_old)
```

Help on function grow_old in module __main__:

```
grow_old(p, n_years)
    Take a person dictionary and update the age and the size to make the person older.

Parameters
-----
p: dictionary
    Person object as defined earlier in the code, with at least 'age' (year)
    and 'size' (cm) keys, to get old.
n_years: integer
    Number of years to be added to the age of the person.

Return
---
person: dictionary
    Person object as defined earlier in the code with age and size updated as
        age  -> age+n_years
        size -> size-n_years*0.13
```

There are several ways to organise the docstring and the example above is based on numpy docstring style. Note that docstring can be also added to a module (in practice, a python file) to document the content, goal and usage of this module.

Arbitrary number of arguments: *args and **kwargs. The example above are relatively simple and generally function takes several arguments. Sometime it is even convenient to have an unfixed number of arguments, so that the function is rather evolutive when the code grows. Python offres a nice way to define such a function thanks to the *packing* and *unpacking* notion, which is describe right below.

Apparte: packing and unpacking. In short, this is the possibility to convert a list into a serie of objects (unpacking) or vis-versa (packing). This way of writing collection makes code developments very concise and fast, especially to call function with a several arguments in a nice way. This also allows to define function with an

arbitrary number of arguments as already mentioned. The following dummy function is used to illustrate the concept of packing/unpacking with both a list and a dictionary.

```
# Test function
def mean(a, b, c):
    return (a+b+c)/3.
```

It is possible to use a list of three numbers to specify the argument values of the `mean(a, b, c)` function, using the unpacking syntax for list: `*list`. This is demonstrated below:

```
# Packing & unpacking with a list (or a tuple): *list
my_numbers = [10, 12, 15]
mean(*my_numbers)
```

12.33333333333334

This is also sometimes convenient to call the argument by their name (mostly to make the code more readable). This type of arguments are called *keyword arguments* and can be packed/unpacked into a dictionary. Each argument name is a key of this dictionary and the value is the value passed to the function. The unpacking is done with `**dict`.

```
# Packing & unpacking with a dictionary: **dict
my_numbers = {'a': 10, 'b': 12, 'c': 15}
mean(**my_numbers)
```

12.33333333333334

Coming back to the initial motivation, i.e. having an arbitrary number of arguments. It is possible to define such a function as follows - which in that case just prints the number and the list of arguments:

```
# Function definition with *args
def test_function(*args):
    print('There are {} arguments: {}'.format(len(args)))
    for a in args:
        print(' -> {}'.format(a))
    print('')
    return
```

```
# Test with different numbers/types of arguments
test_function()
test_function('hoho')
test_function('hoho', 3)
test_function('hoho', 3, [1, 'banana'], {'mood': 'happy', 'state': 'holidays'})
```

There are 0 arguments:

There are 1 arguments:

-> hoho

There are 2 arguments:

-> hoho
-> 3

There are 4 arguments:

-> hoho
-> 3
-> [1, 'banana']
-> {'mood': 'happy', 'state': 'holidays'}

```
# Function definition with kwargs
def test_function(**kwargs):
    print('There are {} arguments: {}'.format(len(kwargs)))
    for k, v in kwargs.items():
        print(' {}={}'.format(k, v))
    print('')
    return
```

```
test_function()
test_function(x='hoho')
test_function(word='hoho', multiplicity=3)
test_function(a='hoho', N=3, shopping=[1, 'banana'], feeling={'mood': 'happy', 'state':
    'holidays'})
```

There are 0 arguments:

There are 1 arguments:
x=hoho

There are 2 arguments:
word=hoho
multiplicity=3

There are 4 arguments:
a=hoho
N=3
shopping=[1, 'banana']
feeling={'mood': 'happy', 'state': 'holidays'}

This can be used to declare argument in a very readable and concise way. This might be helpful for some cosmetic argument of plot that can be common to several plots (but not all). We'll see some concrete example later in the lecture. In the meanwhile, here is the equivalent of last call from the code above:

```
# Pack all keyword arguments in a dictionary first
my_args = {
    'a': 'hoho',
    'N': 3,
    'shopping': [1, 'banana'],
    'feeling': {'mood': 'happy', 'state': 'holidays'}
}

# Then call the function
test_function(**my_args)
```

There are 4 arguments:

```
a=hoho
N=3
shopping=[1, 'banana']
feeling={'mood': 'happy', 'state': 'holidays'}
```

1.7 File manipulation

File handling is quite important since it enable interaction between your code and input/output data (called I/O). There are several features related to file handing in python and this short section just give few basic practices.

Open/close a file. Python has native methods to open and close file. While closing a file doesn't allow for many variations, the opening can be done in different mode, depending if we want to read, write or append to the opened file. The basic syntax is:

```
# Open
f = open('my_file_name.txt', option)

# Close
f.close()
```

where option is a one letter string which can be:
+ r read (default): to just read the file
+ a append: to add content at the end of an existing file
+ w write: to write content in a file (it creates a new file)
+ x create: to create a new file

Write a file.

```
# Text to be written (can be one line string 'my text' or multiple lines string - docstring)
text = '''Gervaise avait attendu Lantier jusqu'à deux heures du matin. Puis,  

toute frissonnante d'être restée en camisole à l'air vif de la fenêtre,  

elle s'était assoupie, jetée en travers du lit, fiévreuse, les joues  

trempées de larmes.  

...  

# Open in write mode
```

```
f = open('test.txt', 'w')

# Write string
f.write(text)

# Close
f.close()
```

Read a file. The following example load the precedent file and loop over each line to analyse its content. One can note several issues: first one sentence can be on two lines, and second, each end of line contain a \n (which is an invisible character meaning “go-to-next-line”). There is a method to clean a line, called `line.strip()` and remove all spaces and invisible characters, unless specified otherwise - see `help(str.strip)`.

```
# Open the file in read mode
f = open('test.txt', 'r')

# Loop over the lines
for line in f:

    # Print a header to make the output clearer
    print('\n\n{}'.format('='*50))

    # Print the line as it is given
    print('This line: {}'.format(line))

    # Split by '.' to isolate sentence
    sentences = line.split('.')
    print('This line has {} sentences: {}'.format(len(sentences)))

    # Split each sentence by ' ' to isolate words
    for i,s in enumerate(sentences):
        sclean = s.strip()
        words = sclean.split(' ')
        print(' - sentence {}: {}'.format(i, words))

f.close()
```

```
=====
This line: Gervaise avait attendu Lantier jusqu'à deux heures du matin. Puis,
This line has 2 sentences:
 - sentence 0: ['Gervaise', 'avait', 'attendu', 'Lantier', 'jusqu'à', 'deux', 'heures', 'du',
'matin']
 - sentence 1: ['Puis,']
```

```
=====
This line: toute frissonnante d'être restée en camisole à l'air vif de la fenêtre,
```

```
This line has 1 sentences:
- sentence 0: ['toute', 'frissonnante', 'd'être', 'restée', 'en', 'camisole', 'à', 'l'air',
  'vif', 'de', 'la', 'fenêtre', ]
```

```
=====
This line: elle s'était assoupie, jetée en travers du lit, fiévreuse, les joues
```

```
This line has 1 sentences:
- sentence 0: ['elle', 's'était', 'assoupie', 'jetée', 'en', 'travers', 'du', 'lit',
  'fiévreuse', 'les', 'joues']
```

```
=====
This line: trempées de larmes.
```

```
This line has 2 sentences:
- sentence 0: ['trempées', 'de', 'larmes']
- sentence 1: []
```

Re-write a modified version of a file into a new file. It can be quite convenient to modify an existing file to correct a systematical mistake automatically, or simply do more complexe operation. The example below shows how to remove all the “e” from the text below and write it in a new file.

```
# Open the in/out files
f_i = open('test.txt', 'r')
f_o = open('test_without_e.txt', 'w')

# Loop over line, remove all "e" for each, and write the result in the output file
for line in f_i:
    line_without_e = line.replace('e', '') # replace "e" by nothing
    f_o.write(line_without_e)

# Close all files
f_i.close()
f_o.close()
```

```
# Open in read mode and check the result
f = open('test_without_e.txt', 'r')
print(f.read())
```

```
Grvais avait attndu Lantir jusqu'à dux hurs du matin. Puis,
tout frissonnant d'êtr rsté n camisol à l'air vif d la fnêtr,
ll s'était assoupi, jté n travrs du lit, fiévrus, ls joues
trmpés d larmes.
```

Read a csv file to get data.

This use case is quite important since it allows to convert a file with data into variables accessible in the code (for some computation, plotting, etc ...). One the most basic format to store data is called *csv* (for comma-separated values) which can import/export from any spreadsheet software (like excel). This format is not necessarily appropriate for large dataset but is quite useful if a large number of situations. one must be able to manipulate it easily, as shown in the example below.

Creation of a csv file on the fly using a docstring

```
# Data taken from kaggle: https://www.kaggle.com/jolasa/waves-measuring-buoys-data-mooloolaba
data_csv_format = '''index,date,height,heightMax,period,energy,direction,temperature
1,01/01/2017 00:00,-99.9,-99.9,-99.9,-99.9,-99.9,-99.9
2,01/01/2017 00:30,0.875,1.39,4.421,4.506,-99.9,-99.9
3,01/01/2017 01:00,0.763,1.15,4.52,5.513,49,25.65
4,01/01/2017 01:30,0.77,1.41,4.582,5.647,75,25.5
5,01/01/2017 02:00,0.747,1.16,4.515,5.083,91,25.45
6,01/01/2017 02:30,0.718,1.61,4.614,6.181,68,25.45
7,01/01/2017 03:00,0.707,1.34,4.568,4.705,73,25.5
8,01/01/2017 03:30,0.729,1.21,4.786,4.484,63,25.5
9,01/01/2017 04:00,0.733,1.2,4.897,5.042,68,25.5
10,01/01/2017 04:30,0.711,1.29,5.019,8.439,66,25.5
11,01/01/2017 05:00,0.698,1.11,4.867,4.584,64,25.55
12,01/01/2017 05:30,0.686,1.14,4.755,5.211,56,25.55
13,01/01/2017 06:00,0.721,1.12,4.843,5.813,67,25.5
14,01/01/2017 06:30,0.679,1.22,4.948,4.71,81,25.45
15,01/01/2017 07:00,0.66,1.08,5.068,5.353,90,25.45
16,01/01/2017 07:30,0.662,1.18,5.263,7.436,67,25.4
17,01/01/2017 08:00,0.653,1.21,5.007,6.001,90,25.45
18,01/01/2017 08:30,0.665,1.17,4.952,6.414,90,25.55
19,01/01/2017 09:00,0.684,1.55,5.022,6.691,88,25.6
20,01/01/2017 09:30,0.679,1.09,4.926,6.804,88,25.65
21,01/01/2017 10:00,0.667,1.12,4.928,6.641,122,25.75
22,01/01/2017 10:30,0.688,1.13,4.808,5.958,91,25.7
23,01/01/2017 11:00,0.644,0.99,4.559,6.691,92,25.9
'''

# Create csv file using these data
f = open('wave_data.csv', 'w')
f.write(data_csv_format)
f.close()
```

Reading the csv file and storing values in python objects. In this example, we'll see how to store all information about the wave in a list of dictionnaries.

```
# Open the file in read mode
f = open('wave_data.csv', 'r')

# Get the first line (calling the readline() method once) to extract the feature names.
features = f.readline().strip().split(',')
print(features)
```

```

# Loop over lines and store the information
data = []
for l in f:
    values = l.strip().split(',')
    data_single_wave = {var: val for var, val in zip(features, values)}
    data.append(data_single_wave)

['index', 'date', 'height', 'heightMax', 'period', 'energy', 'direction', 'temperature']

# helper function for a nice printing
def print_wave(w):
    tmp = 'Wave {} ({} {}) had a heigh of {}m with a temperature of {} degree'
    print(tmp.format(w['index'], w['date'], w['height'], w['heightMax'], w['temperature']))

# Print the first 5 waves
for wave in data[:5]:
    print_wave(wave)

```

Wave 1 (01/01/2017 00:00) had a heigh of -99.9m with a temperature of -99.9 degree
 Wave 2 (01/01/2017 00:30) had a heigh of 0.875m with a temperature of -99.9 degree
 Wave 3 (01/01/2017 01:00) had a heigh of 0.763m with a temperature of 25.65 degree
 Wave 4 (01/01/2017 01:30) had a heigh of 0.77m with a temperature of 25.5 degree
 Wave 5 (01/01/2017 02:00) had a heigh of 0.747m with a temperature of 25.45 degree

At the stage, the problem is that the type of object which are stored is string and not numbers ... so no computation can be made. Typically, the following code will crash because the division between string is not defined:

```

# Compute the average height
heights = [w['height'] for w in data]
average = sum(heights)/len(heights)

```

One has to cast (i.e. change type) the object stored into the dictionnaries. They can all casted as float but the date. The index makes more sense as integer as well. So the following can work:

```

# Manage string to time object conversion
def str_to_time(date_str):
    import datetime
    return datetime.datetime.strptime(date_str, '%d/%m/%Y %H:%M')

# Container with properly converted data
DATA = []

# Loop over waves and make the proper conversion depending on the feature name
for w in data:

```

```
wgood = w.copy()
for k in w:
    if k == 'index':      # Cast string into an integer
        wgood[k] = int(w[k])
    elif k == 'date':     # cast string into a datetime object
        wgood[k] = str_to_time(w[k])
    else:                 # cast string into a float
        wgood[k] = float(w[k])
DATA.append(wgood)
```

Computing the averaged heigh of wave now works but give a quite strange result:

```
# Compute the average height
heights = [w['height'] for w in DATA]
average = sum(heights)/len(heights)
print('Averaged waveheight is {:.1f} m'.format(average))
```

Averaged waveheight is -3.7 m

This is due to the first row which has all values at -99. Removing it (using indexing technics) gives a more sensible result:

```
heights = [w['height'] for w in DATA[1:]]
average = sum(heights)/len(heights)
print('Averaged waveheight is {:.1f} m'.format(average))
```

Averaged waveheight is 0.7 m

1.8 Plotting data: the very first step

Graphical representation of data is a key element of data analysis: it allows to get some intuition (and then ideas), or just perform visual checks. You might find the terminology “Exploratory Data Analysis (EDA)” in the litterature, which correspond to plot data in all possible way to extract exploitable information from data. The EDA is an entire field which will not cover in this lecture. We will simply gives some basic examples, which will provide a starting point to expand your knowledge. The standard library to produce plots is `matplotlib`, but it exist many other tools that we will not introduce (e.g. `seaborn`, `bokeh` for browser-interactive plots, `cartopy` for geographic data analysis/plots, etc.).

```
# Prepare data to plot: wave height v.s. wave energy (removing point with -99 values)
height = [w['height'] for w in DATA if w['height'] > -99]
energy = [w['energy'] for w in DATA if w['energy'] > -99]
```

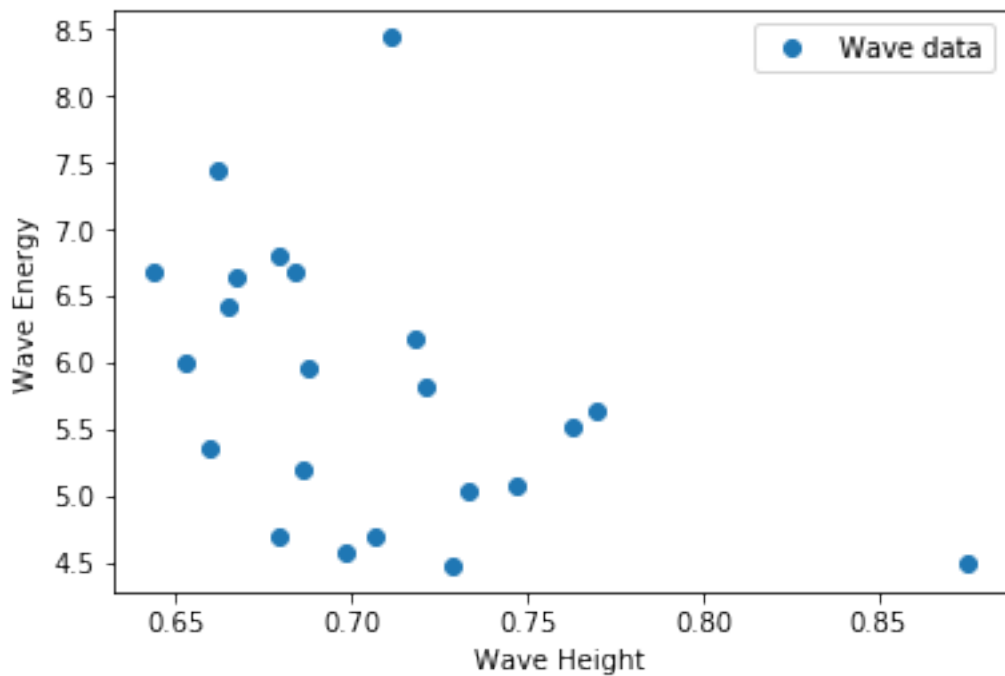
```
# Import the key part of the package: pyplot
import matplotlib.pyplot as plt

# Display matplotlib output in the notebook
%matplotlib inline

# Call the simplest function to plot x vs y
plt.plot(height, energy, linewidth=0, marker='o', label='Wave data')

# Set x and y axis axis labels
plt.xlabel('Wave Height')
plt.ylabel('Wave Energy')

# Adding a legend based on label keyword
plt.legend();
```



Chapter 2

Basic introduction to NumPy

Why numpy? Numpy stands for *numerical python* and is highly optimized (and then fast) for computations in python. Numpy is one of the core package on which many others are based on, such as scipy (for *scientific python*), matplotlib or pandas (described at the end of this chapter). A lot of other scientific tools are also based on numpy and that justifies to have - at least - a basic understanding of how it works. Very well, but one could also ask why using python?

Why python? Depending on your preferences and your purposes, python can be a very good option or not (of course this is largely a matter of taste and not everyone agrees with this statement). In any case, many tools are available in python, scanning a very broad spectrum of applications, from machine learning to web design or string processing.

2.1 The core object: arrays

The core of numpy is the called numpy array. These objects allow to efficiently perform computations over large dataset in a very concise way from the language point of view, and very fast from the processing time point of view. The price to pay is to give up explicit *for* loops. This lead to somehow a counter intuitive logic - at first.

2.1.1 Main differences with usual python lists

The first point is to differentiate numpy array from python list, since they don't behave in the same way. Let's define two python lists and the two equivalent numpy arrays.

```
import numpy as np
l1, l2 = [1, 2, 3], [3, 4, 5]
a1, a2 = np.array([1, 2, 3]), np.array([3, 4, 5])
print(l1, l2)
```

```
[1, 2, 3] [3, 4, 5]
```

First of all, all mathematical operations act element by element in a numpy array. For python list, the addition acts as a concatenation of the lists, and a multiplication by a scalar acts as a replication of the lists:

```
# obj1+obj2
print('python lists: {}'.format(l1+l2))
print('numpy arrays: {}'.format(a1+a2))
```

```
python lists: [1, 2, 3, 3, 4, 5]
numpy arrays: [4 6 8]
```

```
# obj*3
print('python list: {}'.format(l1*3))
print('numpy array: {}'.format(a1*3))
```

```
python list: [1, 2, 3, 1, 2, 3, 1, 2, 3]
numpy array: [3 6 9]
```

One other important difference is about the way to access element of an array, the so called slicing and indexing. Here the behaviour of python list and numpy arrays are closer expect that numpy array supports few more features, such as indexing by an array of integer (which doesn't work for python lists). Use cases of such indexing will be heavily illustrated in the next chapters.

```
# Indexing with an integer: obj[1]
print('python list: {}'.format(l1[1]))
print('numpy array: {}'.format(a1[1]))
```

```
python list: 2
numpy array: 2
```

```
# Indexing with a slicing: obj[slice(1,3)]
print('python list: {}'.format(l1[slice(1,3)]))
print('numpy array: {}'.format(l1[slice(1,3)]))
```

```
python list: [2, 3]
numpy array: [2, 3]
```

```
# Indexing with a list of integers: obj[[0,2]]
print('python list: IMPOSSIBLE')
print('numpy array: {}'.format(a1[[0,2]]))
```

```
python list: IMPOSSIBLE
numpy array: [1 3]
```

2.1.2 Main characteristics of an array

The strength of numpy array is to be multidimensional. This enables a description of a whole complex dataset into a single numpy array, on which one can do operations. In numpy, dimension are also called *axis*. For example, a set of 2 position in space \vec{r}_i can be seen as 2D numpy array, with the first axis being the point $i = 1$ or $i = 2$, and the second axis being the coordinates (x, y, z) . There are few attributes which describe multidimensional arrays:

- `a.dtype`: type of data contained in the array
- `a.shape`: number of elements along each dimension (or axis)
- `a.size`: total number of elements (product of `a.shape` elements)
- `a.ndim`: number of dimensions (or axis)

```
points = np.array([[ 0,  1,  2],
                  [ 3,  4,  5]])

print('a.dtype = {}'.format(points.dtype))
print('a.shape = {}'.format(points.shape))
print('a.size  = {}'.format(points.size))
print('a.ndim  = {}'.format(points.ndim))
```

```
a.dtype = int64
a.shape = (2, 3)
a.size  = 6
a.ndim  = 2
```

2.2 The three key features of NumPy

2.2.1 Vectorization

The *vectorization* is a way to make computations on numpy array **without explicit loops**, which are very slow in python. The idea of vectorization is to compute a given operation *element-wise* while the operation is called on the array itself. An example is given below to compute the inverse of 100000 numbers, both with explicit loop and vectorization.

```
a = np.random.randint(low=1, high=100, size=100000)

def explicit_loop_for_inverse(array):
    res = []
    for a in array:
        res.append(1./a)
    return np.array(res)
```

```
# Using explicit loop
%timeit explicit_loop_for_inverse(a)
```

186 ms ± 13.8 ms per loop (mean ± std. dev. of 7 runs, 10 loops each)

```
# Using list comprehension
%timeit [1./x for x in a]
```

150 ms ± 13.2 ms per loop (mean ± std. dev. of 7 runs, 10 loops each)

```
# Using vectorization
%timeit 1./a
```

106 µs ± 2.65 µs per loop (mean ± std. dev. of 7 runs, 10000 loops each)

The suppression of explicit *for* loops is probably the most unfamiliar aspect of numpy - according to me - and deserves a bit a of practice. At the end, lines of codes becomes relatively short but ones need to properly think how to implement a given computation in a *pythonic way*.

Many standard functions are implemented in a vectorized way, they are call the *universal functions*, or ufunc. Few examples are given below but the full description can be found in [numpy documentation](#).

```
a = np.random.randint(low=1, high=100, size=3)
print('a      : {}'.format(a))
print('a^2    : {}'.format(a**2))
print('a/(1-a^a): {}'.format(a/(1-a**a)))
print('cos(a)  : {}'.format(np.cos(a)))
print('exp(a)  : {}'.format(np.exp(a)))
```

```
a      : [72 29 76]
a^2    : [5184 841 5776]
a/(1-a^a): [ 7.2000000e+01 -5.24473513e-18  7.6000000e+01]
cos(a)  : [-0.96725059 -0.74805753  0.82433133]
exp(a)  : [1.85867175e+31 3.93133430e+12 1.01480039e+33]
```

All these ufunc can work for n-dimension arrays and can be used in a very flexible way depeding on the axis you are refering too. Indeed the mathematical operation can be performed over a different axis of the array, having a totally different meaning. Let's give a simple concrete example with a 2D array of shape (5,2), i.e. 5 vectors of three coordinates (x, y, z) Much more examples will be discussed in the section 2.

```
# Generate 5 vectors (x,y,z)
positions = np.random.randint(low=1, high=100, size=(5, 3))

# Average of the coordinate over the 5 observations
pos_mean = np.mean(positions, axis=0)
```

```

print('mean = {}'.format(pos_mean))

# Distance to the origin sqrt(x^2 + y^2 + z^2) for the 5 observations
distances = np.sqrt(np.sum(positions**2, axis=1))
print('distances = {}'.format(distances))

mean = [50.2 42.6 69.6]
distances = [ 74.16872656  86.68909966 124.83989747  98.04590761 123.59611644]

```

Note on matrix product. Numpy arrays can be used to describe and manipulate matrices. There is a special way to do a matrix product instead of element-wise product. You can use `np.dot(a, b)` (or `a.dot(b)`), even if several other syntaxes are possible (like `a@b`, or equivalently `np.matmul(a, b)`). If you are interested into these features, I would recommand to read in detail the [np.dot documentation](#), because different syntax dont really correspond to the same mathematical operation (for instance, `np.matmul(a, b)` allows *broadcasting* for 2×2 matrix product - cf. latter).

```

a = np.array([[1, 1],
              [1, 0]], dtype=np.int)

b = np.array([[2, 4],
              [1, 1]], dtype=np.int)

print(np.dot(a, b))

[[3 5]
 [2 4]]

```

2.2.2 Broadcasting

The *broadcasting* is a way to compute operation between arrays of having different sizes in a implicit (and concise) manner. One concrete example could be to translate three positions $\vec{r}_i = (x, y)_i$ by a vector \vec{d}_0 simply by adding `points+d0` where `points.shape=(3, 2)` and `d0.shape=(2,)`. Few examples are given below but more details are give in [this documentation](#).

```

# operation between shape (3) and (1)
a = np.array([1, 2, 3])
b = np.array([5])
print('a+b = \n{}'.format(a+b))

a+b =
[6 7 8]

```

```

# operation between shape (3) and (1, 2)
a = np.array([1, 2, 3])
b = np.array([

```

```
[4],
[5],
])
print('a+b = \n{}'.format(a+b))

a+b =
[[5 6 7]
 [6 7 8]]

# Translating 3 2D vectors by d0=(1,4)
points = np.random.normal(size=(3, 2))
d0 = np.array([1, 4])
print('points:\n {}'.format(points))
print('points+d0:\n {}'.format(points+d0))

points:
[[ 0.24333406 -0.80020589]
 [ 1.06124037  0.11580338]
 [ 0.52607555 -1.94077365]]

points+d0:
[[1.24333406 3.19979411]
 [2.06124037 4.11580338]
 [1.52607555 2.05922635]]
```

Not all shapes can be combined together and there are *broadcasting rules*, which are (quoting the [numpy documentation](#)):

When operating on two arrays, NumPy compares their shapes element-wise. It starts with the trailing dimensions, and works its way forward. Two dimensions are compatible when

1. they are equal, or
2. one of them is 1

In a failing case, one can then add a new *empty axis* `np.newaxis` to an array to make their dimension equal and then the broadcasting possible. Here is a very simple example:

```
a = np.arange(10).reshape(2,5)
b = np.array([10,20])

try:
    res = a+b
    print('Possible for {} and {}:'.format(a.shape, b.shape))
    print('a+b = \n{}'.format(res))
except ValueError:
    print('Impossible for {} and {}'.format(a.shape, b.shape))
```

Impossible for (2, 5) and (2,)

```
c = b[:, np.newaxis]
try:
    res = a+c
    print('Possible for {} and {}'.format(a.shape, c.shape))
    print('a+c = \n{}'.format(res))
except ValueError:
    print('Broadcasting for {} and {}'.format(a.shape, c.shape))
```

Possible for (2, 5) and (2, 1):

```
a+c =
[[10 11 12 13 14]
 [25 26 27 28 29]]
```

2.2.3 Working with sub-arrays: slicing, indexing and mask (or selection)

As mentioned earlier, *slicing and indexing* are ways to access elements or sub-arrays in a smart way. Python allows slicing with `Slice()` object but numpy allows to push the logic much further with what is called *fancy indexing*. Few examples are given below and for more details, please have a look to [this documentation page](#).

Rule 1: the syntax is `a[i]` to access the *i*th element. It is also possible to go from the last element using negative indices: `a[-1]` is the last element.

```
a = np.random.randint(low=1, high=100, size=10)
print('a = {}'.format(a))
print('a[2] = {}'.format(a[2]))
print('a[-1] = {}'.format(a[-1]))
print('a[[1, 2, 5]] = {}'.format(a[[1, 2, 5]]))

a = [26 15 60 38 79 78 12 27 86 14]
a[2] = 60
a[-1] = 14
a[[1, 2, 5]] = [15 60 78]
```

Rule 2: numpy also support array of indices. If the index array is multi-dimensional, the returned array will have the same dimension as the indices array.

```
# Small n-dimensional indices array: 3 arrays of 2 elements
indices = np.arange(6).reshape(3,2)
print('indices =\n{}'.format(indices))
print('a[indices] =\n{}'.format(a[indices]))
```

```
indices =
[[0 1]
 [2 3]]
```

```
[4 5]
a[indices] =
[[26 15]
[60 38]
[79 78]]

# Playing with n-dimensional indices array: 2 arrays of (10, 10) arrays
indices_big = np.random.randint(low=0, high=10, size=(2, 3, 2))
print('indices_big =\n{}'.format(indices_big))
print('a[indices_big] =\n{}'.format(a[indices_big]))
```



```
indices_big =
[[[3 1]
[4 2]
[1 7]]

[[4 6]
[7 6]
[0 4]]]

a[indices_big] =
[[[38 15]
[79 60]
[15 27]

[[79 12]
[27 12]
[26 79]]]
```

Rule 3: There is a smart way to access sub-arrays with the syntax `a[min:max:step]`. In that way, it's for example very easy to take one element over two (`step=2`), or reverse the order of an array (`step=-1`). This syntax works also for n-dimensional array, where each dimension is sperated by a comma. An example is given for a 1D array and for a 3D array of shape (5, 2, 3) - that can considered as 5 observations of 2 positions in space.

```
# 1D array
a = np.random.randint(low=1, high=100, size=10)
print('full array a          = {}'.format(a))
print('from 0 to 1: a[:2]     = {}'.format(a[:2]))
print('from 4 to end: a[4:]   = {}'.format(a[4:]))
print('reverse order: a[::-1] = {}'.format(a[::-1]))
print('all even elements: a[::2] = {}'.format(a[::2]))
```



```
full array a          = [10 31 61 38  1 17 59 24  3 61]
from 0 to 1: a[:2]     = [10 31]
from 4 to end: a[4:]   = [ 1 17 59 24  3 61]
reverse order: a[::-1] = [61  3 24 59 17  1 38 61 31 10]
all even elements: a[::2] = [10 61  1 59  3]
```

```
# 3D array
a = np.random.randint(low=0, high=100, size=(5, 2, 3))
print('a = \n{}'.format(a))
```

```
a =
[[[20  0  7]
 [93 24  4]

 [[43 65 46]
 [97 59 57]

 [[86 2 85]
 [31 0 57]

 [[46 1 94]
 [98 65  6]

 [[81 15  3]
 [88 47 90]]]
```

Let's say, one wants to take only the (x,y) coordinates for the first vector for all 5 observations. This is how each axis will be sliced: - first axis (=5 observations): `:`, i.e. takes all - second axis (=2 vectors): `1` i.e. only the 2nd element - third axis (=3 coordinates): `0:2` i.e. from 0 to 2 – 1 = 1, so only (x,y)

```
# Taking only the x,y values of the first vector for all observation:
print('a[:, 0, 0:2] =\n{}'.format(a[:, 0, 0:2]))
```

```
a[:, 0, 0:2] =
[[20  0]
 [43 65]
 [86 2]
 [46 1]
 [81 15]]
```

```
# Reverse the order of the 2 vector for each observation:
print('a[:, ::-1, :] =\n{}'.format(a[:, ::-1, :]))
```

```
a[:, ::-1, :] =
[[[93 24  4]
 [20 0  7]

 [[97 59 57]
 [43 65 46]

 [[31 0 57]
 [86 2 85]]]
```

```
[[98 65 6]
 [46 1 94]]
```

```
[[88 47 90]
 [81 15 3]]]
```

Rule 4: The last part of indexing is about *masking* array or in a more common language, *selecting* sub-arrays/elements. This allows to get only elements satisfying a given criteria, exploiting the indexing rules described above. Indeed, a boolean operation applied to an array such as `a>0` will directly return an array of boolean values True or False depending if the corresponding element satisfies the condition or not.

```
a = np.random.randint(low=-100, high=100, size=(5, 3))
mask = a>0
print('a = \n{}'.format(a))
print('\nmask = \n{}'.format(mask))
```

```
a =
[[ 19 -52  77]
 [ 36 -68  88]
 [-61 -18 -71]
 [ 64 -46  58]
 [-83 -12  18]]
```

```
mask =
[[ True False  True]
 [ True False  True]
 [False False False]
 [ True False  True]
 [False False  True]]
```

```
print('\na[mask] = \n{}'.format(a[mask])) # always return 1D array
print('\na*mask = \n{}'.format(a*mask)) # preserves the dimension (False=0)
print('\na[~mask] = \n{}'.format(a[~mask])) # ~mask is the negation of mask
print('\na*~mask = \n{}'.format(a*~mask)) # working for a product too.
```

```
a[mask] =
[19 77 36 88 64 58 18]
```

```
a*mask =
[[19  0 77]
 [36  0 88]
 [ 0  0  0]
 [64  0 58]
 [ 0  0 18]]
```

```
a[~mask] =
[-52 -68 -61 -18 -71 -46 -83 -12]
```

```
a*~mask =
[[ 0 -52   0]
 [ 0 -68   0]
 [-61 -18 -71]
 [ 0 -46   0]
 [-83 -12   0]]
```

Note the case of boolean arrays as indices has then a special treatment in numpy (since the result is always a 1D array). There is actually a dedicated numpy object called *masked array* (cf. [documentation](#)) which allows to keep the whole array but without considering some elements in the computation (e.g. CCD camera with dead pixel). Note however that when a boolean array is used in an mathematical operation (such as `a*mask`) then `False` is treated as 0 and `True` as 1:

```
print('a+mask = \n{}'.format(a+mask))
```

```
a+mask =
[[ 20 -52  78]
 [ 37 -68  89]
 [-61 -18 -71]
 [ 65 -46  59]
 [-83 -12  19]]
```

This boolean arrays are also very useful to *replace a category of elements* with a given value in a very easy, concise and readable way:

```
a = np.random.randint(low=-100, high=100, size=(5, 3))
print('Before: a=\n{}'.format(a))

a[a<0] = a[a<0]**2
print('\nAfter: a=\n{}'.format(a))
```

```
Before: a=
[[ 42  81  68]
 [ -5  89 -83]
 [-23  77  93]
 [ 78  18 -29]
 [-99 -38  54]]
```

```
After: a=
[[ 42  81  68]
 [ 25  89 6889]
 [ 529  77  93]
 [ 78  18 841]
 [9801 1444  54]]
```

2.3 Few useful NumPy tips

This short section is presenting few handy features to know about NumPy, which can help beginners. For a slightly more complete view of “everyday NumPy”, I would recommend to have a look to the [cheat sheet](#) from DataCamp.

Dummy array initialization.

```
x = np.zeros(shape=(3, 2))           # Only 0
x = np.ones(shape=(3, 2))            # Only 1
x = np.full(shape=(3, 2), fill_value=10) # Only 10
x = np.eye(2)                      # Create identity matrix (only return 2D array)
```

Create sequence of numbers.

```
# Linear interval from 0 to 10
x = np.linspace(0, 10, 10) # 10 numbers between 0 and 1
x = np.arange(0, 10, 1.0) # One number every 1.0 between 0 and 1
x = np.logspace(0, 10, 10) # 10 numbers between 10**0 and 10**10
```

Shape-based manipulation of arrays.

```
a = np.arange(0, 18).reshape(3, 3, 2)
x = a.ravel()                      # Return a flat array
x = a.reshape(9, 2)                 # change the shape
x = a.T                            # transpose array: a.T[i, j, k] = a[k, j, i]
x = np.concatenate([a,a], axis=0)   # concatenate arrays along a given axis (shape is (6, 3,
                                  # 2))
x = np.stack([a, a], axis=0)        # group arrays along a given axis (shape is (2, 3, 3, 2))
```

Compare arrays.

```
# Making dummy arrays for comparisons
a = np.arange(-6, 6).reshape(3, 4)
b = np.abs(a)
c = np.append(b, [[1, 2, 3, 4]], axis=0)

# Print the arrays
print('a = {}'.format(a))
print('b = {}'.format(b))
print('c = {}'.format(c))
```

```
a = [[-6 -5 -4 -3]
      [-2 -1  0   1]
      [ 2   3   4   5]]
```

```
b = [[6 5 4 3]
```

```
[2 1 0 1]
[2 3 4 5]]
```

```
c = [[6 5 4 3]
 [2 1 0 1]
 [2 3 4 5]
 [1 2 3 4]]
```

```
# Arrays with the same shapes
print(np.equal(a, b))           # Return array with element-wise True/False
print(np.all(a==b))             # Return true if all element is true
print(np.allclose(a, b, rtol=10)) # same as all function with relative/absolute precision
print(np.any(a==b))             # Return true if any of the element is true
```

```
[[False False False False]
 [False False  True  True]
 [ True  True  True  True]]
False
True
True
```

```
# Arrays with the possibly different shapes
print(np.array_equal(a, c)) # True if a and b have the same shape and np.equal(a, b)
print(np.array_equiv(a, b)) # True if a and b have broadcastable shapes and same elements
```

```
False
False
```

```
# Example of equivalent arrays
a = np.array([1, 2])
b = np.array([[1, 2], [1, 2]])
np.array_equiv(a, b)
```

```
True
```

2.4 Example of simple gradient descent: NumPy v.s. pure python

2.4.1 Gradient descent: what (for) is this?

The gradient descent is a method allowing to numerically find a the minum of a function $f(p_0, \dots, p_n)$. Finding minumum is needed for most of machine learning problems (I include usual model fitting in the machine learning category here): ones always want to find the *best set of parameter describing a dataset, assuming a given function*. Let's assume, you have n couple of measured values (x_i, y_i) and you want to be able to predict the mathematical relationship between x and y for all points: $y_i = \text{model}(x_i)$. Usually the mathematical

function “model” will depend on some unknown parameters p_0, \dots, p_N . In that case, the function to minimize is often the *error* function (or *cost*):

$$f(p_0, \dots, p_N) = \frac{1}{n} \sum_{i=1}^n (y_i - \text{model}(x_i; p_0, \dots, p_N))^2$$

Finding the minimum of an error (cost) function is rather general to any (supervised) learning algorithm. These notions will be described in more details in other lectures.

How does gradient descent work? At each iteration (or *epoch*), parameters are updated using a step value μ along the oposit direction of the gradient, evaluated at the present point:

$$(p_0, \dots, p_N)^{i+1} \leftarrow (p_0, \dots, p_N)^i - \mu \left(\frac{\partial f}{\partial p_0}, \dots, \frac{\partial f}{\partial p_N} \right) |_{(p_0, \dots, p_N)^i}$$

This assume that the value of the gradient is known. There are some technics to numerically estimate the gradient for arbitrary function. In the example below, we consider a much simpler situation - which actually has an exact solution: a linear model. In other words, there are only two parameters and we assume that:

$$\text{model}(x) = p_0 + p_1 x$$

with the following loss function gradient:

$$\frac{\partial f}{\partial p_0} = -\frac{2}{n} \sum_{i=0}^{i=n} (y_i - p_0 - p_1 x_i) \quad (2.1)$$

$$\frac{\partial f}{\partial p_1} = -\frac{2}{n} \sum_{i=0}^{i=n} ((y_i - p_0 - p_1 x_i) \times x_i) \quad (2.2)$$

From a coding point of view, we will then introduce an array `delta = [yi - p0 - p1*xi]` which will be used to compute both the two gradient components and the loss function. The next two section describe the pure python implementation and a numpy implementation, in order to compare performances. What follows is higly inspired from a [RealPython post](#) on performance comparison. Before entering in the discussion, let's define our fake dataset:

```
# Fake (xi, yi) data definition
n = 1000
x = np.linspace(0, 2, n)
xfine = np.linspace(0, 2, 1000) # to draw a line
y = 3 + 2 * x + 0.1*np.random.randn(n)

# Linear model definition
def model(x, p0, p1):
    return p0 + p1*x

# Loss function definition
def loss_function(p0, p1):
    return np.mean((y - model(x, p0, p1))**2)
```

```
# Vectorize the loss function for many parameters
loss_function = np.vectorize(loss_function)
```

The above `numpy.vectorize()` function allow to make several calls of the same function much faster using vectorization (cf. [this documentation page](#)). The next function `plot_model(p)` follows simply represent the data, the model, the loss function evolution and the trajectory in the (p_0, p_1) space which is followed by the gradient descent. This function is based in `matplotlib` librairy which will be discussed in later chapters of this lecture.

```
# Plotting function (data vs fit, loss function, gradient descent)
def plot_model(p):
    """
    Producing three plots from the list of the 2
    parameters for all epochs: p.shape = (Nepochs, 2)
    """

    import matplotlib.pyplot as plt
    plt.figure(figsize=(30, 7))

    # Get Best parameters (last ones), ymodel and lost functions
    p0, p1 = p[-1, 0], p[-1, 1]
    ymodel = model(x, p0, p1)
    loss = loss_function(p[:, 0], p[:, 1])

    # Plot (xi,yi) data and overlay (x, model(x, p)) points
    plt.subplot(1, 3, 1)
    plt.plot(x, y, 'o', alpha=0.3, markersize=5, label='data')
    plt.plot(xfine, ymodel, linewidth=3, color='tab:red', label='model')
    plt.xlabel('$x$'); plt.ylabel('$y$')
    plt.legend()

    # Plot the loss function v.s. epoch
    plt.subplot(1, 3, 2)
    plt.semilogy(loss, linewidth=3)
    plt.xlabel('Epochs'); plt.ylabel('Loss function')

    # Plot the gradient in the (p0, p1) space and the descent trajectory
    plt.subplot(1, 3, 3)
    P0, P1 = np.meshgrid(np.linspace(0, 4, 300), np.linspace(0, 3, 300))
    plt.imshow(np.log(loss_function(P0, P1)), extent=[0, 4, 0, 3], aspect='auto',
               origin='lower', cmap='Greys')
    plt.plot(p[:, 0], p[:, 1], linewidth=5, color='tab:red', alpha=0.8, label='Trajectory')
    plt.xlabel('$p_0$'); plt.ylabel('$p_1$')
    plt.colorbar(label='log(loss function)')
    for t in plt.legend().get_texts():
        t.set_color('white')

    return
```

2.4.2 Pure python implementation

The pure python implementation don't use any of the vectorized features of numpy. The loop and sum over the data points are explicit, using `zip()`, `sum()` and comprehension syntax. The function return an array of all parameters for each epoch (for later convenience, we simply return a numpy array - but mathematical operations are not done with numpy in this function).

```
def python_linear_descent(x, y, mu, N_epochs):

    # Length of data
    n = len(x)

    # Initialize predictions, errors, parameters and gradients.
    ym = [0] * n
    para = [[0, 0]] * N_epochs
    grad = [0, 0]
    pm = [0, 0]

    # Looping over iterations (epochs)
    for i_epoch in range(0, N_epochs):
        delta = tuple(i - j for i, j in zip(y, ym))
        grad[0] = -2/n * sum(delta)
        grad[1] = -2/n * sum(i * j for i, j in zip(delta, x))
        pm = [i - mu * j for i, j in zip(pm, grad)]
        ym = (model(i, pm[0], pm[1]) for i in x)

        # Save all parameters
        para[i_epoch] = pm

    # Return numpy array of the 2 paramters for all epochs
    return np.array(para)
```

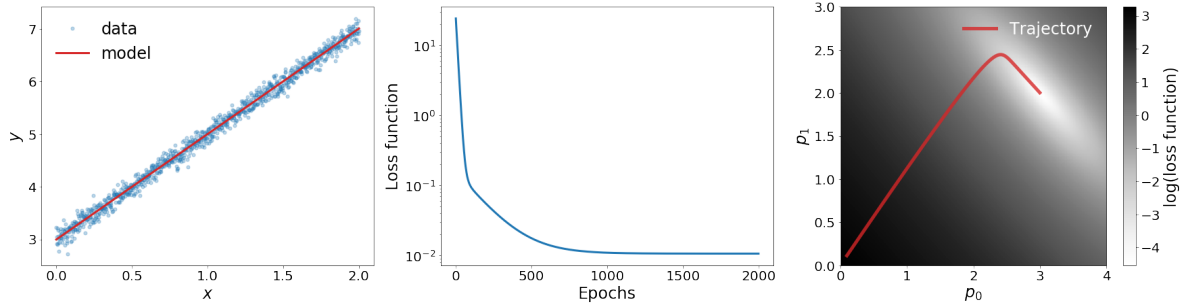
We can then try to time this function using a step of 0.01 and 2000 epochs for our 1000 data points.

```
%timeit python_linear_descent(x, y, mu=0.01, N_epochs=2000)
```

```
1.12 s ± 34.8 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)
```

This takes approximatly 1 second to run. What follows shows the best model prediction for the best parameters, the evolution of the loss function as well as the descent trajectory in the parameter space:

```
parameters = python_linear_descent(x, y, mu=0.01, N_epochs=2000)
plot_model(parameters)
```



2.4.3 Numpy implementation

The numpy implementation makes a full use of the vectorization feature discussed several time in this chapter, but also broadcasting. This lead to a significantly clearer piece of code and also much faster. We can note in particular the different syntax used to update the model parameters.

```
def numpy_linear_descent(x, y, mu, N_epochs):

    # To define the lost function
    n = x.shape[0]

    # Initialize predictions, errors, parameters and gradients.
    ym = np.zeros(n)
    para = np.zeros((N_epochs, 2))
    pm, grad = np.zeros(2), np.zeros(2)

    # Looping over iterations (epochs)
    for i_epoch in range(0, N_epochs):
        delta = y - ym
        grad = -2/n * np.array([np.sum(delta), np.sum(delta*x)])
        pm = pm - mu*grad
        ym = model(x, pm[0], pm[1])

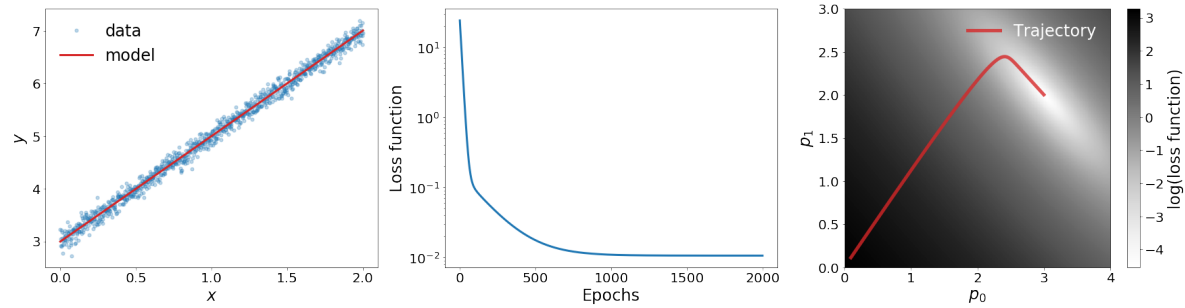
        # Save all parameters
        para[i_epoch] = pm

    return para

%timeit numpy_linear_descent(x, y, mu=0.01, N_epochs=2000)
```

37.3 ms ± 3.47 ms per loop (mean ± std. dev. of 7 runs, 10 loops each)

```
parameters = numpy_linear_descent(x, y, mu=0.01, N_epochs=2000)
plot_model(parameters)
```



We check that the result are indeed the same for an execution about 25 times faster ...

Chapter 3

Three important tools to know

3.1 matplotlib: graphical representation of data

Matplotlib is an extremely rich library for data visualization and there is no way to cover all its features in this note. The goal of this section is just to give short and practical examples to plot data. Much more details can be obtained on the [webpage](#). The following shows how to quickly make *histograms*, *graph*, *2D and 3D scatter plots*.

The main object of matplotlib is `matplotlib.pyplot` imported as `plt` here (and usually). The most common functions are then called on this objects, and often takes numpy arrays in argument (possibly with more than one dimension) and a lot of `kwargs` to define the plotting style.

```
import numpy as np
import matplotlib.pyplot as plt
%matplotlib inline
```

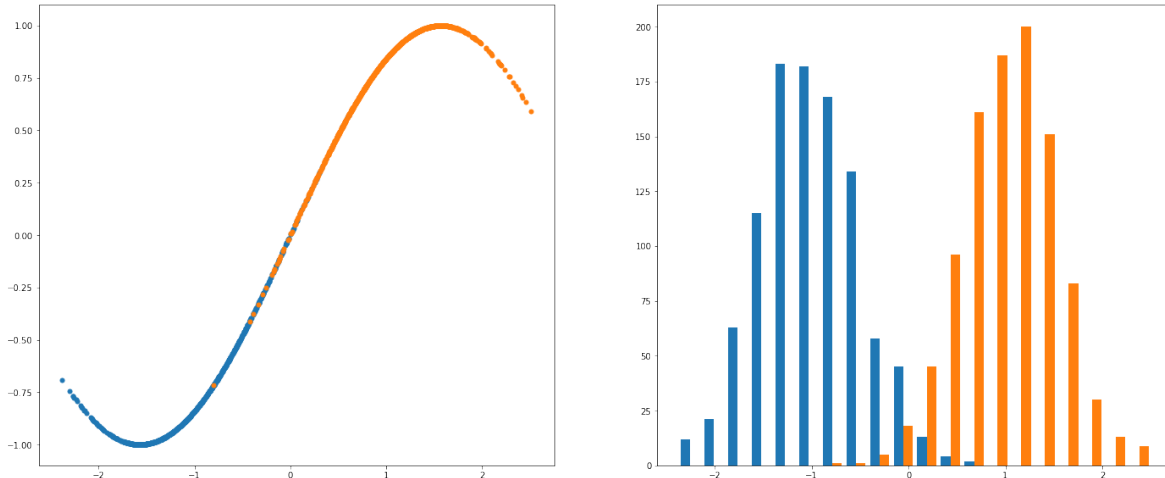
3.1.1 Example of 1D plots and histograms

To play with data, we generate 2 samples of 1000 values distributed according to a normal probability density function with $\mu = -1$ and $\mu = 1$ respectively, and $\sigma = 0.5$. These data are stored in a numpy array `x` of shape `x.shape=(1000, 2)`. We then simply compute and store the sinus of all these values into a same shape array `y`:

```
x = np.random.normal(loc=[-1, 1], scale=[0.5, 0.5], size=(1000,2))
y = np.sin(x)
```

The next step is to plot these data in two ways: first we want `y` v.s. `x`, second we want the histogram of the `x` values. We need to first create a figure, then create two *subplots* (specifying the number of line, column, and subplot index). Note that matplotlib take always the first dimension to define the numbers to plot, while higher dimensions are considered as other plots - automatically overlaid.

```
plt.figure(figsize=(24, 10))
plt.subplot(121) # 121 means 1 line, 2 column, 1st plot
plt.plot(x, y, marker='o', markersize=5, linewidth=0.0)
plt.subplot(122) # 122 means 1 line, 2 column, 2nd plot
plt.hist(x, bins=20);
```



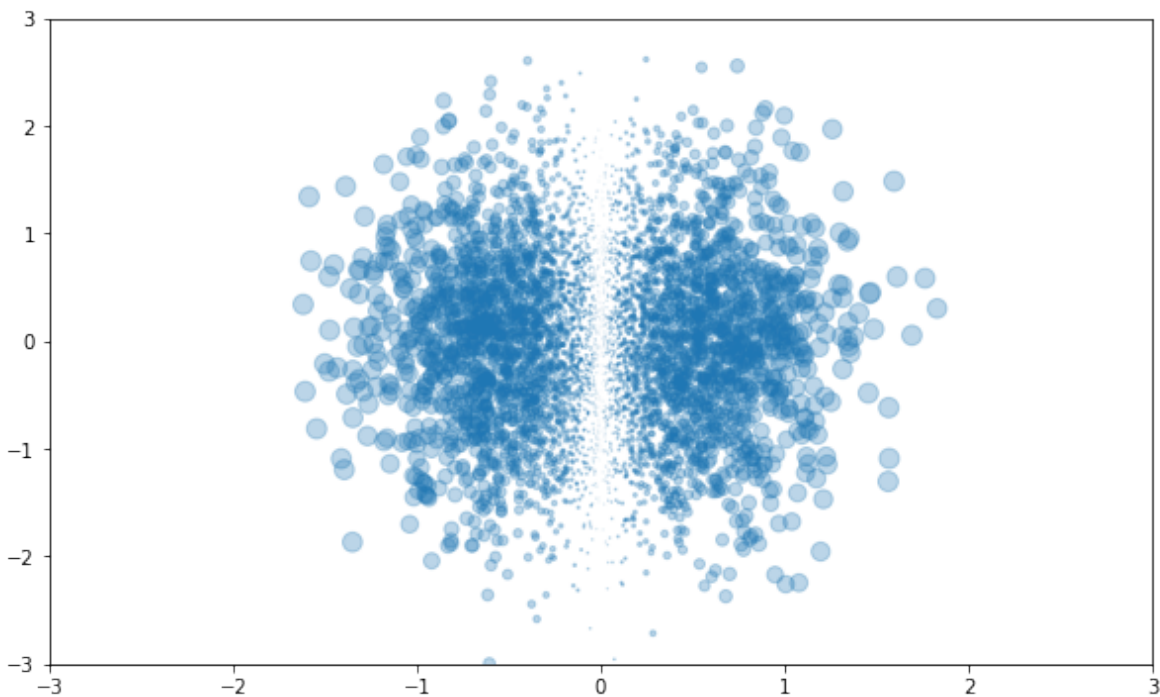
3.1.2 Example of 2D scatter plot

A scatter plot allows to draw marker in a 2D space and a third information is encoded into the marker size. In order to play, we generated two set of 5000 numbers distributed according to uncorrelated gaussians of $(\mu_0 = \mu_1 = 0)$ and $(\sigma_1, \sigma_2) = (0.5, 0.8)$ in a numpy array `points` of shape `points.shape=(5000, 2)`. These two sets of numbers are then interpreted as (x, y) positions being loaded in two $(5000, 1)$ arrays `x` and `y`:

```
points = np.random.normal(loc=[0, 0], scale=[0.5, 0.8], size=(5000,2))
x, y = points[:, 0], points[:, 1]
```

We can then plot the 5000 points in the 2D plan, and here we specify the marker size at $100 \times \sin^2(x)$ using the argument `s` of the `plt.scatter()` function (note that the array `x`, `y` and `s` must have the same shape):

```
plt.figure(figsize=(10,6))
plt.scatter(x, y, s=100*(np.sin(x))**2, marker='o', alpha=0.3)
plt.xlim(-3, 3)
plt.ylim(-3, 3);
```



3.1.3 Example of 3D plots

For 3D plots, one can generate 1000 positions in space, and operate a translation by a vector \vec{r}_0 using broadcasting:

```
data = np.random.normal(size=(1000, 3))
r0 = np.array([1, 4, 2])
data_trans = data + r0
```

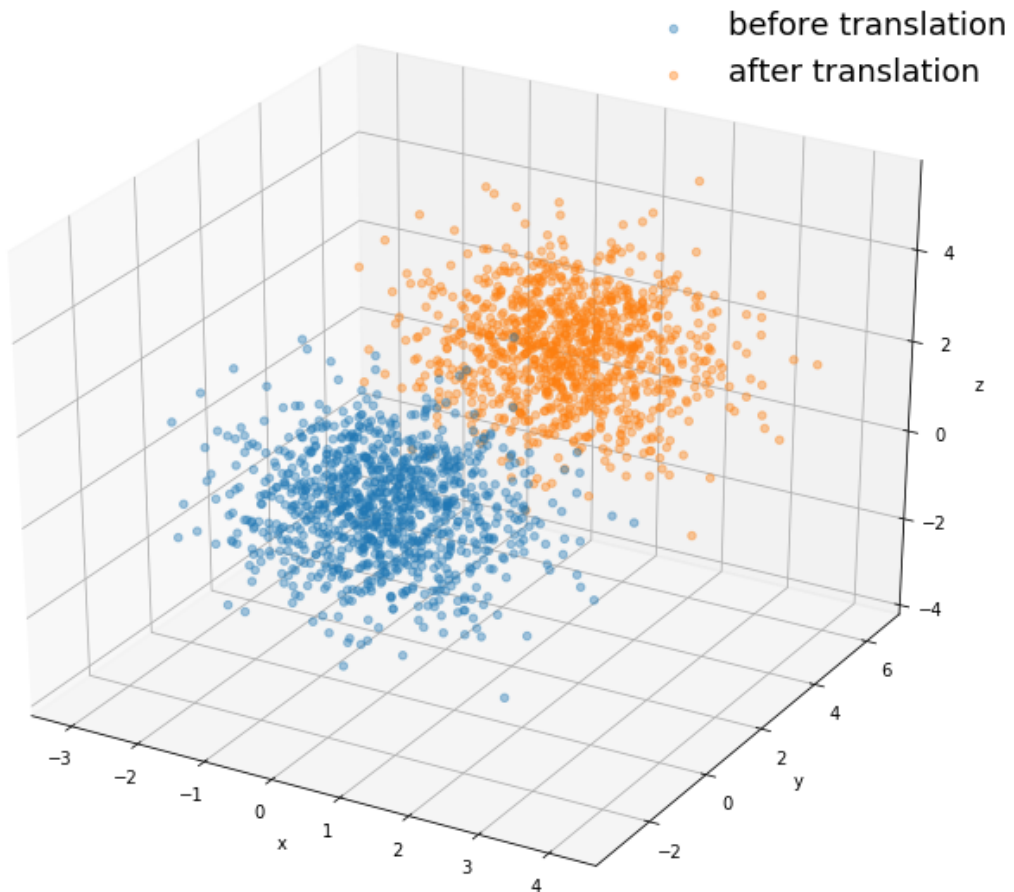
It is then easy to get back the spatial initial (*i.e.* before translation) and final (*i.e.* after translation) coordinates:

```
xi, yi, zi = data[:,0], data[:,1], data[:,2]
xf, yf, zf = data_trans[:,0], data_trans[:,1], data_trans[:,2]
```

An additional module must be imported in order to plot data in three dimensions, and the projection has to be stated. Once it's done, a simple call to `ax.scatter3D(x, y, z)` does the plot. Note that we call a function of `ax` and not `plt` as before. This is due to the `ax = plt.axes(projection='3d')` command which is needed for 3D plotting. More details are available on the [matplotlib 3D tutorial](#).

```
from mpl_toolkits import mplot3d
plt.figure(figsize=(12,10))
ax = plt.axes(projection='3d')
ax.scatter3D(xi, yi, zi, alpha=0.4, label='before translation')
ax.scatter3D(xf, yf, zf, alpha=0.4, label='after translation')
ax.set_xlabel('x')
```

```
ax.set_ylabel('y')
ax.set_zlabel('z')
ax.legend(frameon=False, fontsize=18);
```



3.1.4 Example of 2D function $z = f(x, y)$: notion of meshgrid

Another typical plot we might want to do is to represent a function of two variables (x, y) in 3D: $z = f(x, y)$. In python this implies the notion of *meshgrid* which is not trivial at first. Let's first define a 2 variable function:

```
def my_surface(x, y):
    x0 = 5*np.sin(y)
    sigma = 5+y
    amp = (10-y)
    return amp*np.exp(-(x-x0)**2/sigma**2)
```

Let's define a (x, y) interval on which we want to describe the surface:

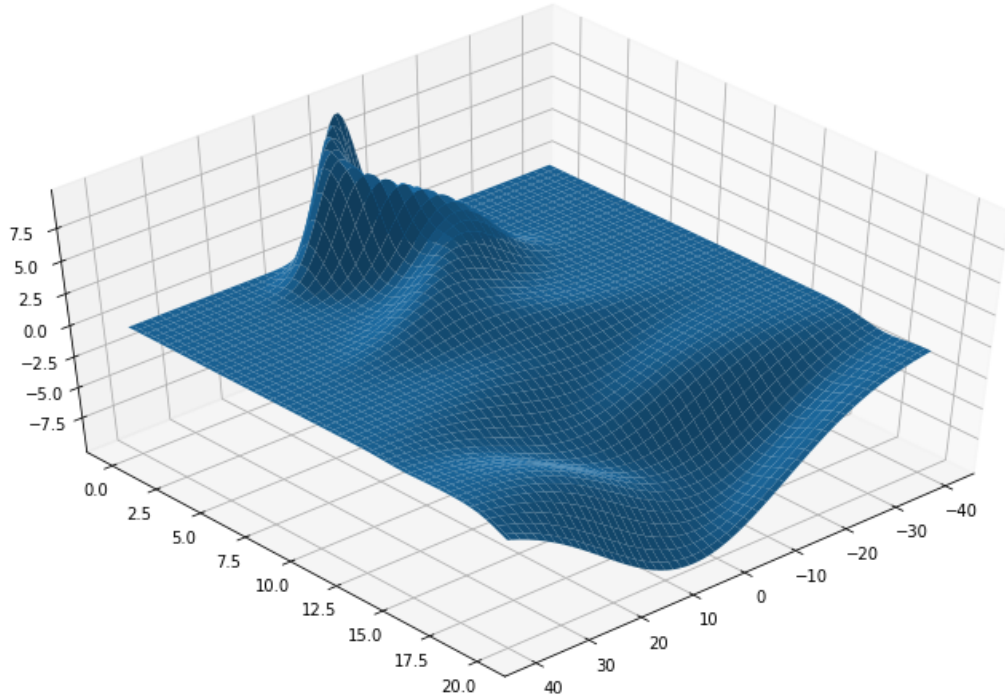
```
x = np.linspace(-40, 40, 100)
y = np.linspace(0, 20, 200)
```

These two numpy arrays don't have the same shape and an explicit loop would be needed to process them - which is very time consuming in python. This is where the *meshgrid* notion comes: it will provide a two arrays *with the same size* and allow then the vectorization:

```
# Meshgrid and function application (see after for more details)
xx, yy = np.meshgrid(x, y)
Z = my_surface(xx, yy)

# Plotting
fig = plt.figure(figsize=(13,8))
ax = fig.gca(projection='3d')
ax.plot_surface(xx, yy, Z)

# Choose the default view
ax.view_init(azim=48, elev=48)
```



Meshgrid explanation. The *meshgrid* consists of two 2D arrays made out of two 1D arrays. The main purpose is to have numpy arrays with the same shape, which can then support vectorized operations. The logic is relatively straightforward and can be understood with two points along each coordinate. Let's assume you want to scan the $\{0, 1\}$ x-values and $\{2, 3\}$ y-values, then you need to build up the four following 2D points: $(0, 2)$, $(1, 2)$, $(0, 3)$, $(1, 3)$. These four points can be encoded in the two arrays $[[0, 1], [0, 1]]$ and $[[2, 2], [3, 3]]$. These two arrays have the same shape, which is similar to the result of $f(x, y)$ computed on this grid, even if there are *not the same numbers of x and y values*.

```
# Create a simple 2-variables function
def f(x, y):
    return x**2+y**2

# Define x-values, y-values and create the 2D
x, y = np.arange(0, 2), np.arange(2, 4)
xx, yy = np.meshgrid(x, y)
zz = f(xx, yy)

# Printing arrays
print('Array values:')
print('xx={}'.format(xx))
print('yy={}'.format(yy))
print('zz={}'.format(zz))
```

```
Array values:
xx=[[0 1]
 [0 1]]
yy=[[2 2]
 [3 3]]
zz=[[ 4  5]
 [ 9 10]]
```

A way to explicit the meshgrid, you can flat xx and yy arrays with the `ravel()` function, and take each pair (with the `zip()` syntax). Every point is indeed formed:

```
for i, j in zip(xx.ravel(), yy.ravel()):
    print('({},{}) = {} ; f({},{}) = {}'.format(i, j, f(i, j)))
```

```
(x,y)=(0,2); f(x,y)=4
(x,y)=(1,2); f(x,y)=5
(x,y)=(0,3); f(x,y)=9
(x,y)=(1,3); f(x,y)=10
```

If you want to read more about this, you can check the [numpy meshgrid documentation](#) and this [stackoverflow post](#). For more advanced readers, there are two similar functions which return slightly different objects: `np.ogrid()` and `np.mgrid()`. For a nice discussion of differences, you can check [this post](#).

3.2 pandas: import and manipulate data as numpy array

The package `pandas` is an very rich interface to read data from different format and produce a `pandas.DataFrame` that can be based on `numpy` (but containing a lot more features). There is no way to fully describe this package here, the goal is simply to give functional and concrete example easily usable. More details, please check the [pandas webpage](#).

3.2.1 Data importation

Many build-in functions are available to import data as pandas dataframe. One, which is particularly convenient, directly reads csv files (one can specify the columns to loads, the row to skip, and many other options ...):

```
import pandas as pd
df = pd.read_csv('../data/WaveData.csv')
print(df.head())
```

	Date/Time	Hs	Hmax	Tz	Tp	Peak Direction	SST
0	01/01/2017 00:00	-99.900	-99.90	-99.900	-99.900		-99.9 -99.90
1	01/01/2017 00:30	0.875	1.39	4.421	4.506		-99.9 -99.90
2	01/01/2017 01:00	0.763	1.15	4.520	5.513		49.0 25.65
3	01/01/2017 01:30	0.770	1.41	4.582	5.647		75.0 25.50
4	01/01/2017 02:00	0.747	1.16	4.515	5.083		91.0 25.45

```
# Rename columns names using df.rename() function
old_new_cols = {
    'Date/Time': 'date',
    'Hs': 'height',
    'Hmax': 'heightMax',
    'Tz': 'period',
    'Tp': 'energy',
    'Peak Direction': 'direction',
    'SST': 'temperature'
}

# The argument `inplace` means the current dataframe is overwritten with the change
df.rename(columns=old_new_cols, inplace=True)
print(df.head())
```

	date	height	heightMax	period	energy	direction	\
2	01/01/2017 01:00	0.763	1.15	4.520	5.513	49.0	
3	01/01/2017 01:30	0.770	1.41	4.582	5.647	75.0	
4	01/01/2017 02:00	0.747	1.16	4.515	5.083	91.0	
5	01/01/2017 02:30	0.718	1.61	4.614	6.181	68.0	
6	01/01/2017 03:00	0.707	1.34	4.568	4.705	73.0	

	temperature	height_normalized	heightMax_normalized	period_normalized	\
2	25.65	-0.898215	-1.047342	-1.184338	
3	25.50	-0.884973	-0.757690	-1.117565	
4	25.45	-0.928484	-1.036201	-1.189723	
5	25.45	-0.983346	-0.534881	-1.083102	
6	25.50	-1.004155	-0.835673	-1.132643	

	energy_normalized	direction_normalized	temperature_normalized	
2	-1.463956	-2.044360	0.762152	

```

3      -1.407891      -0.973294      0.694918
4      -1.643866      -0.314176      0.672506
5      -1.184467      -1.261658      0.672506
6      -1.802020      -1.055683      0.694918

```

3.2.2 Cleaning the dataset using numpy syntax

This is possible to clean the dataframe using some masking syntax. First, let's check how many default values are stored for each column (all but the date):

```

# Check which wave has -99 values for every variables
for c in ['height', 'heightMax', 'period', 'energy', 'direction', 'temperature']:
    n = np.count_nonzero(df[c]<=-99)
    print('{}: {} wave have <=-99'.format(c, n))

```

```

height: 85 wave have <=-99
heightMax: 85 wave have <=-99
period: 85 wave have <=-99
energy: 85 wave have <=-99
direction: 271 wave have <=-99
temperature: 262 wave have <=-99

```

```

# Simply take value above -99
print(df[df>-99].head())

```

	date	height	heightMax	period	energy	direction	temperature
0	01/01/2017 00:00	NaN	NaN	NaN	NaN	NaN	NaN
1	01/01/2017 00:30	0.875	1.39	4.421	4.506	NaN	NaN
2	01/01/2017 01:00	0.763	1.15	4.520	5.513	49.0	25.65
3	01/01/2017 01:30	0.770	1.41	4.582	5.647	75.0	25.50
4	01/01/2017 02:00	0.747	1.16	4.515	5.083	91.0	25.45

```

# Removing all entry (line) which has at least one default value
for c in ['height', 'heightMax', 'period', 'energy', 'direction', 'temperature']:
    df = df[df[c]>-99]

print(df.head())

```

	date	height	heightMax	period	energy	direction	temperature
2	01/01/2017 01:00	0.763	1.15	4.520	5.513	49.0	25.65
3	01/01/2017 01:30	0.770	1.41	4.582	5.647	75.0	25.50
4	01/01/2017 02:00	0.747	1.16	4.515	5.083	91.0	25.45
5	01/01/2017 02:30	0.718	1.61	4.614	6.181	68.0	25.45
6	01/01/2017 03:00	0.707	1.34	4.568	4.705	73.0	25.50

We can now check how many default value we get:

```
for c in ['height', 'heightMax', 'period', 'energy', 'direction', 'temperature']:
    n = np.count_nonzero(df[c]<=-99)
    print('{}: {} wave have <=-99'.format(c, n))
```

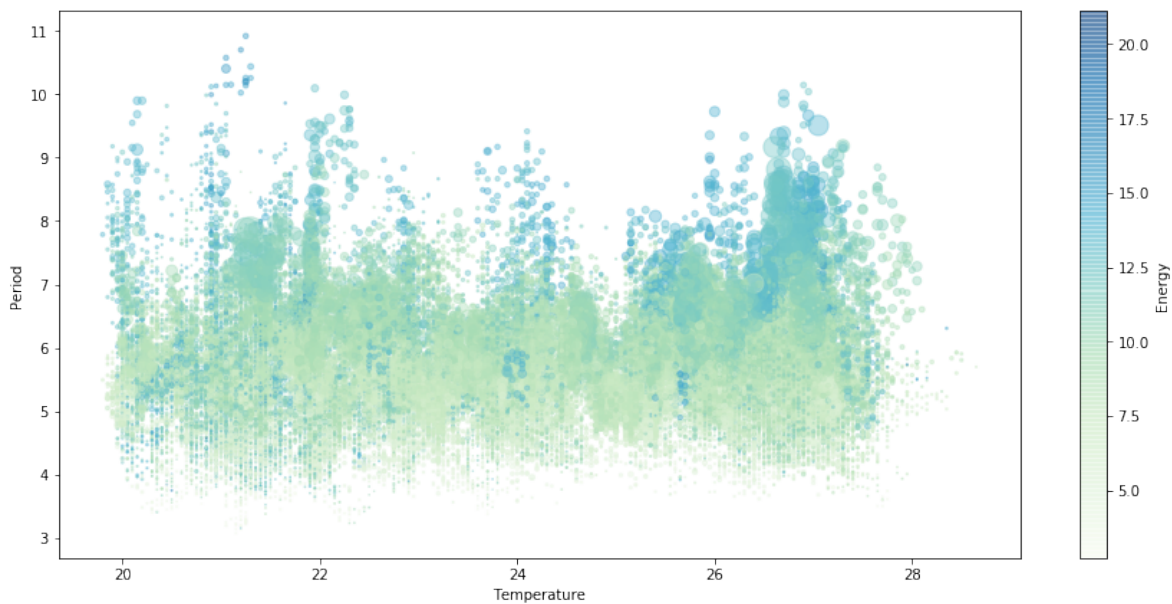
```
height: 0 wave have <=-99
heightMax: 0 wave have <=-99
period: 0 wave have <=-99
energy: 0 wave have <=-99
direction: 0 wave have <=-99
temperature: 0 wave have <=-99
```

3.2.3 Extracting numpy arrays and plotting

This is possible to perform some computation using pandas columns directly, but it can be useful to extract numpy arrays in case of more complex broadcasting or indexing. This can be done using `df[col].values` command:

```
# Get numpy array for further manipulations
T = df['temperature'].values
P = df['period'].values
H = df['heightMax'].values
E = df['energy'].values
```

```
# Plot temperature vs period vs max_height vs energy
plt.figure(figsize=(15, 7))
plt.scatter(T, P, s=H**3, c=E, cmap='GnBu', alpha=0.4)
plt.colorbar(label='Energy')
plt.xlabel('Temperature')
plt.ylabel('Period');
```



3.2.4 Add information in a dataframe

One of the nice feature of pandas is to be able to easily store the result of a computation as a new column. For instance, it's a common practice in machine learning to *normalize* the input variables, *i.e.* transform them to have a mean of 0 and a variance of 1.0. The following example shows how to add new column which are normalized:

```
def add_normalized_variable_to_df(col):

    # Get a numpy arrays
    v = df[col].values

    # Replace NaN by 0.0
    v[np.isnan(v)] = 0

    # Compute quantities
    v_mean = np.mean(v)
    v_rms = np.sqrt(np.mean((v-v_mean)**2))

    # Add them into the pandas dataframe
    df[col+'_normalized'] = (v-v_mean)/v_rms

    return

for c in ['height', 'heightMax', 'period', 'energy', 'direction', 'temperature']:
    add_normalized_variable_to_df(c)

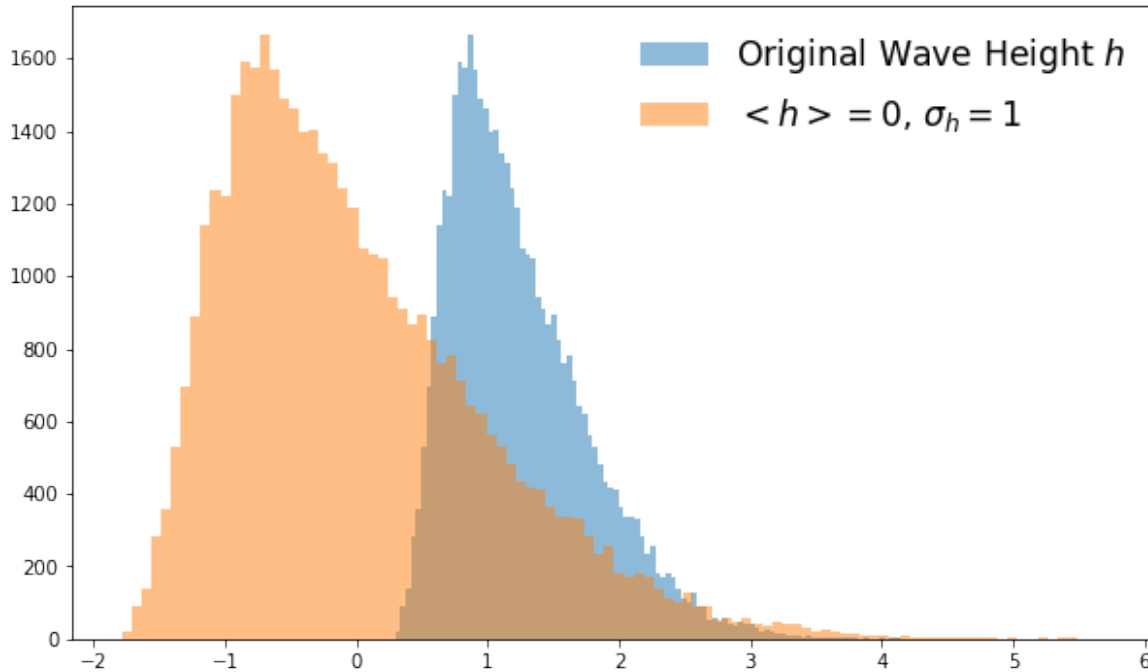
# Get only normalized column
normalized_cols = [c for c in df.columns.tolist() if '_normalized' in c]
print(df[normalized_cols].head())

      height_normalized  heightMax_normalized  period_normalized \
2        -0.898215          -1.047342         -1.184338
3        -0.884973          -0.757690         -1.117565
4        -0.928484          -1.036201         -1.189723
5        -0.983346          -0.534881         -1.083102
6       -1.004155          -0.835673         -1.132643

      energy_normalized  direction_normalized  temperature_normalized
2        -1.463956          -2.044360          0.762152
3        -1.407891          -0.973294          0.694918
4        -1.643866          -0.314176          0.672506
5        -1.184467          -1.261658          0.672506
6       -1.802020          -1.055683          0.694918
```

One can simply plot the content of a pandas dataframe using the name of the column (more direct alternative than extracting numpy array). For instance, one can compare the evolution of the wave height after the transformation:

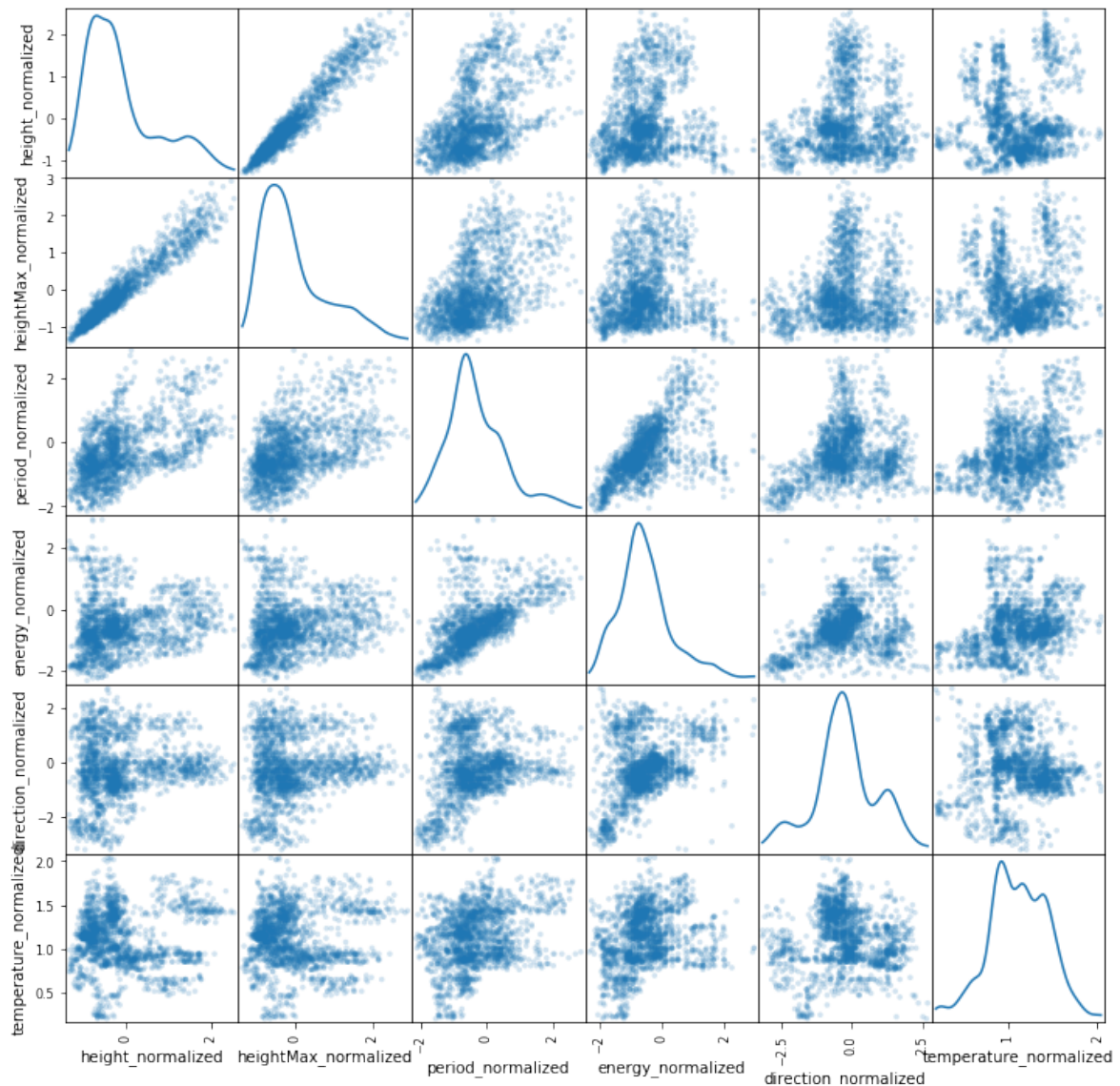
```
plt.figure(figsize=(10, 6))
plt.hist(df['height'], bins=100, alpha=0.5, label='Original Wave Height $h$')
plt.hist(df['height_normalized'], bins=100, alpha=0.5, label='$\langle h \rangle = 0$, $\sigma_h = 1$')
plt.legend(frameon=False, fontsize='xx-large');
```



3.2.5 Data visualization with pandas

There are also many plotting function already included into the pandas library. To show only one example (all functions are described in the [pandas visualization tutorial](#)), here is the *scatter matrix* between variables (defined as a subset of the ones stored the dataframe) obtained in a single line of code:

```
from pandas.plotting import scatter_matrix
scatter_matrix(df[normalized_cols][:2000], figsize=(12, 12), alpha=0.2, s=50,
               diagonal='kde');
```



3.3 scipy: mathematics, physics and engineering

The [scipy](#) project is python-based ecosystem of open-source software for mathematics, science, and engineering. In particular, the following core package are part of it: NumPy, matplotlib, pandas, [scipy library](#) (very quickly introduced here) and [SymPy](#) (symbolic calculations with mathematical expressions *a la* mathematica).

3.3.1 General overview

Obviously, there is no way to extensively present the `scipy` library in this short introduction, but one can quickly summarize few features and illustrate one with a concrete and useful example: fitting data points with a function. Among the main features, the `SciPy` library contains:

- Integration (`scipy.integrate`): integrals, differential equations, etc ...

- Optimization (`scipy.optimize`): minimization, fits, etc ...
- Interpolation (`scipy.interpolate`): smoothing methods, etc ...
- Fourier Transforms (`scipy.fftpack`): spectral analysis, etc ...
- Signal Processing (`scipy.signal`): transfer functions, filtering, etc ...
- Linear Algebra (`scipy.linalg`): matrix operation, diagonalisation, determinant, etc ...
- Statistics (`scipy.stats`): random number, probability density function, cumulative distribution, etc ...

3.3.2 Curve fitting example

```
from scipy import optimize
from scipy import stats
```

Let's now show how to perform a fit of data with error bar using one particular function of `scipy.optimize`. First, we need to generate some data where we choose 20 measurements, with some noise of ~30% and an combined uncertainty of an absolute 0.1 uncertainty and 10% relative uncertainity:

```
Npoints, Nsampling = 20, 1000
xcont = np.linspace(-5.0, 3.5, Nsampling)
x = np.linspace(-5, 3.0, Npoints)
y = 2*(np.sin(x/2)**2 + np.random.random(Npoints)*0.3)
dy = np.sqrt(0.1**2 + (0.1*y)**2)
```

Then we need to define functions with which we want to fit our data, for example a degree 1 polynoms. The syntax has to be `func(x, *pars)`:

```
def pol1(x, p0, p1):
    return p0 + x*p1
```

The following lines actually perform the fit and return both the optimal parameters and the covariances for the dgree 1 polynom:

```
p, cov = optimize.curve_fit(pol1, x, y, sigma=dy)
```

One can then generalize the procedure by plotting the result of the fit for polynoms of several degrees, after having plotted the data. This is a good way to compare different models for the same data. First, we define an arbitrary degree polynom `plo_func()` and we *vectorize* it using `np.vectorize` so that it can accept NumPy arrays:

```
def pol_func(x, *coeff):
    '''Arbitrary degree polynom: f(x) = a0 + a1*x + a2*x^2 + ... aN*x^N'''
    a = np.array([coeff[i]*x**i for i in range(len(coeff))])
    return np.sum(a)

pol_func = np.vectorize(pol_func)
```

In the previous call for `optimize.curve_fit()`, we didn't use additional argument. For this example, we need to specify at least the starting point of the parameters `p0` because *the number of parameter will be assessed using `len(p0)`* (it's not known *a priori* since it is dynamically allocated). Other options can be specified, such as the minimum and maximum allowed values of parameters. Here is a wrap-up function performing the fit for an arbitrary polynomial degree:

```
def fit_poly(degree):
    nPars = degree+1
    p0, pmin, pmax = [1.0]*nPars, [-10]*nPars, [10]*nPars
    fit_options = {'p0': p0, 'bounds': (pmin, pmax), 'check_finite': True}
    par, cov = optimize.curve_fit(pol_func, x, y, sigma=dy, **fit_options)
    return par, cov

degree_max = 12
```

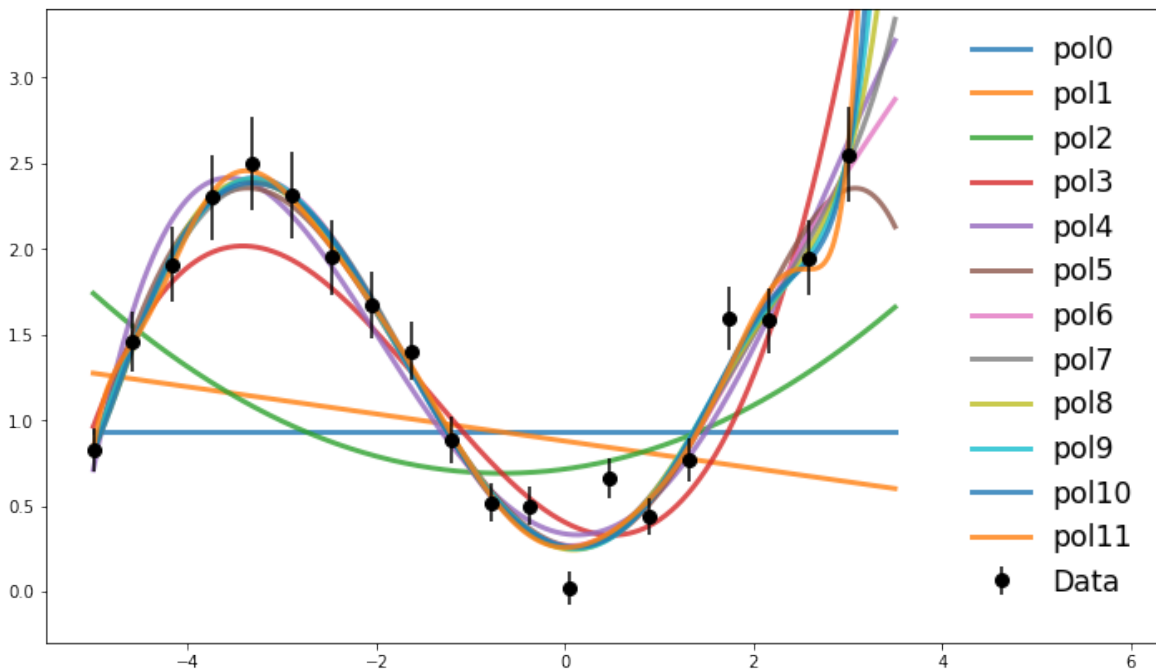
The following code try every polynomial functions up to a degree `degree_max`, perform the fit and overlay the result for each together with the experimental data on the same figure:

```
# Figure for the result
fig = plt.figure(figsize=(12,7))

# Fitting & plotting
for d in np.arange(0, degree_max):
    par, cov = fit_poly(d)
    plt.plot(xcont, pol_func(xcont, *par), label='pol{}'.format(d),
              linewidth=3, alpha=0.8)

# Plotting data
style = {'marker': 'o', 'color': 'black', 'markersize': 8,
         'linestyle': '', 'zorder': 10, 'label': 'Data'}
plt.errorbar(x, y, yerr=dy, **style)

# Plot cosmetics
plt.xlim(-5.5, 6.3)
plt.ylim(-0.3, 3.4)
plt.legend(frameon=False, fontsize='xx-large');
```



It is possible to quantify how well a given model explain the observations, computing what we call the *goodness of fit*. In a frequentist approach, this can be assessed by the fraction of pseudo-data coming from - in principle - repeating the exact same experiment, with to a worst agreement for a given model. The agreement can be quantified using $\chi^2 = \sum_{i=1}^n \frac{(y_i - f(x_i))^2}{\sigma_i^2}$ and its probability density function (PDF) directly gives access to the fraction of “worst pseudo-data” (by integrating the PDF from χ^2 to ∞). More precisely, one can use the cumulative distribution function (CDF) of χ^2 computed with n degrees of freedom, for instance `Npoints`, i.e. `len(x)`. More details can be found, for example, in the [statistics review of the Particle Data Group](#). The following two functions allow to compute the goodness of fit:

```
def get_chi2_nDOF(y, dy, yfit):
    r = (y-yfit)/dy
    return np.sum(r**2), len(y)

def get_pvalue(chi2, nDOF):
    return 1-stats.chi2.cdf(chi2, df=nDOF)
```

We can now perform all these fits and extract the goodness of fit (χ^2 and *p*-value) for each model:

```
# Fitting and getting p-value
degree, chiSquare, pvalue = [], [], []
for d in np.arange(degree_max):
    par, cov = fit_poly(d)
    c2, n = get_chi2_nDOF(y, dy, pol_func(x, *par))
    degree.append(d), chiSquare.append(c2), pvalue.append(get_pvalue(c2, n))
```

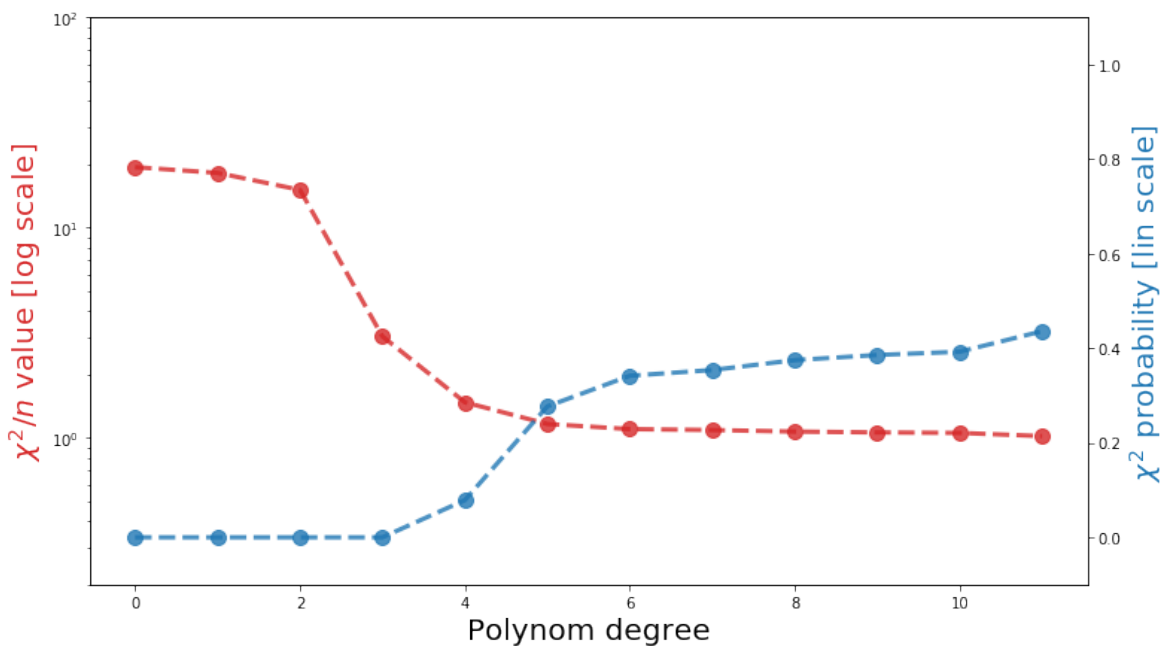
The following piece of code plot both the χ^2 and the *p*-value versus the degree of the polynomial using two different y-axis. This gives another way to use matplotlib by defining explicit object such as `ax` and `fig` and

call methods on those (called *stateless approach*), instead of using function on plt (called *stateful approach*). For more details on these different approaches, see this [RealPython post](#).

```
# Plotting the result with 2 different axis
fig, ax1 = plt.subplots(figsize=(12,7))
ax1.set_xlabel('Polynom degree', fontsize=20)
style = {'marker': 'o', 'markersize': 10, 'alpha': 0.8,
         'linestyle': '--', 'linewidth': 3}

# Plot chi2/n
ax1.semilogy(degree, np.array(chiSquare)/Npoints, color='tab:red', **style)
ax1.set_yscale('log')
ax1.set_ylim(0.2, 100)
ax1.set_ylabel('$\chi^2/n$ value [log scale]', color='tab:red', fontsize=20)

# Plot p-values
ax2 = ax1.twinx()
ax2.plot(degree, pvalue, color='tab:blue', **style)
ax2.set_yscale('linear')
ax2.set_ylim(-0.1, 1.1)
ax2.set_ylabel('$\chi^2$ probability [lin scale]', color='tab:blue', fontsize=20);
```



Chapter 4

High dimensional data manipulation

The present chapter makes use of the concept previously introduced to perform computation that one would do with high dimensional data. Typically, if a given dataset consist of several 3D positions for each observation, one has to deal with many numbers. It is possible that grouping these vectors by pairs is relevant to understand the problem. Or maybe other operation within these various 3D vector is useful. Since we want to use the full power of numpy, all these computations cannot be done with an explicit loop over observations and/or over vectors.

This chapter consider few of these typical use cases and their implementation using numpy, using a simple toy dataset made by hand. Most likely, you will never face such a situation for machine learning algorithm, but it is good to go trough these examples to show some of the limitation of not being able to loop overs observations.

Let's first perform the usual imports:

```
import itertools
import numpy as np
import matplotlib.pyplot as plt
import matplotlib as mpl
%matplotlib inline
```

Then, one can setup he default matplotlib style for all following plots (more details on available option can be found on [how to customize matplotlib](#)):

```
mpl.rcParams['legend.frameon'] = False
mpl.rcParams['legend.fontsize'] = 'xx-large'
mpl.rcParams['xtick.labelsize'] = 16
mpl.rcParams['ytick.labelsize'] = 16
mpl.rcParams['axes.titlesize'] = 18
mpl.rcParams['axes.labelsize'] = 18
mpl.rcParams['lines.linewidth'] = 2.5
mpl.rcParams['figure.figsize'] = (10, 7)
```

4.1 Data model and goals

We consider 1 millions observations, each defined by ten 3D vectors (r_0, \dots, r_9) where $r_i = (x, y, z)$ (arrow for vector will be omitted from now on). These pseudo-data can represent position in space or RGB colors for an image. This is just an example to play with and apply numpy concepts for both simple computations (element-by-element functions, statistics calculations) and more complex computation exploiting the multi-dimensional structure of the data. For example, one might want to compute the distance between all pairs (r_i, r_j) , which has to be done without loop.

Using the `np.random` module, it is possible to generate n-dimensional arrays easily. In our case, we want to generate an array containing our observations with have 3 dimensions (or *axis* in numpy language), and the size along each of these axis will have the following value and meaning:

- `axis=0`: over 1 million events
- `axis=1`: over 10 vectors
- `axis=2`: over 3 coordinates

```
r = np.random.random_sample((1000000, 10, 3))
```

It is possible to print the first two observations as follow:

```
print(r[0:2])
```

```
[[[3.12646048e-01 4.55257576e-01 7.62120920e-01]
 [7.57904883e-01 5.75783716e-01 9.85730373e-01]
 [8.58351019e-01 9.97112982e-01 6.94945548e-02]
 [3.97010641e-01 3.30452282e-01 4.76705513e-01]
 [1.90192515e-01 8.46642981e-01 9.44922049e-01]
 [2.28626376e-01 5.32713270e-01 5.86119632e-02]
 [8.78240385e-01 4.84309389e-01 5.41506300e-01]
 [9.04149582e-01 4.92954799e-01 2.21837932e-01]
 [7.92243462e-01 9.92160857e-01 5.22952886e-01]
 [5.90601463e-01 8.57334963e-01 5.76432781e-01]]

[[4.89732288e-01 2.45947658e-01 5.24605965e-01]
 [2.79684077e-01 1.75887137e-01 5.96777979e-01]
 [6.36118572e-01 8.08656904e-01 5.67401037e-01]
 [5.01315803e-01 3.79415584e-01 9.73566504e-05]
 [8.04499847e-01 4.01132808e-01 2.73607136e-01]
 [4.62946717e-01 8.46061510e-01 8.82090460e-01]
 [2.30961828e-02 6.79642827e-01 4.79125888e-01]
 [7.51716668e-01 9.00985276e-01 6.87922769e-01]
 [9.47722015e-01 3.27310436e-01 1.49407680e-02]
 [3.36817391e-01 2.63926051e-01 6.90325842e-01]]]
```

4.2 Mean over the differents axis

4.2.1 Mean over observations (axis=0)

This mean will average all observations *i.e.* over the first dimension, returning an array of dimension (10, 3) corresponding to the average $r_i = (x_i, y_i, z_i)$ over the observations.

```
m0 = np.mean(r, axis=0)
print(m0.shape)
```

(10, 3)

Note the computation time of 30ms for 30 averages over a million number:

```
%timeit np.mean(r, axis=0)
```

29.9 ms ± 165 µs per loop (mean ± std. dev. of 7 runs, 10 loops each)

While it takes 10 times longer for a *single mean* over a million number with an explicit loop, so **the gain of vectorization is a factor 300**:

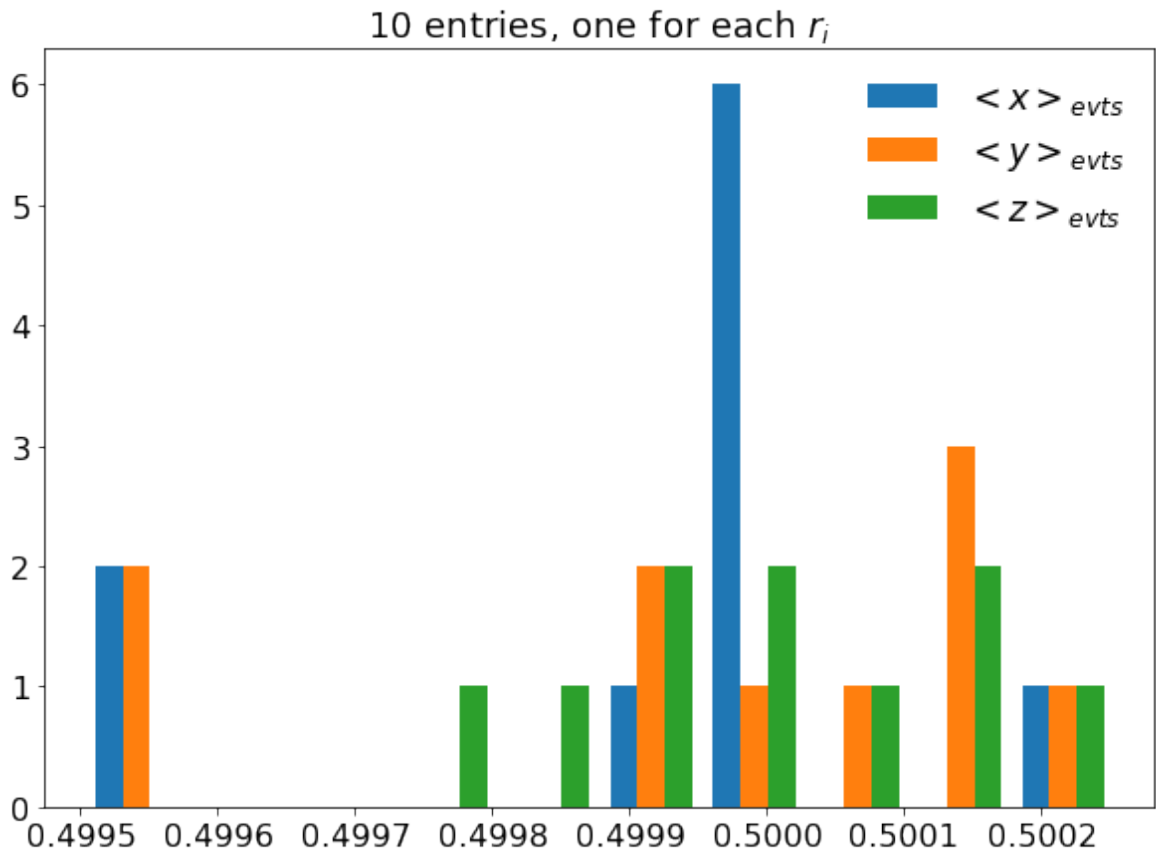
```
def explicit_loop(array):
    res=0
    for a in array:
        res += a/len(array)
```

```
%timeit explicit_loop(np.random.random_sample(size=1000000))
```

286 ms ± 1.32 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)

The distributions of m0 obtained with plt.hist() results into three separate histograms (one for each x, y, z) each having 10 entries (one per r_i):

```
plt.hist(m0, label=['$<x>_{evts}$', '$<y>_{evts}$', '$<z>_{evts}$'])
plt.title('10 entries, one for each $r_i$')
plt.legend();
```



4.2.2 Mean over the 10 vectors (axis=1)

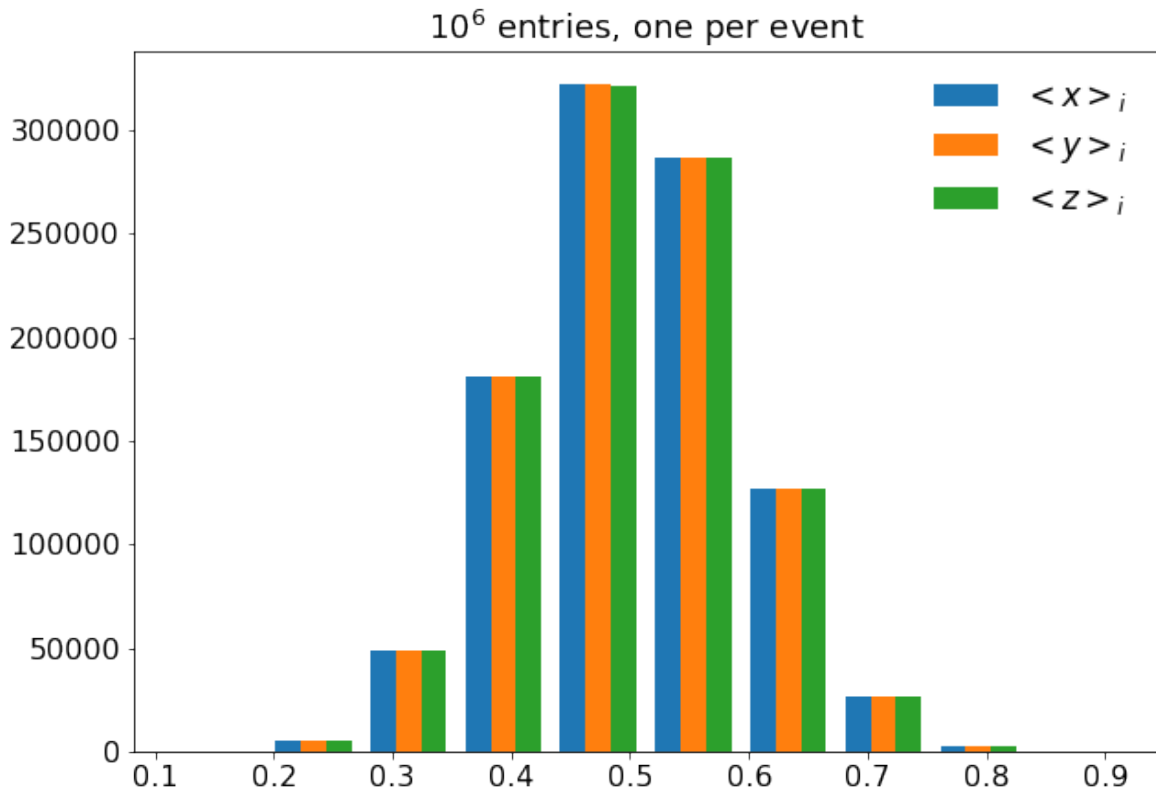
This one will compute the average over the 10 vectors, for each observations, reducing into a (1000000, 3) shape array, as seen below. This is 3D barycenter of each observation.

```
m1 = np.mean(r, axis=1)
print(m1.shape)
```

(1000000, 3)

One can plot the obtained array `m1` using `plt.hist()`, which results into 3 histograms of a million entry each:

```
plt.hist(m1, label=['$\langle x \rangle_{\text{evts}}$', '$\langle y \rangle_{\text{evts}}$', '$\langle z \rangle_{\text{evts}}$'])
plt.title('$10^6$ entries, one per event')
plt.legend();
```



4.2.3 Mean over the coordinates (axis=2)

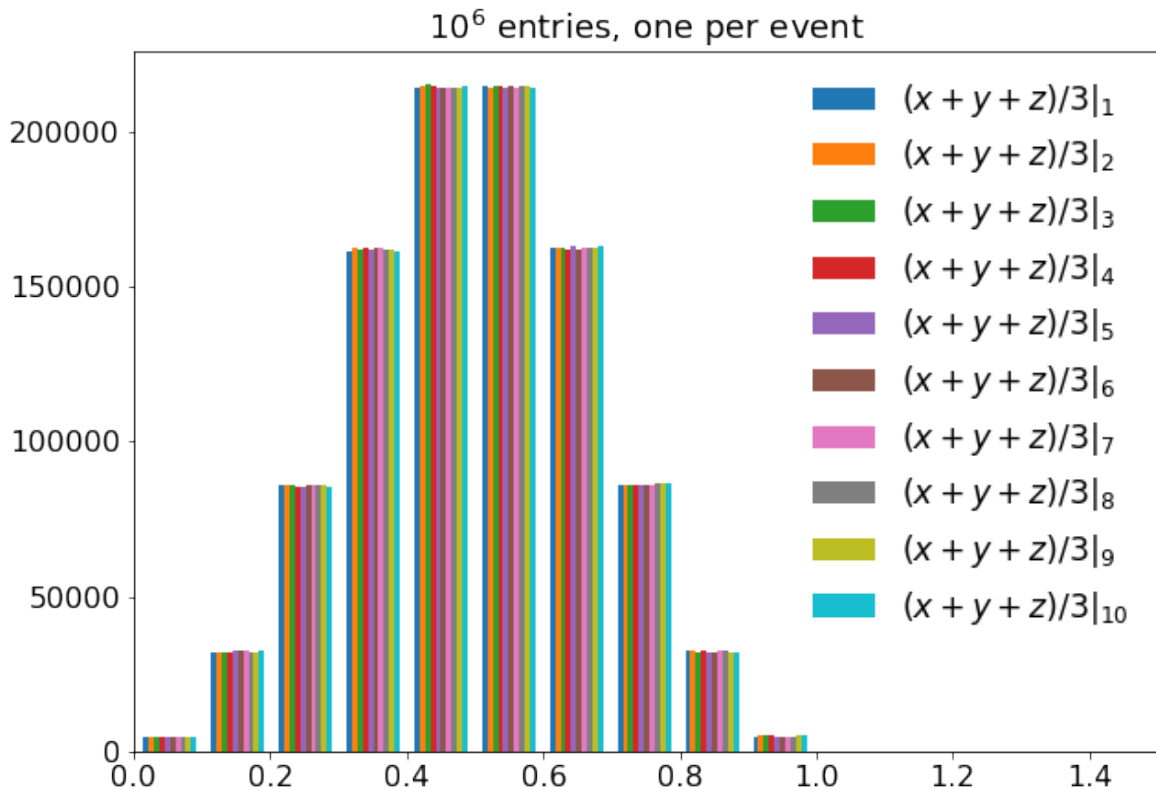
This directly computes the average over the three coordinates $(x + y + z)/3$ for each vector of each event, resulting in 10 values per event:

```
m2 = np.mean(r, axis=2)
print(m2.shape)
```

(1000000, 10)

The `plt.hist()` of the resulting array `m2` corresponds then to 10 histograms of a million entries each:

```
names = ['$({x+y+z})/3|_{+{}}'.format(i)+'}$' for i in range(1, 11)]
plt.hist(m2, label=names)
plt.title('$10^6$ entries, one per event')
plt.xlim(0, 1.5)
plt.legend();
```



4.3 Distance computation

Computing particular distances inside a given event is relevant for many applications (distances here can be seen as any type of metric). For example, these computations are crucial in learning algorithms based on nearest neighbor approach. In collider physics, it's always useful to compute angle between two objects (tracks, deposit, particles, ...) in order to compute invariant masses, or isolation in a given cone, etc ...

4.3.1 Distance to a reference r_0

We can start simple by defining a new origin r_0

```
r0 = np.array([1, 2, 1])
```

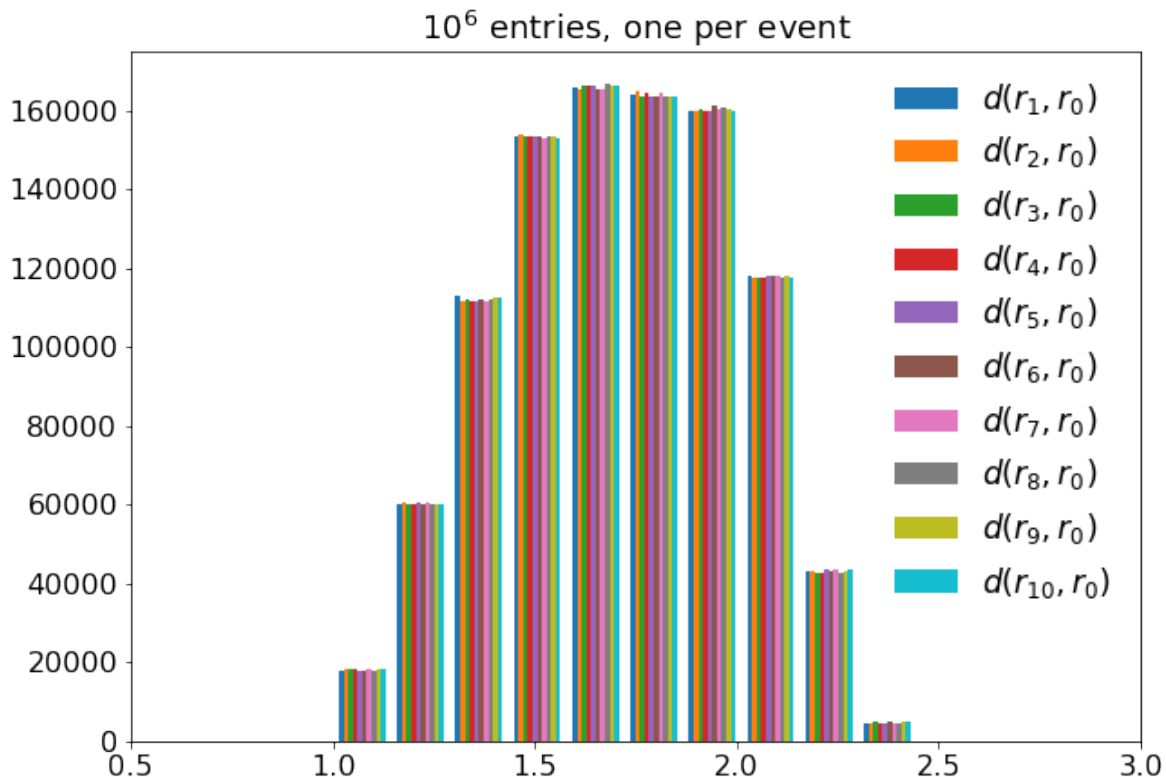
and compute the distance to this new origin for all points, using $\star\star 2$ to square all numbers, perform the sum over the coordinate (`axis=2`) and square-root everything with $\star\star 0.5$:

```
d = np.sum((r-r0)**2, axis=2)**0.5
print(d.shape)
```

(1000000, 10)

As expected the result is 10 numbers for each of the events, which can be easily plotted:

```
names = ['$d(r_{'+str(i)}).format(i),r_0)' for i in range(1, 11)]
plt.hist(d, label=names)
plt.title('10^6 entries, one per event')
plt.xlim(0.5, 3)
plt.legend();
```



4.3.2 Distance between r_i and $\langle r \rangle_i$ for each event

Another calculation is to compute the averaged position for each event and see how distant each vector is from this position. To perform such a calculation, we will use numpy array broadcasting. Let's first compute the average position for every events:

```
r_mean = np.mean(r, axis=1)
```

Now, let's broadcast this array of shape $(1e6, 3)$ with the full dataset, *i.e.* an array of shape $(1e6, 10, 3)$, by computing the distance for each point:

```
try:
    d_to_mean = np.sum((r-r_mean)**2, axis=2)**0.5
except ValueError:
    print('Impossible for {} and {}'.format(r.shape, r_mean.shape))
```

```
Impossible for (1000000, 10, 3) and (1000000, 3)
```

There is one missing dimension, describing the 10 positions, which has to be created so that the array can be copied 10 times over along this dimension:

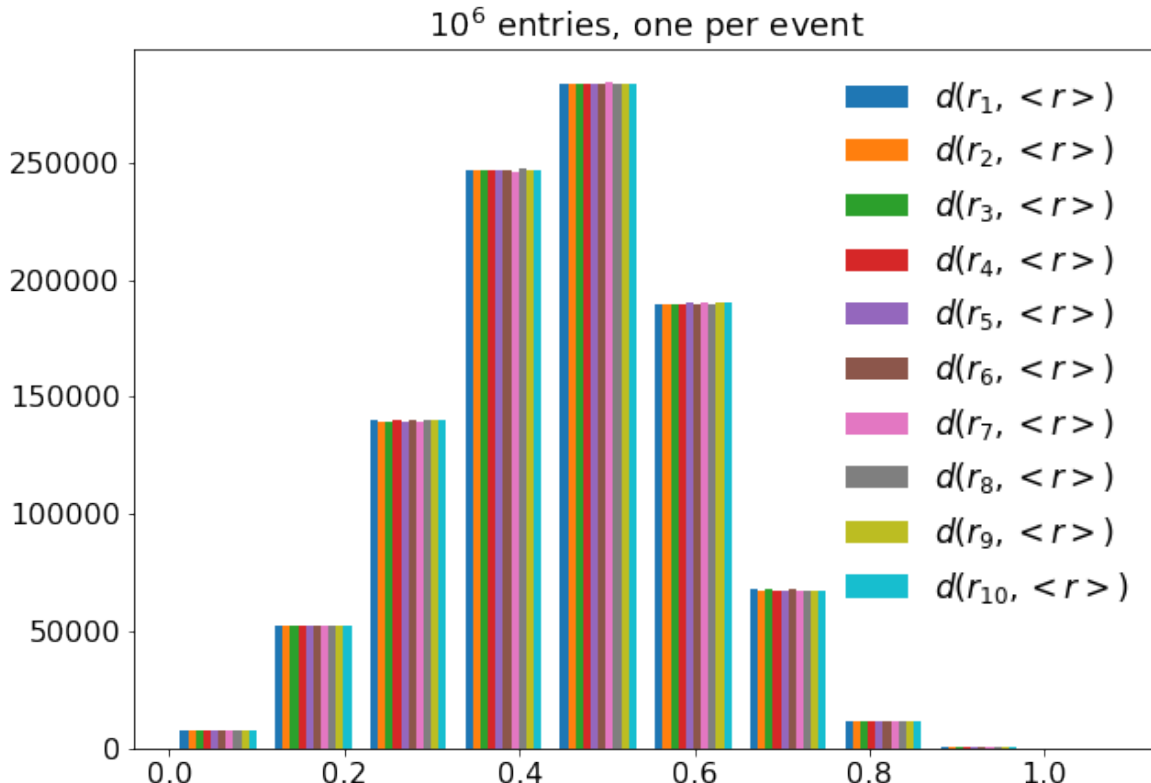
```
r_mean_3d = r_mean[:, np.newaxis, :]
```

We can now retry the operation:

```
try:
    d_to_mean = np.sum((r - r_mean_3d)**2, axis=2)**0.5
    print('Possible for {} and {}'.format(r.shape, r_mean_3d.shape))
except ValueError:
    print('Impossible for {} and {}'.format(r.shape, r_mean_3d.shape))
```

```
Possible for (1000000, 10, 3) and (1000000, 1, 3)
```

```
names = ['$d(r_{'+str(i)+'})'.format(i)+r')$' for i in range(1, 11)]
plt.hist(d_to_mean, label=names)
plt.title('$10^6$ entries, one per event')
plt.legend();
```



4.4 Pairing 3D vectors for each observation, without a loop

Being able to pair objects is obviously important for many type of calculations. This allows to probe correlations at the first order, to identify sub-systems, etc ... In a traditional way, a pairing would involve a *for* loop in which the combinatorics can be done for each event. Working with numpy, one has to perform the combinatorics in a vectorized way and return a new numpy array containing all the pairs. Once done, one can perform many types of computations on this new array.

4.4.1 Finding all possible (r_i, r_j) pairs for all events

One solution to perform such a task without *for* loop was found on [stackoverflow](#). The idea is to simply work on indices to build the pairs (since it doesn't really matter what are the nature of the objects), and use numpy *fancy* indexing. Let proceed step by step with a smallest array to understand the procedure (namely 2 observations of 5 positions):

```
a = r[0:2,0:5]
print(a)

[[[3.12646048e-01 4.55257576e-01 7.62120920e-01
 [7.57904883e-01 5.75783716e-01 9.85730373e-01]
 [8.58351019e-01 9.97112982e-01 6.94945548e-02]
 [3.97010641e-01 3.30452282e-01 4.76705513e-01]
 [1.90192515e-01 8.46642981e-01 9.44922049e-01]]

 [[4.89732288e-01 2.45947658e-01 5.24605965e-01]
 [2.79684077e-01 1.75887137e-01 5.96777979e-01]
 [6.36118572e-01 8.08656904e-01 5.67401037e-01]
 [5.01315803e-01 3.79415584e-01 9.73566504e-05]
 [8.04499847e-01 4.01132808e-01 2.73607136e-01]]]
```

Since we want to work with the indices of the 5 vectors, we create a numpy array of integer going from 0 to 4 (`a.shape[1]` is the number of elements along the second dimension, i.e. 5):

```
array_indices = np.arange(a.shape[1])
print(array_indices)
```

```
[0 1 2 3 4]
```

Then, we use the package `itertools` to deal with the combinatorics. This will return an *iterator* that can be turned into a numpy array using `np.fromiter()`. But this function requires to specify the data type `dt`, which is done using a structured array synthax here (i.e. `[(varName1, type1), (varName2, type2)]`). For more details on data type, check this [documentation page](#).

```
dt = np.dtype([('index1', np.intp), ('index2', np.intp)])
print(dt)
```

```
[('index1', '<i8'), ('index2', '<i8')]
```

```
array_indice_comb = np.fromiter(itertools.combinations(array_indices, 2), dt)
print(array_indice_comb)
```

```
[(0, 1) (0, 2) (0, 3) (0, 4) (1, 2) (1, 3) (1, 4) (2, 3) (2, 4) (3, 4)]
```

The next step is to format these numbers in a indices array with the proper dimension, so that when we do `a[indices]`, we get all the pairs. For instance, we need to have all 10 pairs, each with two elements corresponding to a shape `indices.shape=(10,2)`. We can achieve this in two steps:

1. `array_indice_comb.view(np.intp)` return the exact same data of `array_indice_comb` as a 1D array of positive integer.
2. we reshape the resulting array with `reshape(-1, 2)`, where -1 means "compute the size of the first dimension to have 2 objects (we want pairs!) in the second dimension.

```
indices = array_indice_comb.view(np.intp).reshape(-1, 2)
print(indices)
```

```
[[0 1]
 [0 2]
 [0 3]
 [0 4]
 [1 2]
 [1 3]
 [1 4]
 [2 3]
 [2 4]
 [3 4]]
```

The final step is to exploit fancy indexing along `axis=1` i.e. the 5 spatial positions. In practice, for each observation `iobs`, we want to have `a[iobs, indices]`. There are two ways to do this: (a) `a[:, indices]` or (b) using the numpy function `np.take(a, indices, axis)` which makes the code more independent from the structure of `a`:

```
a_pairs = np.take(a, indices, axis=1)
print(a_pairs.shape)
```

```
(2, 10, 2, 3)
```

```
a_pairs = a[:,indices]
print(a_pairs.shape)
```

```
(2, 10, 2, 3)
```

We have now 2 events, each having 10 pairs, each having 2 objects (still a pair!), each having 3 coordinates (spatial positions). We can print all the 10 pairs for the first observation:

```
print(a_pairs[0])

[[[0.31264605 0.45525758 0.76212092]
 [0.75790488 0.57578372 0.98573037]]

 [[0.31264605 0.45525758 0.76212092]
 [0.85835102 0.99711298 0.06949455]]

 [[0.31264605 0.45525758 0.76212092]
 [0.39701064 0.33045228 0.47670551]]

 [[0.31264605 0.45525758 0.76212092]
 [0.19019251 0.84664298 0.94492205]]

 [[0.75790488 0.57578372 0.98573037]
 [0.85835102 0.99711298 0.06949455]]

 [[0.75790488 0.57578372 0.98573037]
 [0.39701064 0.33045228 0.47670551]]

 [[0.75790488 0.57578372 0.98573037]
 [0.19019251 0.84664298 0.94492205]]

 [[0.85835102 0.99711298 0.06949455]
 [0.39701064 0.33045228 0.47670551]]

 [[0.85835102 0.99711298 0.06949455]
 [0.19019251 0.84664298 0.94492205]]

 [[0.39701064 0.33045228 0.47670551]
 [0.19019251 0.84664298 0.94492205]]]
```

Once understood, we can wrap-up this code into a function where we generalize the number of objects we want to group `n` and the axis along which we want to group `axis`:

```
def combs_nd(a, n, axis=0):
    i = np.arange(a.shape[axis])
    dt = np.dtype([('i', np.intp)]*n)
    i = np.fromiter(itertools.combinations(i, n), dt)
    i = i.view(np.intp).reshape(-1, n)
    return np.take(a, i, axis=axis)
```

As a sanity check, we can re-compute `a_pair` and compare with the previous results:

```
a_pairs = combs_nd(a=r[0:2,0:5], n=2, axis=1)
print(a_pairs[0])
```

```
[[[0.31264605 0.45525758 0.76212092]
 [0.75790488 0.57578372 0.98573037]]

 [[0.31264605 0.45525758 0.76212092]
 [0.85835102 0.99711298 0.06949455]]

 [[0.31264605 0.45525758 0.76212092]
 [0.39701064 0.33045228 0.47670551]]

 [[0.31264605 0.45525758 0.76212092]
 [0.19019251 0.84664298 0.94492205]]

 [[0.75790488 0.57578372 0.98573037]
 [0.85835102 0.99711298 0.06949455]]

 [[0.75790488 0.57578372 0.98573037]
 [0.39701064 0.33045228 0.47670551]]

 [[0.85835102 0.99711298 0.06949455]
 [0.39701064 0.33045228 0.47670551]]

 [[0.85835102 0.99711298 0.06949455]
 [0.19019251 0.84664298 0.94492205]]

 [[0.39701064 0.33045228 0.47670551]
 [0.19019251 0.84664298 0.94492205]]]
```

It can be interesting to see that this operation takes less than a second for a million observations of 10 vectors, meaning 45 pairs:

```
%timeit combs_nd(a=r, n=2, axis=1)
```

```
966 ms ± 33.3 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)
```

4.4.2 Computing (minimum) distances on these pairs

Once we have these pairs, we can for example compute all the distances and find which pair has the closest objects. Starting with the pairs:

```
pairs = combs_nd(a=r, n=2, axis=1)
```

We can then define the vectorial difference between the two position of a pair, and compute the euclidian distance:

```
dp = pairs[:, :, 0] - pairs[:, :, 1]
distances = (np.sum(dp**2, axis=2))**0.5
```

And get the minimum distance for each event:

```
smallest_distance = np.min(distances, axis=1)
print(smallest_distance.shape)
```

(1000000,)

All these instructions can be put into a function which can be timed:

```
def compute_dr_min(a):
    pairs = combs_nd(a, 2, axis=1)
    i1 = tuple([None, None, 0, None])
    i2 = tuple([None, None, 1, None])
    return np.min(np.sum((pairs[i1] - pairs[i2])**2, axis=2)**0.5, axis=1)
```

```
%timeit compute_dr_min(r)
```

980 ms ± 122 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)

Note that doing all operations in the less possible amount of lines can significantly speed up the process. Let's define another function where the difference between the pair elements is done separately:

```
def compute_dr_min_more_steps(a):
    pairs = combs_nd(a, 2, axis=1)
    dp = pairs[:, :, 0] - pairs[:, :, 1]
    return np.min(np.sum(dp**2, axis=2)**0.5, axis=1)
```

And let's compare the performance on 0.2 million observations:

```
%timeit compute_dr_min(a=r[:200000])
%timeit compute_dr_min_more_steps(a=r[:200000])
```

194 ms ± 1.77 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)
606 ms ± 5.13 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)

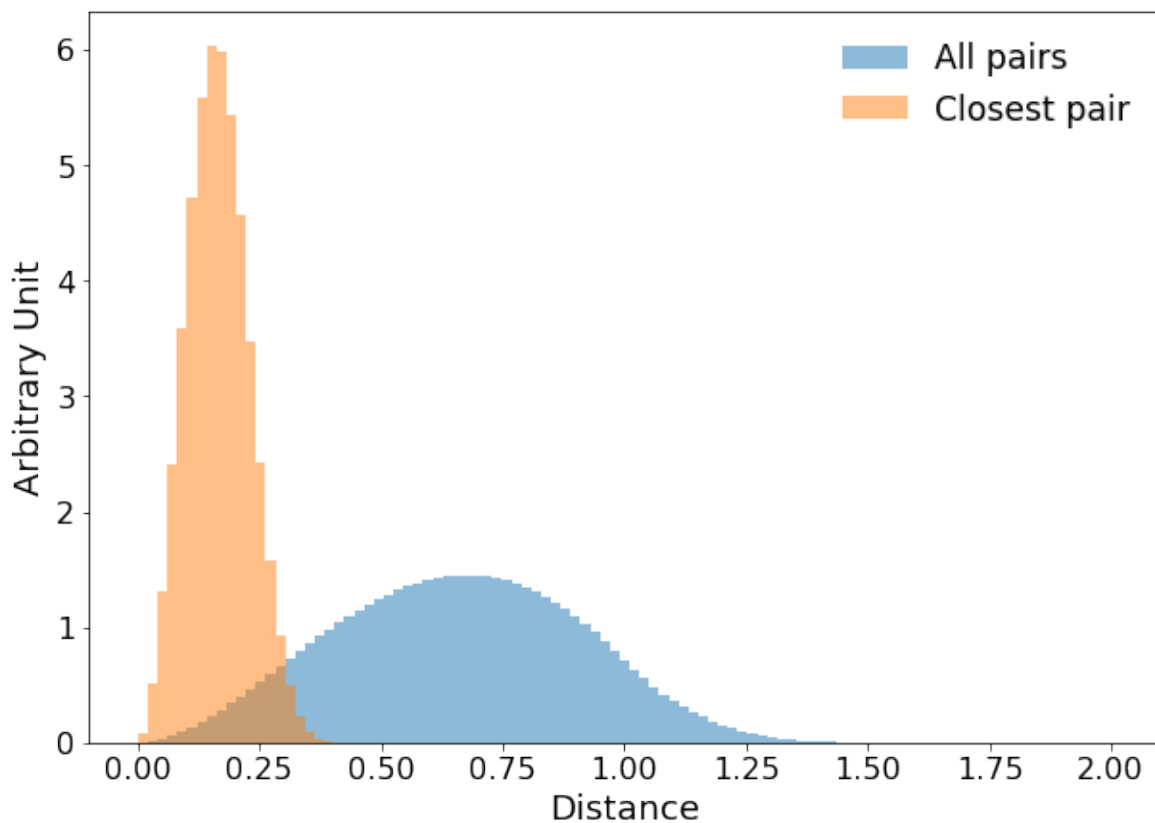
Let's now plot the distributions of all distances for all the pairs (using `flatten()` function which returns a 1D array), and only the pair having the smallest distances:

```

plot_style = {
    'bins': np.linspace(0, 2, 100),
    'alpha': 0.5,
    'density': True,
}

plt.hist(distances.flatten(), label='All pairs', **plot_style)
plt.hist(smallest_distance, label='Closest pair', **plot_style)
plt.xlabel('Distance')
plt.ylabel('Arbitrary Unit')
plt.legend();

```



4.5 Selecting a subset of r_i based on (x, y, z) values, without loop

The next step in our exploration “loop-less calculations” is to be able to perform the same kind of computation described above but only on a subset of positions, selected according to a given criteria. For example, we might want to keep particles only if there have positive charge. Many obvious application can be found in other physics field and/or machine learning. Let’s start with accessing the three arrays of coordinates in order to select points based on some easy criteria.

```
x, y, z = r[:, :, 0], r[:, :, 1], r[:, :, 2]
```

4.5.1 Counting number of points among the 10 with $x_i > y_i$ in each event

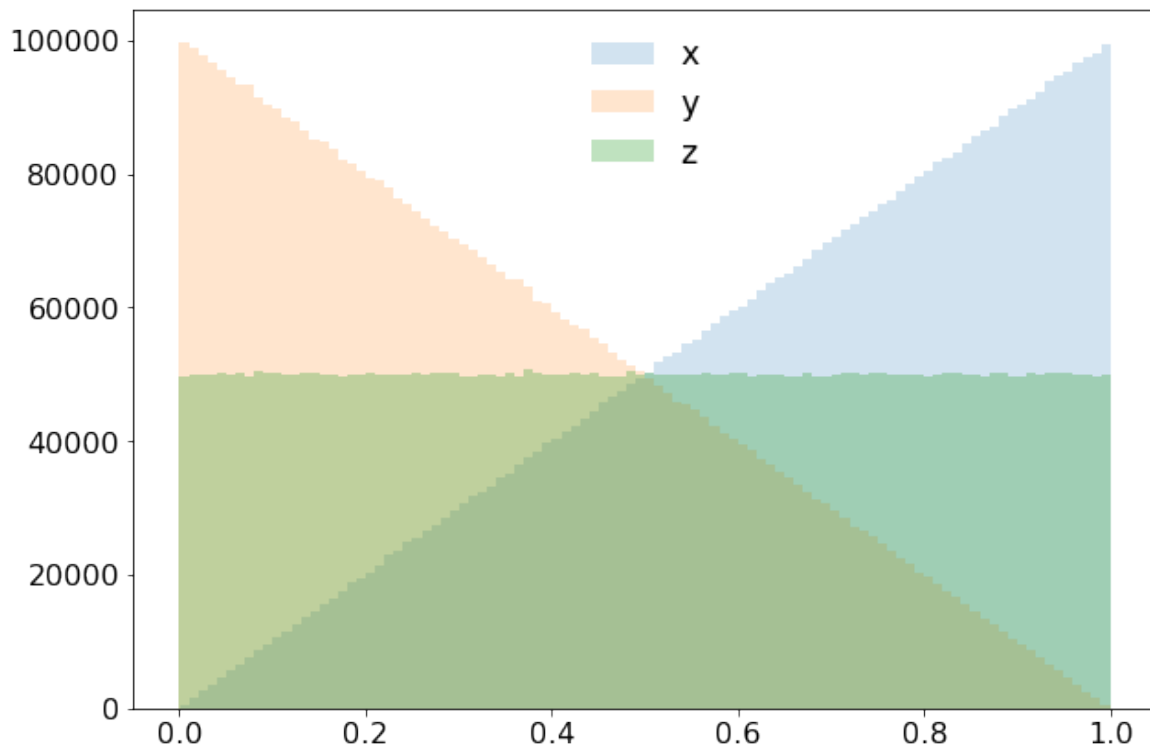
We will use the numpy masking feature described in the first chapter, defining a index of boolean based on x and y arrays:

```
idx = x > y
print(idx.shape)
```

```
(1000000, 10)
```

We can quickly check the distribution for the selected coordinates: x and y are anti correlated - as expected - while z is flat - as expected.

```
plt.hist(x[idx], bins=100, alpha=0.2, label='x')
plt.hist(y[idx], bins=100, alpha=0.2, label='y')
plt.hist(z[idx], bins=100, alpha=0.3, label='z')
plt.legend();
```



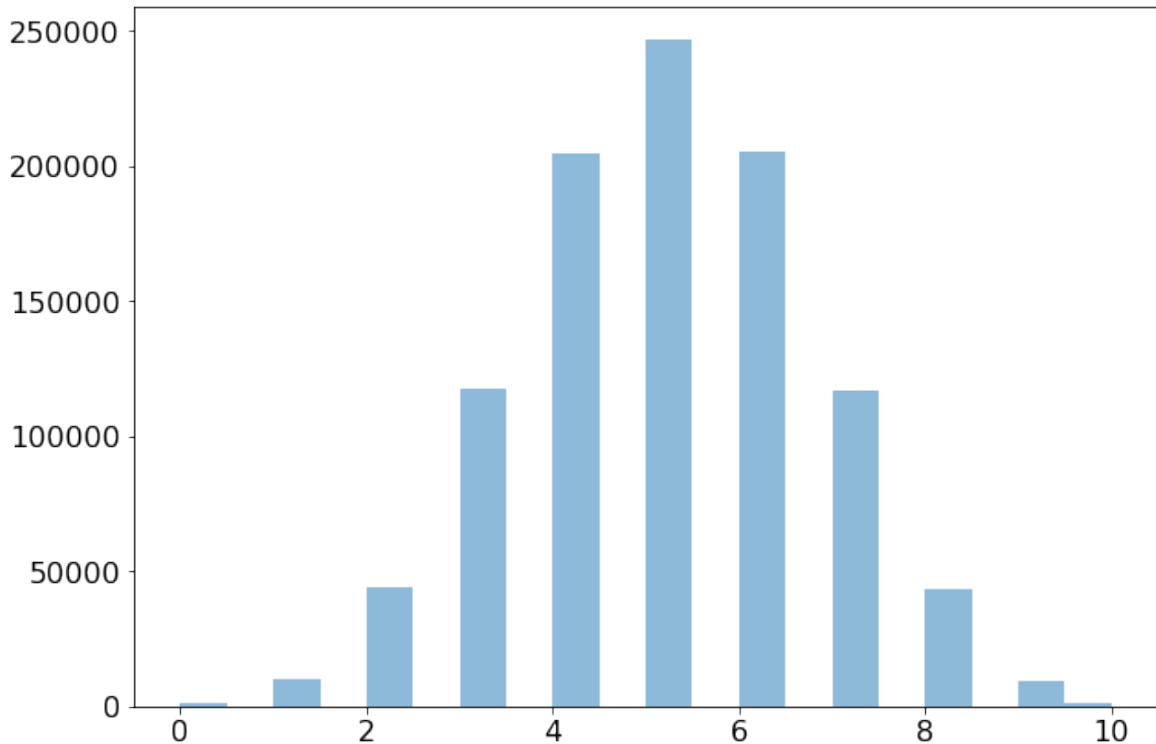
If we want to better understand how this selection affect our data, one might want to count the number of points per event satisfying this selection, using `np.count_nonzero()` on the boolean array along the axis representing the 10 vectors `axis=1`:

```
c = np.count_nonzero(idx, axis=1)
print(c.shape)
```

(1000000,)

We can then plot the distribution of this number over all the events:

```
plt.hist(c, bins=20, alpha=0.5);
```



4.5.2 Plotting z for the two types of population ($x > y$ and $x < y$)

This is obviously useful to inspect the different populations - something we want to do very often. For the plotting purpose, let's consider only the 500 first observations that we dump into sx , sy , sz (s for small):

```
sx, sy, sz = x[0:500, ...], y[0:500, ...], z[0:500, ...]
```

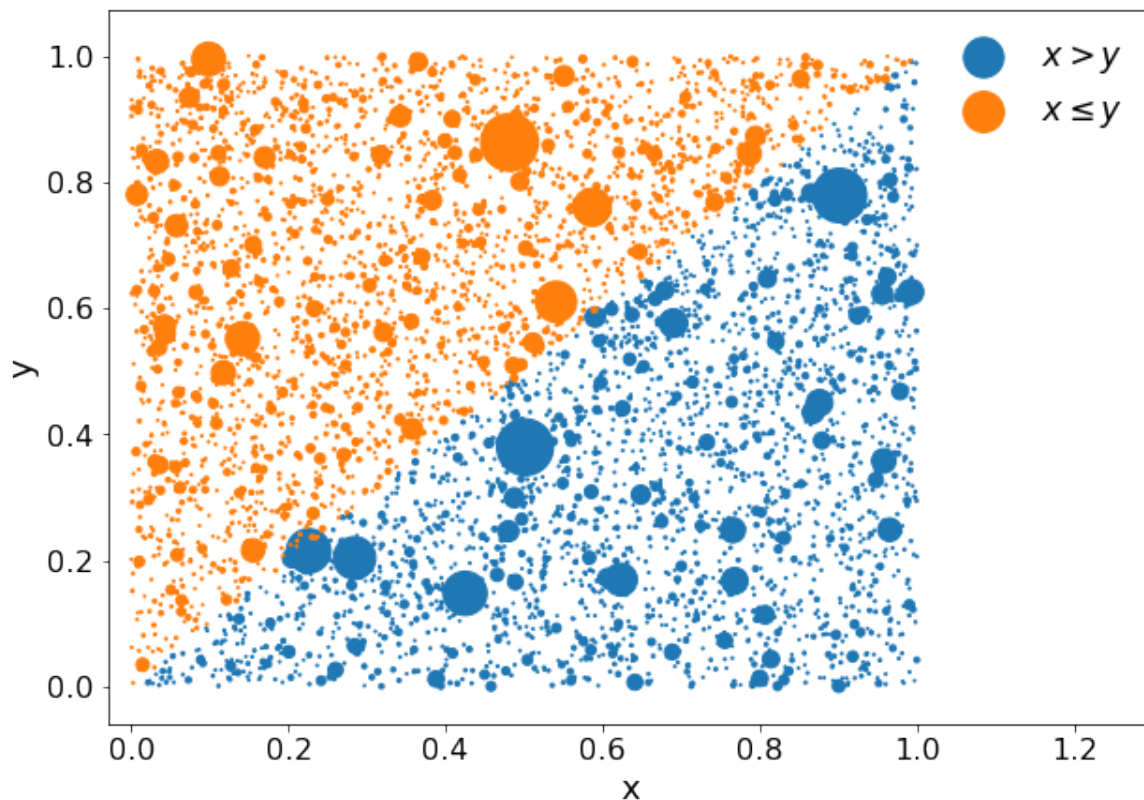
We define the mask computed on these small arrays $smask$:

```
smask = sx>sy
```

And we can plot the result in the 2D plane (x, y) with the z coordinate as marker size, for instance $1/(z + 10^{-3})$. The two populations are defined using both $smask$ and $\sim smask$ to make sure the union of the two is the original dataset:

```

plt.scatter(sx[smask], sy[smask], s=(sz[smask]+1e-3)**-1, label='$x>y$')
plt.scatter(sx[~smask], sy[~smask], s=(sz[~smask]+1e-3)**-1, label='$x\leq y$')
plt.xlabel('x')
plt.ylabel('y')
plt.xlim(-0.03, 1.3)
plt.legend();
    
```



4.5.3 Computation of $x_i + y_i + z_i$ sum over a subset of the 10 positions

Once we are able to isolate a subset of points, we might want to compute new numbers only based on those. This is what is proposed here with the sum of the three coordinates. Let's first compute the sum, called `ht`, over all the 10 points:

```

ht1 = np.sum(x+y+z, axis=1)
print(ht1.shape)
    
```

(1000000,)

Apply now a selection, which multiply the value by 0 (i.e. `False`) if the condition is not satisfied:

```
selection = x>y
ht2 = np.sum((x+y+z)*selection, axis=1)
```

Of course, this works only for computation which is not affected by a 0: if we want to compute the product of coordinate, this approach will obviously not work.

```
prod = np.product((x+y+z)*selection, axis=1)
eff = np.count_nonzero(prod>0)/len(prod)
print('Efficiency of prod>0: {:.5f}'.format(eff))
```

Efficiency of prod>0: 0.00093

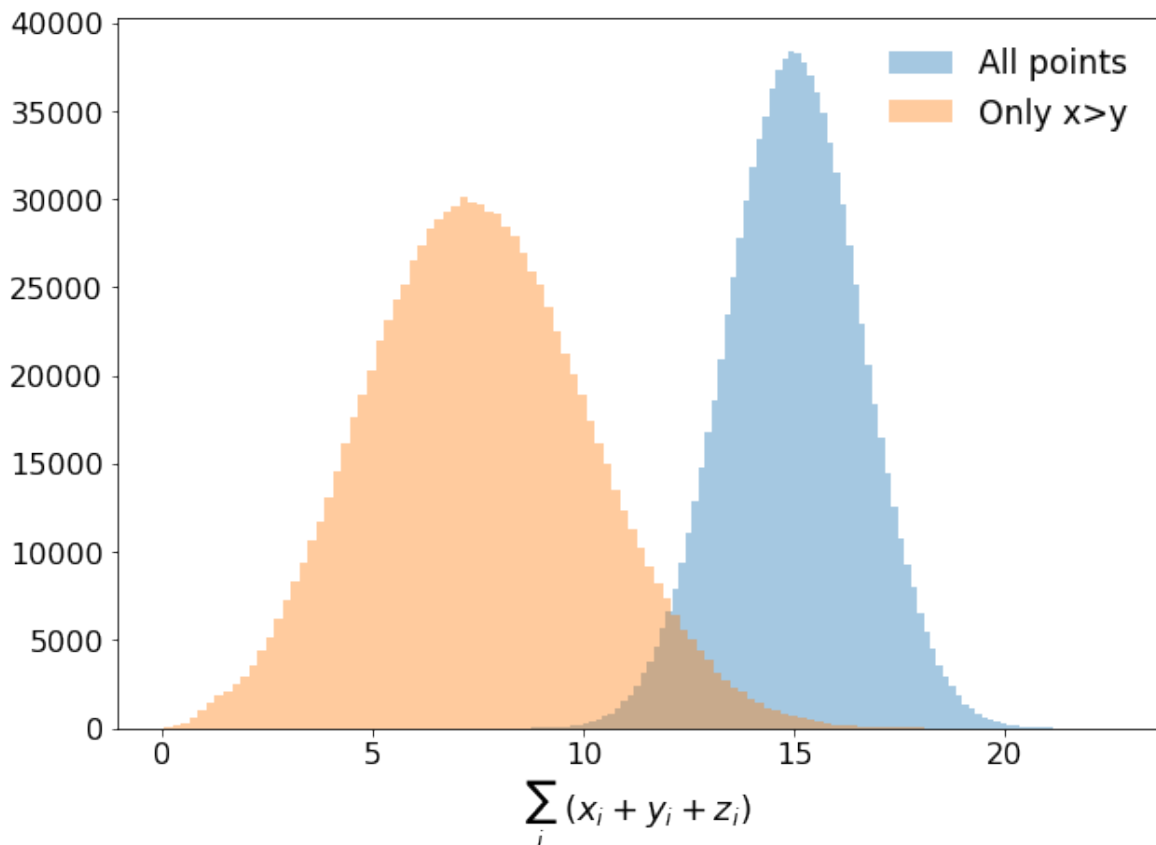
In a more general manner, we should use *masked arrays* which completely remove the masked elements from any computations:

```
mx = np.ma.array(x, mask=selection)
my = np.ma.array(y, mask=selection)
mz = np.ma.array(z, mask=selection)
prod = np.product((mx+my+mz), axis=1)
eff = np.count_nonzero(prod>0)/len(prod)
print('Efficiency of prod>0: {:.5f}'.format(eff))
```

Efficiency of prod>0: 1.00000

Finally one can plot the result, removing the observation with $ht2==0$ (case where all the 10 points have $x \leq y$):

```
plt.hist(ht1, bins=100, alpha=0.4, label='All points')
plt.hist(ht2[ht2>0], bins=100, alpha=0.4, label='Only x>y')
plt.xlabel('$\sum_i (x_i + y_i + z_i)$')
plt.legend();
```



4.5.4 Pairing with a subset of r_i verifying $x_i > y_i$ only

Another computation would be to redo the pairing on the subset of selected position. In order to do so, we follow the same logic, expect that we will directly replace removed values by `nan` in order to be easily identifiable in after the pairing. It's very important to copy the original data with the module `copy`, otherwise, the original data will be modified in the following piece of code:

```
import copy
selection = x>y
selected_r = copy.copy(r)
selected_r[selection] = np.nan
print(selected_r[0])
```

```
[[0.31264605 0.45525758 0.76212092]
 [      nan         nan         nan]
 [0.85835102 0.99711298 0.06949455]
 [      nan         nan         nan]
 [0.19019251 0.84664298 0.94492205]
 [0.22862638 0.53271327 0.05861196]
 [      nan         nan         nan]
 [      nan         nan         nan]
 [0.79224346 0.99216086 0.52295289]
 [0.59060146 0.85733496 0.57643278]]
```

One can now call the pairing function on the filtered dataset:

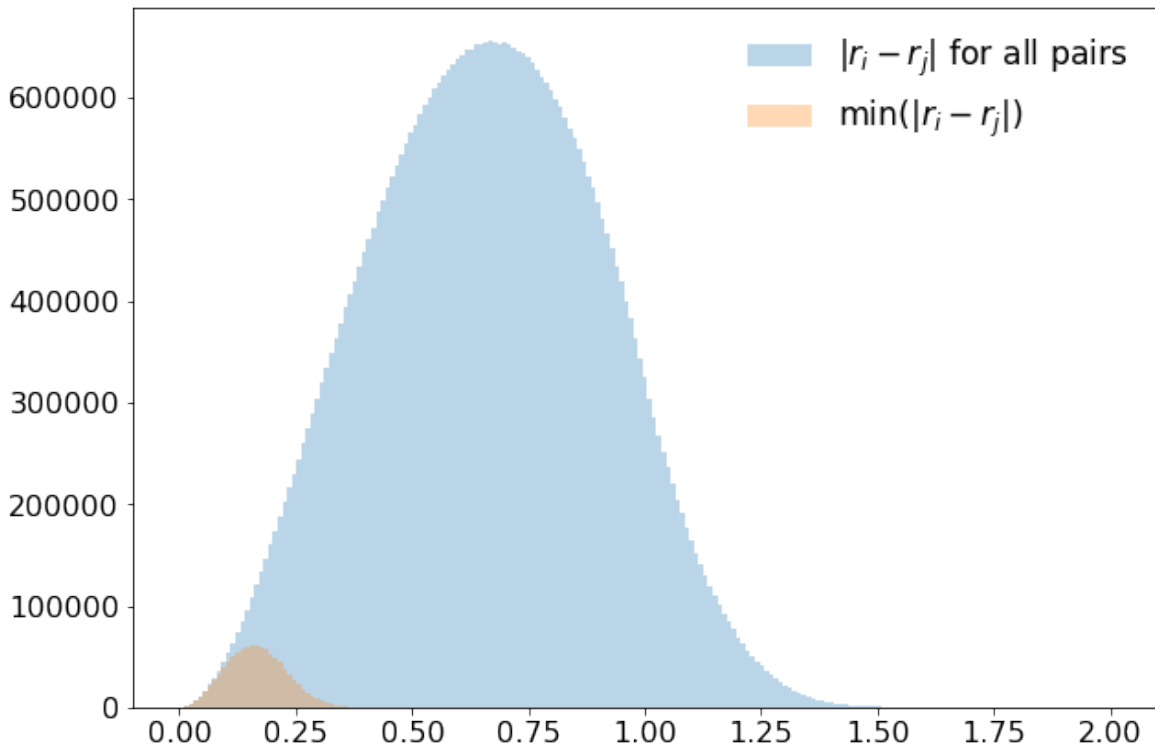
```
selected_pairs = combs_nd(selected_r, n=2, axis=1)
```

And compute the distances, but replacing back the `np.nan` by a default value that will not be seen on a plot.

```
p1, p2 = pairs[:, :, 0, :], pairs[:, :, 1, :]
dp = np.sum((p1-p2)**2, axis=2)**0.5
dp[np.isnan(dp)] = 999
```

And plotting the distributions of both all distances and minimum distances for pairs made out of points verifying $x > y$:

```
plot_style = {'bins': np.linspace(0, 2, 200), 'alpha':0.3}
plt.hist(dp.flatten(), label='|ri-rj| for all pairs', **plot_style)
plt.hist(np.min(dp, axis=1), label='min(|ri-rj|)', **plot_style)
plt.legend();
```



4.6 Some comments

Manipulating numpy array is quite powerful and fast for both computation and plotting, at the condition that we use numpy optimization, namely vectorization, indexing and broadcasting. This is often possible when this has also some limitations as we saw above. Namely, we need to play a bit with “patchwork approaches” to

achieve what we want without loops in the last two sections. Typically, what will work for one computation will not work for another (replacing rejected values by 0 works for an addition and not for a product). For the pairing as well, we had to replace all rejected values by `np.nan` in order to filter them later on. This kind of practice makes things less readable when complexity increases - according to me. Or maybe there are smarter ways to do things.

Chapter 5

Introduction to image processing

Image processing has an important role in data science and in science in general. The typical digit recognition problem is one (classic) example of image processing. The notion of *convolutional neural networks* (CNN) is also a key point in image processing based on machine learning algorithm. Another one, more recent, is the *generative adversarial neural networks* (GAN) which are able to *generate* image of a given nature, after being properly trained. You can check out <https://thispersondoesnotexist.com> which shows generated image of people ... which doesn't exist.

The very first step is to understand how an image is encoded in numpy and how to manipulate it - even without talking about sophisticated algorithms. This is the goal of this notebook, which is split into three different sections:

1. **Basic investigations:** load/plot/write an image, get image histogram, grey scale, cropping, ...
2. **Numerical operations:** addition, subtraction, masking some pixel based on a given condition, ...
3. **Applying basic filters:** image split in blocks versus windows, blurring, sharpening, edge detection, ...

Note that there are few python packages dedicated to image processing, such as [Pillow](#) or [scikit-image](#). The package [scipy](#) also has a module [ndimage](#) dedicated to image processing (cf. this [online lecture](#)). I choose to not use these tools here in order to not increase the number of libraries (very easy to do in python), so *only NumPy and matplotlib will be used in this chapter*. However, if you are interested in doing intensive image processing, I would recommend to look at pillow. Another tool, more oriented toward machine learning and computer vision, is [OpenCV](#) - good to keep in mind depending on your applications.

5.1 Basic investigations

An image is a numpy array of 8-bits integers with a shape ($N_x, N_y, 3$) with N_x (N_y) pixels in the x (y) direction and three colors being an integer in [0, 255] interval (RGB). The low pixel values correspond to dark areas while values close to 255 are clear pixels. An usual image format (png, jpg, ...) can be loaded as numpy array using `plt.imread()` function:

```
# Usual import
import numpy as np
import matplotlib.pyplot as plt

# Load a test image
im = plt.imread('../data/image_test.jpg')
```

Indeed, we can investigate the numpy array we loaded:

```
# Print characteristics
print('Object      : {}'.format(type(im)))
print('Shape       : {}'.format(im.shape))
print('Data type   : {}'.format(im.dtype))

# Print the values of four first pixels along x (for y=0):
print('pixels(x<5, y=0): \n{}'.format(im[0:4, 0, :]))
```

```
Object      : <class 'numpy.ndarray'>
Shape       : (3008, 4008, 3)
Data type   : uint8
pixels(x<5, y=0):
[[18 18  8]
 [21 21  9]
 [31 31 19]
 [29 29 17]]
```

5.1.1 Plotting

The obvious first thing we want to do with an image is to see it! This can be achieved using `plt.imshow()` function. Below, we write a function which uses `plt.imshow()`, with arbitrary figure size keeping the figure ratio:

```
def plot_image(im, h=5, **kwargs):
    lx, ly = im.shape[:2]
    w = (ly/lx)*h
    plt.figure(figsize=(w,h))
    plt.imshow(im, interpolation=None, **kwargs)
    return
```

```
plot_image(im, h=7)
```



5.1.2 Histograms

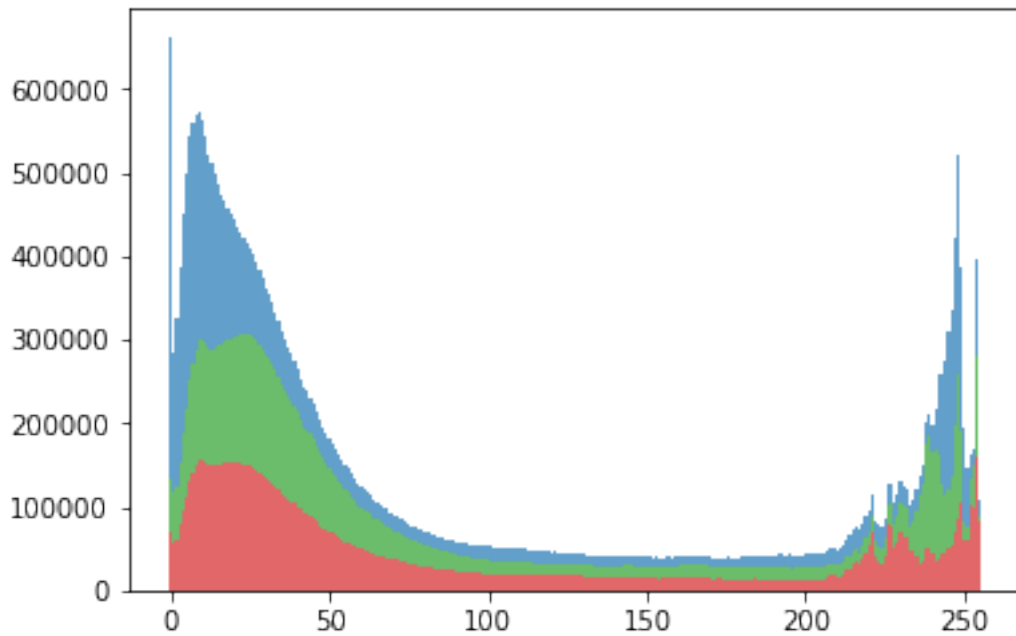
The histogram of an image is often shown on a camera or on post-processing picture software. This allow to appreciate how all pixels are distributed in term of intensity. This can be obtained with a rather straightforward function:

```
def plot_histogram(image):

    # Get the 2D array for each of the three colors, and flat it.
    pixels = [p.ravel() for p in np.array_split(image, 3, axis=2)]

    # Produce the histogram for each color and stack them
    style_hist = {'bins': np.arange(-0.5, 256.5, 1.0), 'alpha': 0.7, 'stacked':True}
    plt.hist(pixels, color=['tab:red', 'tab:green', 'tab:blue'], **style_hist)
    return

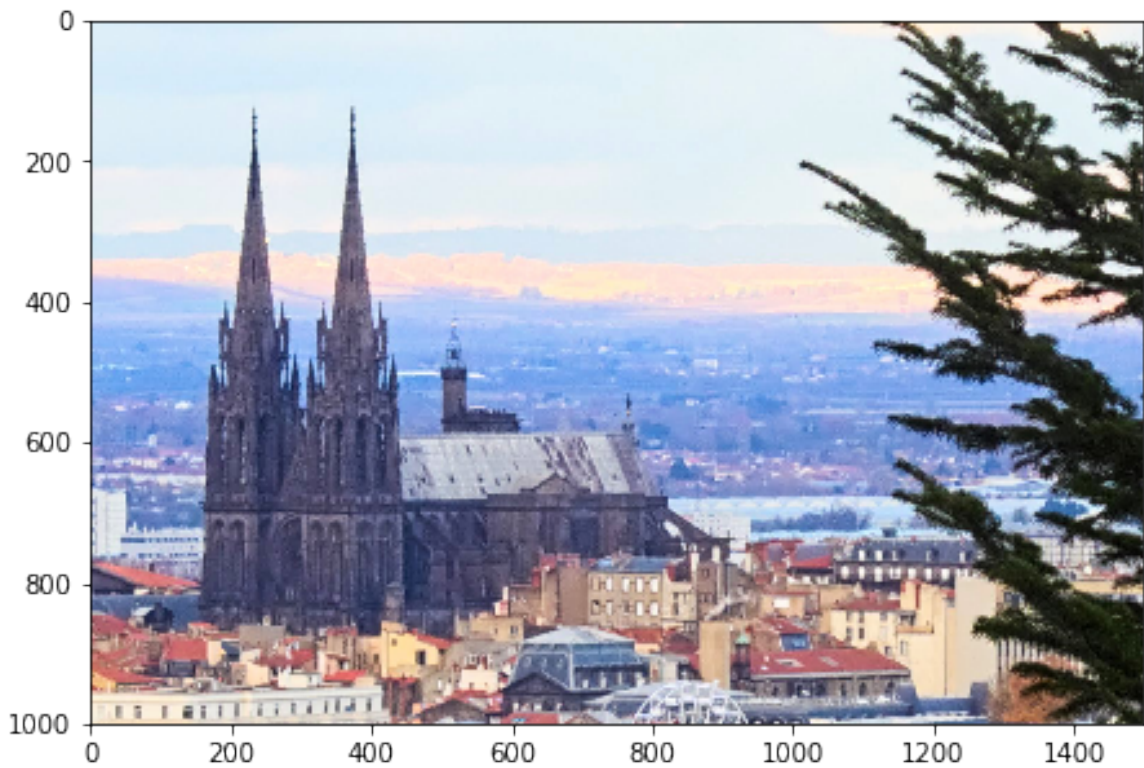
plot_histogram(im)
```



5.1.3 Color and gray scale

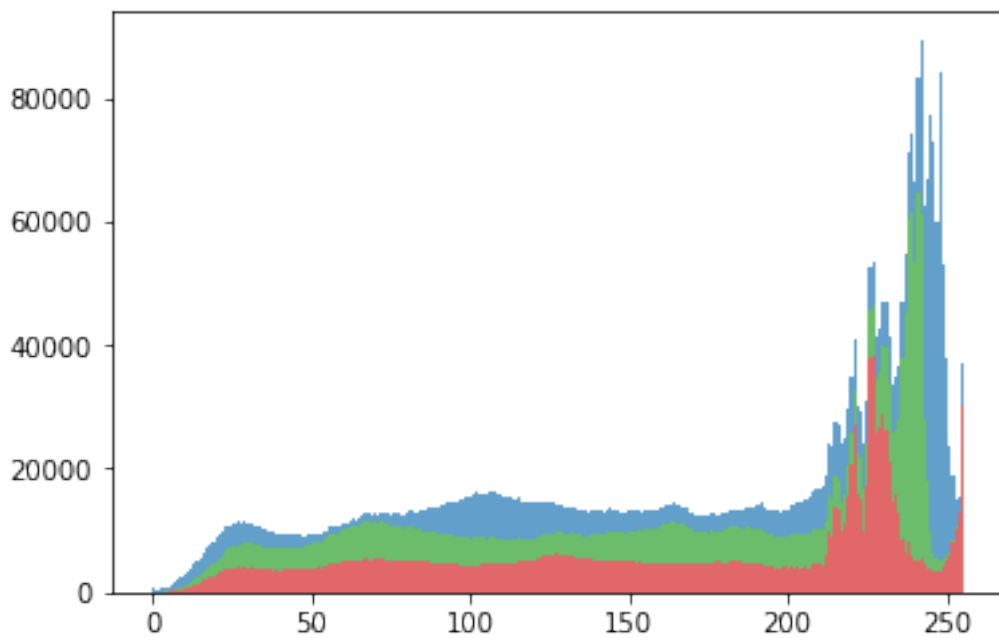
This is also possible to plot each channel (color) separately. First, we can crop the picture to focus on the interesting part of it: the Clermont-Ferrand cathedral. Since the image is a numpy array, cropping is just taking a sub-array using numpy indexing:

```
image = im[500:1500, 1500:3000]
plot_image(image)
```



For instance, we can plot the histogram of this new image and check that the dark parts corresponding to surrounding trees (low values) are significantly decreased:

```
plot_histogram(image)
```



In order to investigate how the colors are spatially distributed, one can plot each channel separately using the appropriated color map. This is what the next function does:

```

def plot_RGB(image):
    '''Plot each color channel'''

    # Get each color channel
    R, G, B = image[:,0], image[:,1], image[:,2]

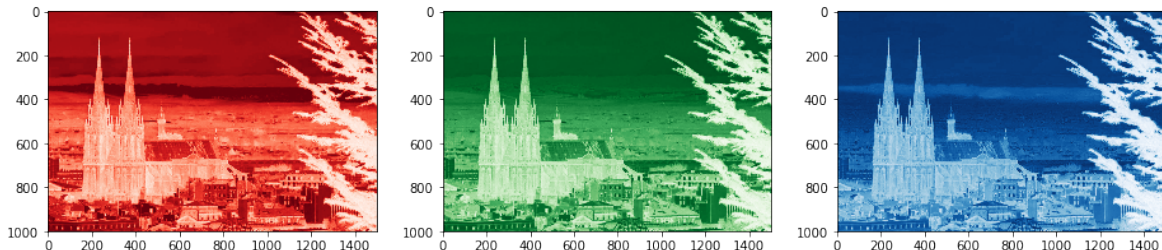
    # Figure shape preserving the image ratio
    lx, ly = image.shape[:2]
    w = (ly/lx)*3
    fig = plt.figure(figsize=(w*3.5,5))

    # One subplot per channel (Nrow, Ncol, Nplot)
    for i, (pixel, color) in enumerate(zip([R, G, B], ['Reds', 'Greens', 'Blues'])):
        plt.subplot(1, 3, i+1)
        plt.imshow(pixel, interpolation=None, cmap=color)

    return

plot_RGB(image)

```



Another usual operation is to switch from colored to gray scale picture. This can be done in several ways (check for e.g. the corresponding [wikipedia article](#)), but one which is relatively simple to implement is based on luminance preservation:

```

def get_gray_scale(image):

    # Get RGB individual values
    R, G, B = image[:,0], image[:,1], image[:,2]

    # Get gray scale from RGB colors: PIX = 0.299 R + 0.587 G + 0.114 B
    pixels = np.array(0.299*R + 0.587*G + 0.114*B, dtype=np.uint8)

    # Replace each channel by this gray scale
    im_gs = np.stack([pixels, pixels, pixels], axis=2)

    # Retrun the gray image
    return im_gs

```

```

# Get the gray scale image
gray_image = get_gray_scale(image)

```

```
# Plot the result
plot_image(gray_image, h=7)
```

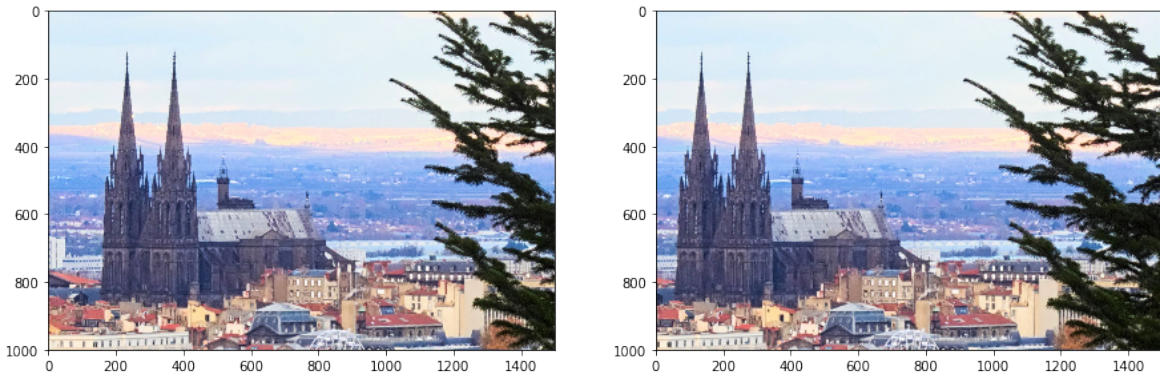


5.2 Numerical operations on images

Since images are numpy arrays, we can perform numerical operations very easily. Not all of them have a proper meaning though, but it is interesting to explore the possibilities. First, we define two images which are the Clermont-Ferrand cathedral slightly shifted:

```
# Create the two images
image1 = im[500:1500, 1500:3000]
image2 = im[500:1500, 1600:3100]

# Plot the two images side-by-side
plt.figure(figsize=(15, 5))
plt.subplot(1, 2, 1)
plt.imshow(image1)
plt.subplot(1, 2, 2)
plt.imshow(image2);
```



5.2.1 Addition & subtraction

What if we add or subtract these pictures? One has to define what happens if the sum (or the difference) is out of the permitted range [0, 255]. Let's take the following convention: if the pixel is below 0, we set it to 0 and if it is above 255, we set it to 255. This can be done by taking the image with float (allowing above values) and operate the truncation after end. The two following functions implement this “image addition/subtraction”:

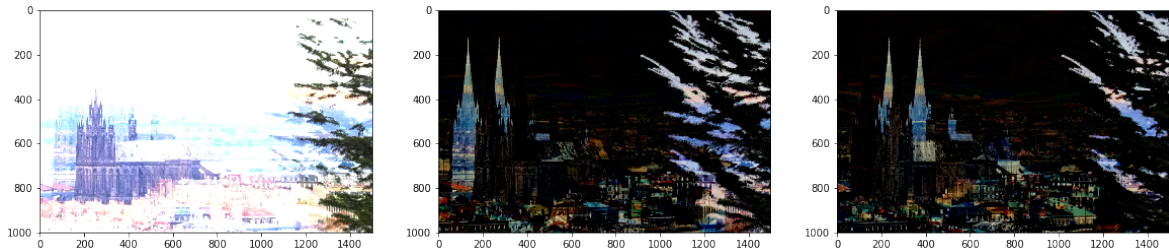
```
def add_pictures(im1, im2):
    s = im1.astype(np.float) + im2.astype(np.float)
    s[s>255] = 255
    s[s<0] = 0
    return s.astype(np.uint8)

def subtract_pictures(im1, im2):
    s = im1.astype(np.float) - im2.astype(np.float)
    s[s>255] = 255
    s[s<0] = 0
    return s.astype(np.uint8)
```

Let's try to plot the added and subtracted images:

```
# Perform the operations
s12 = add_pictures(image1, image2)
d12 = subtract_pictures(image1, image2)
d21 = subtract_pictures(image2, image1)

# Plot the results side-by-side
plt.figure(figsize=(20, 5))
plt.subplot(1, 3, 1)
plt.imshow(s12)
plt.subplot(1, 3, 2)
plt.imshow(d12)
plt.subplot(1, 3, 3)
plt.imshow(d21);
```



When we sum the two picture, we get something very bright (as expected) and we see the echo of the cathedral. After the subtraction (middle), we still see the echo but we get something every dark. The last plot show the other difference, looking quite cool especially at the bottom of the Cathedral!

5.2.2 Modifying certain pixels

Another useful operation we can easily do with NumPy is to mask pixel satisfying a given condition. Let say we want to mask all pixel which as its red level higher than it's blue level summed to the green level:

```
# Get the copy of colors (to be modified latter)
r, g, b = image[...,0].copy(), image[..., 1].copy(), image[..., 2].copy()

# Get the mask
th = add_pictures(g, b)
to_black = r>=th

# Perform the mask
r[to_black] = 0
g[to_black] = 0
b[to_black] = 0
```

One can also decide to set to white the too dark regions, say $r+g+b \leq 60$, without touching the previous pixels:

```
# Get the new indices to set to white
to_white = r+b+g<=60

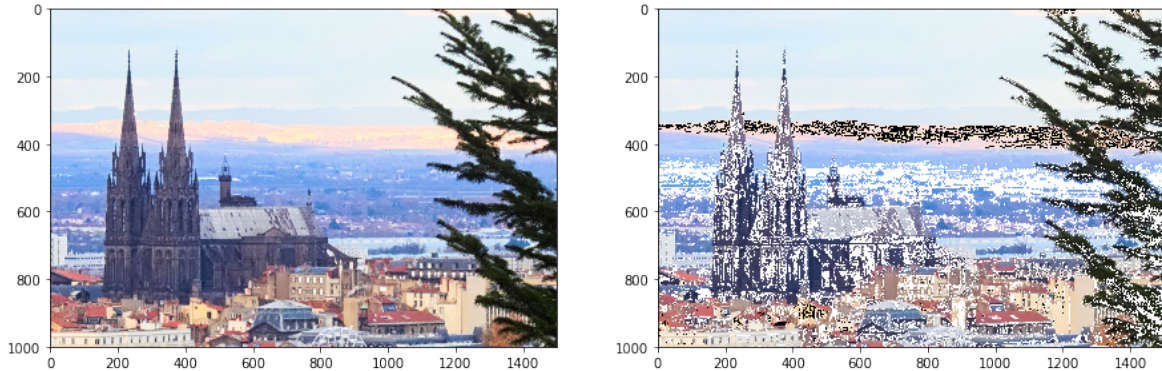
# White only pixel that were didn't touch by the previous mask
to_white = to_white * ~to_black

# Perform the whitening
r[to_white] = 255
g[to_white] = 255
b[to_white] = 255

# Combine the three colors together
image_masked = np.stack([r, g, b], axis=2)

# Plot the results side-by-side
plt.figure(figsize=(15, 5))
```

```
plt.subplot(1, 2, 1)
plt.imshow(image1)
plt.subplot(1, 2, 2)
plt.imshow(image_masked);
```



5.2.3 Modifying regions

We can use the fancy indexing of numpy to define a shape and modify only pixels which are within this shape. For the example, let's consider a circle with a given center position $r_0 = (x_0, y_0)$ and radius R in which we will decrease the luminosity - i.e scale down all pixels together. To start, let's write a function which return a boolean 2D array which tell us whether a give (x, y) -pixel is in the circle or not. This will, again, involve a meshgrid:

```
def idx_in_circle(im, x0, y0, R):
    lx, ly = im.shape[:2]
    X, Y = np.meshgrid(np.arange(0, ly), np.arange(0, lx))
    radius = ((X-x0)**2 + (Y-y0)**2)**0.5
    return radius <= R
```

We will now use this function to change to add 10 random shadowed circles that we generate using `np.random.randint()` function for the center and the radius of circles. Note the use packing/unpacking of arguments to call the `idx_in_circle()` function in a more concise and clear way (together with the `zip()` syntax):

```
# Copy original image and modify it
result = image.copy()

# Get 10 random circles (in position and radius)
x0 = np.random.randint(low=100, high=1000, size=10)
y0 = np.random.randint(low=100, high=1500, size=10)
R0 = np.random.randint(low=50, high=300, size=10)

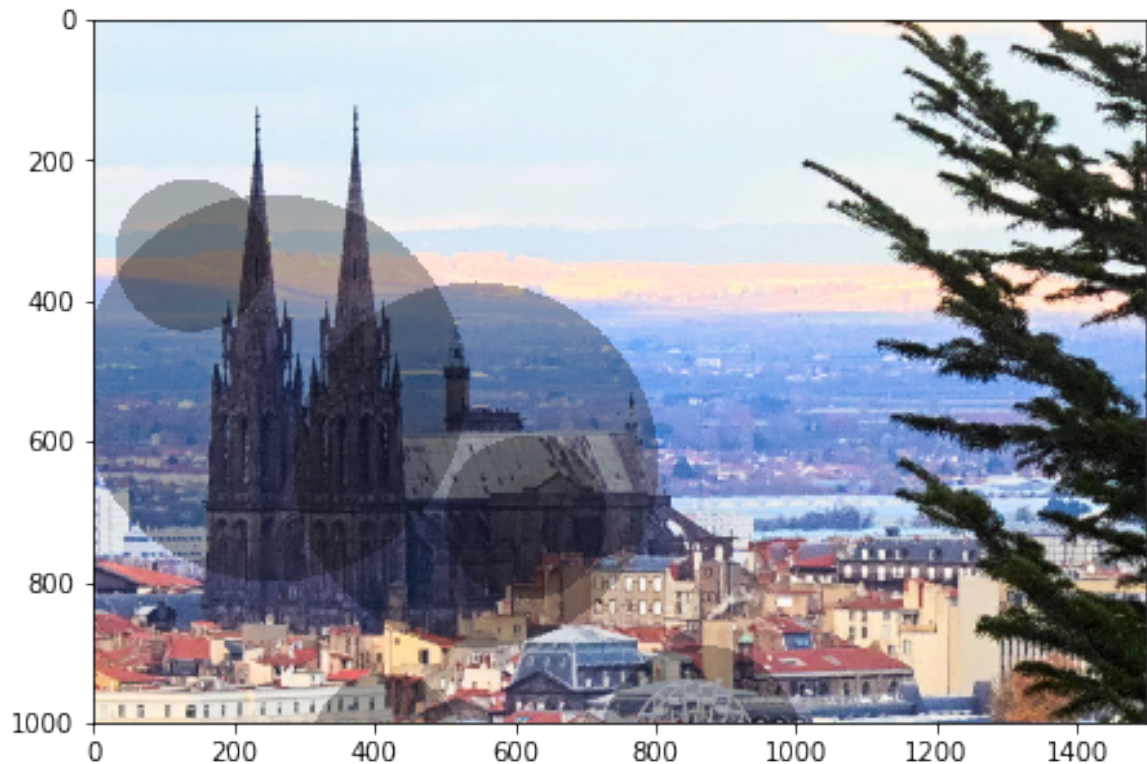
# For each of them, modify the picture
for circle in zip(x0, y0, R0):
```

```

idx = idx_in_circle(image, *circle)
result[idx] = result[idx]*0.7

# Plot the result
plt.figure(figsize=(10, 5))
plt.imshow(result);

```



5.3 Image filters with NumPy

5.3.1 Kernels, image blocks v.s. windows

In image processing, a filter is a small 2D array $n \times n$ (also called *kernel*) which is used to modify the value of each pixel using a *convolution* between a portion of the image and the kernel. These portions can be either use every pixel only once - split the image in $(n \times n)$ blocks - or they can use every pixel several times - sliding $(n \times n)$ overlapping windows. The mathematical operation behind the name convolution is a simple sum over all elements from the window, weighted by the elements of the kernel.

In order to better understand the concept of kernels, blocks and windows, let's now take an example of a 12x12 image and build up both the blocks and sliding windows:

```

# Image definition
image = np.arange(12*12).reshape(12, 12)
print('image = \n{}'.format(image))

```

```

# Build-up 3x3 independant blocks
step3 = range(0, 12, 3)
blocks_3x3 = np.array([image[i:i+3, j:j+3] for i in step3 for j in step3])
blocks_3x3 = blocks_3x3.reshape(4, 4, 3, 3) # Organize the 16 blocks into a 4x4 grid
print('\nBlock[1, 1] = \n{}'.format(blocks_3x3[1, 1]))


# Built-up 3x3 windows for pixel far from the border (to avoid technical issues)
windows_3x3 = np.array([image[i-1:i+2, j-1:j+2] for i in range(1, 11) for j in range(1, 11)])
windows_3x3 = windows_3x3.reshape(10, 10, 3, 3) # Organize the 100 blocks into a 10x10 grid
print('\nWindow[3, 2] = \n{}'.format(windows_3x3[3, 2]))


image =
[[ 0   1   2   3   4   5   6   7   8   9   10  11]
 [ 12  13  14  15  16  17  18  19  20  21  22  23]
 [ 24  25  26  27  28  29  30  31  32  33  34  35]
 [ 36  37  38  39  40  41  42  43  44  45  46  47]
 [ 48  49  50  51  52  53  54  55  56  57  58  59]
 [ 60  61  62  63  64  65  66  67  68  69  70  71]
 [ 72  73  74  75  76  77  78  79  80  81  82  83]
 [ 84  85  86  87  88  89  90  91  92  93  94  95]
 [ 96  97  98  99  100 101 102 103 104 105 106 107]
 [108 109 110 111 112 113 114 115 116 117 118 119]
 [120 121 122 123 124 125 126 127 128 129 130 131]
 [132 133 134 135 136 137 138 139 140 141 142 143]]


Block[1, 1] =
[[39 40 41]
 [51 52 53]
 [63 64 65]]


Window[3, 2] =
[[38 39 40]
 [50 51 52]
 [62 63 64]]

```

For instance, the number 39 can only be on a edge of a block (used once) while it can be everywhere for the sliding windows (used several times). Let's now define a 3x3 kernel and apply it to `blocks_3x3[1, 1]`:

```

# Definition
kernel = np.arange(9).reshape(3, 3)/20
print('kernel = \n{}'.format(kernel))

# Convolution with the block[1, 1]
this_block = blocks_3x3[1, 1]
new_pixel = np.sum(kernel * this_block)
print('\nProduct of elements = \n{}'.format(kernel * this_block))
print('\nNew pixel = {:.1f} (vs an old pixel of {})'.format(new_pixel, this_block[1, 1]))

```

```

kernel =
[[0.  0.05 0.1 ]
 [0.15 0.2  0.25]
 [0.3  0.35 0.4 ]]

Product of elements =
[[ 0.    2.    4.1 ]
 [ 7.65 10.4  13.25]
 [18.9  22.4  26.  ]]

New pixel = 104.7 (vs an old pixel of 52)

```

Let's now apply the kernel defined above to both blocks and sliding windows. We can also represent the image before filter, after block-based filter and window-based filter.

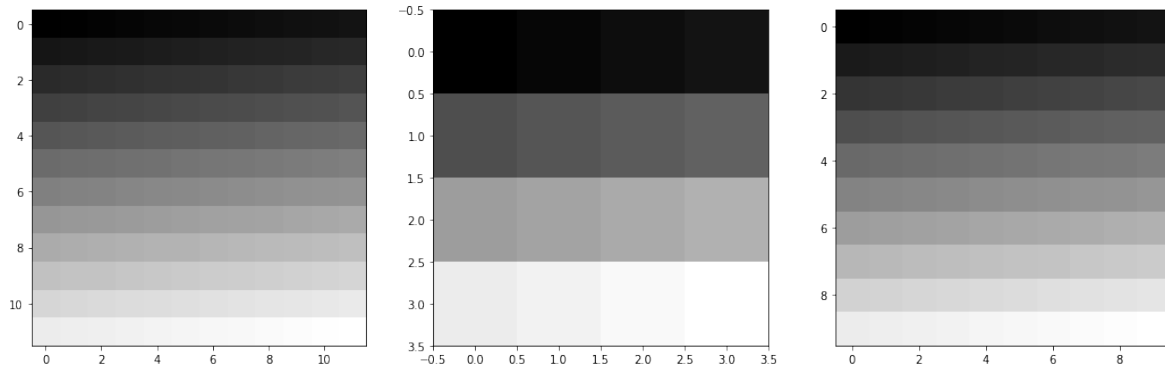
```

# Apply block-based filter
block_filtered = np.sum(blocks_3x3*kernel[np.newaxis, np.newaxis], axis=(2, 3))
block_filtered[block_filtered>255]=255
block_filtered[block_filtered<0]=0

# Apply window-based filter
window_filtered = np.sum(windows_3x3*kernel, axis=(2, 3))
window_filtered>window_filtered>255]=255
window_filtered>window_filtered<=0]=0

# Plot the results side-by-side
plt.figure(figsize=(18, 7))
plt.subplot(1, 3, 1)
plt.imshow(image, cmap='gray')
plt.subplot(1, 3, 2)
plt.imshow(block_filtered, cmap='gray')
plt.subplot(1, 3, 3)
plt.imshow(window_filtered, cmap='gray');

```



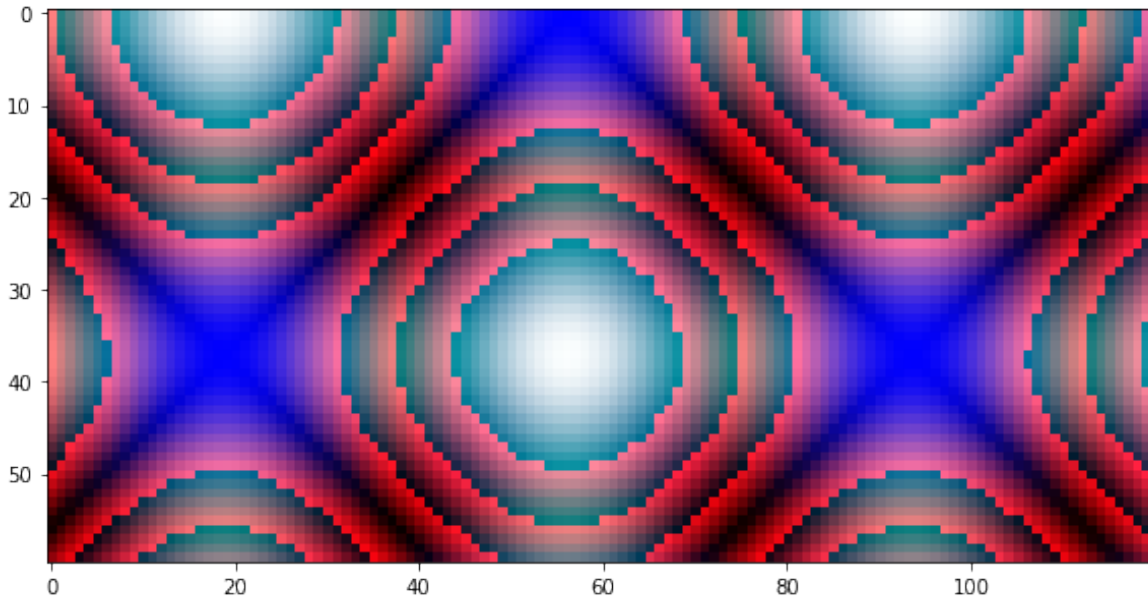
Important comment. The block view is not too gridy in memory but the the windows view can explode quite rapidly. Indeed, for a kernel of $n \times n$, the windows view is n^2 larger than the original array. If you manipulate millions of images this can be problematic. For this reason, we will use a buildin `scipy.signal` function to

perform the “sliding windows application”, called `convolved2D()`. However in the next section, we will study how efficiently perform the “block approach” using a deeper numpy feature: the strides.

Generalisation to RGB image. Before moving forward, we need to consider the 3 colors of an image to apply a filter, which has some implication in term of broadcasting structure. First, let’s define a dummy RGB image using a meshgrid (careful x and y are reversed wrt `imshow`) and three function for each color:

```
# Create a shaped image
def get_dummy_image(nx=600, ny=1200):
    X, Y = np.meshgrid(np.linspace(0, 10, ny), np.linspace(0, 5, nx))
    f = lambda n: np.abs(np.sin(X)**n+np.cos(Y)**n)
    im = np.stack([2*f(1), 0.5*f(3), 0.5*f(2)], axis=2)*255
    return im.astype(np.uint8)

im = get_dummy_image(nx=60, ny=120)
plot_image(im)
```



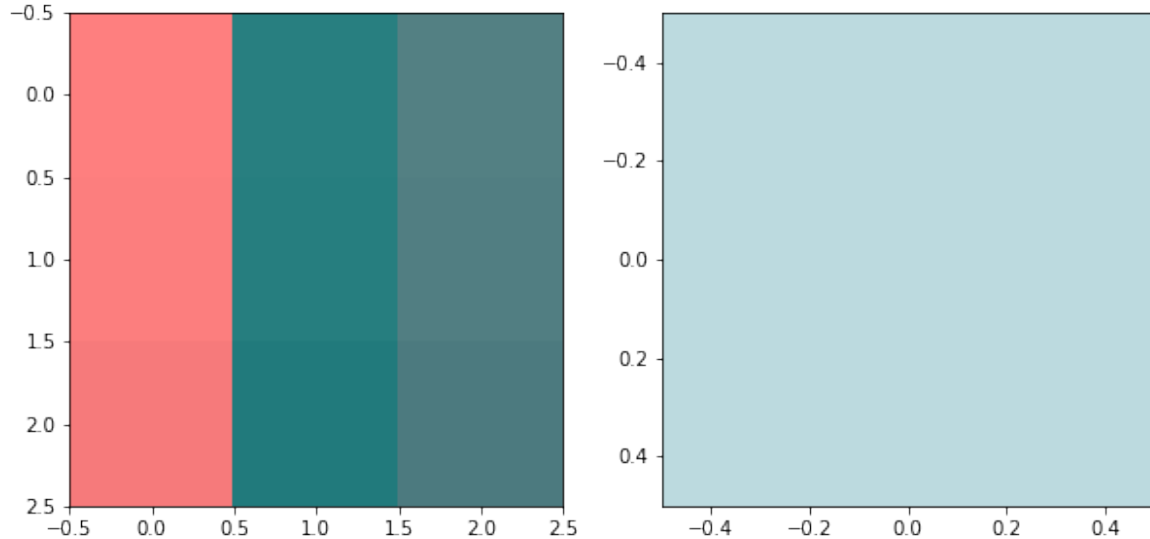
In order to get the proper broadcasting, we need to extend the kernel with a third dimension for the colors, which can be done *via* the syntax `kernel[:, :, np.newaxis]`. In that way, the kernel will be automatically duplicated for each color and its application can be properly vectorized.

```
# Kernel application with the proper broadcasting over colors only for the first block
kernel = np.arange(9).reshape(3, 3)/20
new_pixel = np.sum(im[0:3, 0:3, :]*kernel[:, :, np.newaxis], axis=(0,1), dtype=np.uint8)
print(new_pixel)

# Plotting the 9 considered pixels and the result
plt.figure(figsize=(10, 5))
plt.subplot(1, 2, 1)
plt.imshow(im[0:3, 0:3, :])
```

```
plt.subplot(1, 2, 2)
plt.imshow(new_pixel.reshape(1, 1, 3));
```

[188 218 223]



5.3.2 Image blocks: intuitive but inefficient approach

The most intuitive approach is to apply the filter to each block, involving an explicit *loop* over all the blocks of the image. Let's follow this approach for now, defining a function which apply the kernel to the block indexed by (i, j) . Note that we didn't handle properly the boundaries, *i.e.* if the size of the image is not exactly n times the size of the kernel.

```
def apply_filter_one_block(im, kn, i, j):
    dx, dy = kn.shape
    start_i, end_i = i*dx, (i+1)*dx
    start_j, end_j = j*dy, (j+1)*dy
    indices = (slice(start_i, end_i), slice(start_j, end_j), slice(None, None))
    pixel = np.sum(im[indices].astype(np.float)*kn[:, :, np.newaxis], axis=(0, 1))
    pixel[pixel<0]=0
    pixel[pixel>255]=255
    return pixel.astype(np.uint8)

# Testing with a dummy image and averaging filter (1 everywhere)
im = get_dummy_image(nx=12, ny=12)
kernel = np.ones(shape=(3, 3))/9.
print('new pixel = {}'.format(apply_filter_one_block(im, kernel, 1, 1)))

new pixel = [98 57 79]
```

Let's now try to applyt the strategy to a real image 1200×1200 with the two kernel sizes 3×3 and 6×6 :

```

# Testing with a real image
image_full = plt.imread('../data/image_test.jpg')
im = image_full[500:1700, 1500:2700]

# 3x3 kernel with all ones
kernel = np.ones(shape=(3, 3))/(3*3)
im3x3 = np.array([[apply_filter_one_block(im, kernel, i, j) for j in range(0, 400)] for i in
                  range(0, 400)])

# 6x6 kernel with all ones
kernel = np.ones(shape=(6, 6))/(6*6)
im6x6 = np.array([[apply_filter_one_block(im, kernel, i, j) for j in range(0, 200)] for i in
                  range(0, 200)])

# 12x12 kernel with all ones
kernel = np.ones(shape=(12, 12))/(12*12)
im12x12 = np.array([[apply_filter_one_block(im, kernel, i, j) for j in range(0, 100)] for i in
                     range(0, 100)])

# Plotting the result
fig = plt.figure(figsize=(40, 10))
for i, this_im in enumerate([im, im3x3, im6x6, im12x12]):
    plt.subplot(1, 4, i+1)
    plt.imshow(this_im)

```



This is also possible to time the loop operation with the %timeit magic command. Let's do it in the full picture:

```

im_test = image_full[:3006, :4008]
kernel = np.ones(shape=(3, 3))/(3*3)
%timeit np.array([[apply_filter_one_block(im_test, kernel, i, j) for j in range(0, 1336)] for
                  i in range(0, 1002)])

```

19.6 s ± 1.27 s per loop (mean ± std. dev. of 7 runs, 1 loop each)

It takes then **~20s** to process a 12 Mpixels image. This shows that this approach is way too long for a systematic treatment, especially if you think of a larger number of image to process. This was expected since we already mentioned that explicit loops in python are just too slow. The goal of the next section is to use the power of numpy to remove the loop and speed-up this computation.

5.3.3 Image blocks : fast numpy-based approach

The idea is to first turn the image array from the dimension ($N_x, N_y, 3$) to ($N_{x_new}, N_{y_new}, N_{x_kernel}, N_{y_kernel}, 3$), where (N_{x_new}, N_{y_new}) is the dimension of the “image of blocks”. Once this is done, one can simply multiply and sum over the axis 2 and 3.

The other approach is quite advance but also quite powerful: `numpy.lib.stride_tricks.as_strided()`. Strides are basically a tuple of bytes of memory to jump from one element to another in each dimension. In other words, it’s the byte-separation between consecutive items for each dimension. This bytes manipulation doesn’t duplicate the data but rather view them as a different way, which is much more efficient from the memory point of view. This is the approach behind the broadcasting and is, by far, *the fastest approach*. Let’s go step by step to understand the strides concept.

```
# Small image as a copy of a sub-image from our favorite test
im_small = image_full[1000:1006, 1000:1006].copy()

# Print few information, including the number of bytes 6x6x3x1 = 108
print('Shape      : {}'.format(im_small.shape))
print('Dtype      : {}'.format(im_small.dtype))
print('Item size  : {}'.format(im_small.itemsize))
print('Nbytes    : {}'.format(im_small.nbytes))
```

```
Shape      : (6, 6, 3)
Dtype      : uint8
Item size  : 1
Nbytes    : 108
```

Reminder: 1 byte (or octet) is 8 bits. One bit is a single number being 0 and 1. The 8-bits image we are looking at in this lecture is then 1 byte per color and per pixel. The above number then makes perfect sense.

Let’s try to compute the byte jump by hand first. In the third axis (color) axis, the jump between two consecutive items is just the item size so 1 byte. For the y-axis you jump 3 by 3 elements to jump over all colors and reach the next position. This leads to a byte jump of 3 bytes. Finally, to jump over the next x-axis value, one needs to loop over all y-values and 3 colors, which makes $6 \times 3 = 18$. So the `im.strides` command should give (18, 3, 1).

```
print(im_small.strides)
```

```
(18, 3, 1)
```

Now, we can manipulate these jumps to organize the numbers differently. Let’s do it for one color only and just 4×4 array, to have a smaller array and follow numbers individually.

```
# 4x4 array and getting strides
a = im_small[:4, :4, 0].copy()
jumps = a.strides
print('a = \n{}'.format(a))
print('strides = {}'.format(jumps))
```

```
a =
[[124 120 114 111]
 [123 119 115 112]
 [122 119 115 113]
 [120 118 116 115]]
strides = (4, 1)
```

Let's say that we want to make an array of 2x2 blocks each 2x2 elements, so a new shape of (2, 2, 2, 2). The first block would look like:

```
[124, 120]
[123, 119]
```

In term of strides, this means that the memory jump between two elements along the before-the-last axis is equivalent to a jump of 4 bytes. This corresponds to the jump between two elements along the first axis of the original array. In order to have the second block like:

```
[114 111]
[115 112]
```

One needs to jump along the second axis by 2 bytes, which is the size of the jump to go from 124 to 114. Finally, for a jump along the first axis, we need to go from 124 to 122, which account for 8 bytes (corresponding to the two first lines of a).

```
# Re-agencement using the strides
new_shape = (2, 2, 2, 2)
new_jumps = (8, 2, 4, 1)

# Create the new array
from numpy.lib.stride_tricks import as_strided
a_new = as_strided(a, strides=new_jumps, shape=new_shape)

# Print the result
print('a_new = \n{}'.format(a_new))
print('strides = {}'.format(new_jumps))
```

```
a_new =
[[[[124 120
    [123 119]

    [[114 111]
     [115 112]]]

    [[[122 119]
     [120 118]]

    [[[115 113]
     [116 115]]]]]

strides = (8, 2, 4, 1)
```

Once we did this by hand, we can try to automatize it given the size of the blocks we want to make and the size of the image. Let's take the example of (100, 100) image with a (3, 3) block:

```
# 100x100 image
im100x100 = image_full[:100, :100, 0].copy()

# Converting each shape into numpy array for element-wise operation
im_shape = np.array(im100x100.shape) # image dimensions
bl_shape = np.array((3, 3))      # block dimensions

# Shape of the "image of blocks", as floor division
im_blocks_shape = im_shape // bl_shape
print(im_blocks_shape)

# Full dimension is a concatenation of corresponding shapes
new_shape = tuple(im_blocks_shape) + tuple(bl_shape)
print(new_shape)
```

```
[33 33]
(33, 33, 3, 3)
```

Now, it is matter to get automatically the new strides knowing the image and block sizes. The jumps corresponding to the two first dimensions are directly the original picture jumps times the block shape. Indeed, we want the following:

- along y-axis (2nd axis): take the 1st element, then the $1+ Ny_block$'s one, etc ... so memory jump is $old_jump_y * Ny_block$.
- along x-axis (1st axis): take the 1st element then the $1+ Nx_block$'s one, so memory jump is also $old_jump_x * Nx_block$

Concerning the two last axis, i.e. the navigation inside a block, this is just the original memory jumps.

```
# Get old strides as numpy array
old_strides = np.array(im100x100.strides)
print(old_strides)

# Form new strides
new_strides = tuple(old_strides*bl_shape) + tuple(old_strides)
print(new_strides)
```

```
[100    1]
(300, 3, 100, 1)
```

```
# Form the blocks and check this is correct for the first few
im100x100_blocked = as_strided(im100x100, strides=new_strides, shape=new_shape,
→ writeable=False)
```

```
# Print the original image
print('Original image: \n{}'.format(im100x100[:6, :6]))

# Print the first few blocks
print('\nFirst 3x3 blocks: \n',format(im100x100_blocked[:2, :2]))
```

Original image:
[[18 18 22 26 23 21]
[21 23 19 21 29 30]
[31 31 17 13 25 24]
[29 24 15 17 28 31]
[25 12 12 19 20 23]
[13 2 12 27 28 38]]

First 3x3 blocks:

[[[[18 18 22]

[21 23 19]

[31 31 17]]]

[[26 23 21]

[21 29 30]

[13 25 24]]]

[[[[29 24 15]

[25 12 12]

[13 2 12]]]

[[[17 28 31]

[19 20 23]

[27 28 38]]]]

We are now ready to build-up a function which apply a kernel per block (accounting for the additional axis for colors - not taken into account above), where the last step is just operating the sum:

```
def apply_filter_strides(image, kernel):
    from numpy.lib.stride_tricks import as_strided

    # Get the new shape
    m_shape, image_shape = np.array(kernel.shape), np.array(image.shape)
    new_shape = tuple(image_shape[:2] // m_shape) + tuple(m_shape) + (image_shape[-1],)

    # Get the new strides
    new_strides = tuple(image.strides[:2] * m_shape) + image.strides

    # Get the new blocked image (Nx_new, Ny_new, Nx_mask, Ny_mask, 3)
    blocked_image = as_strided(image, shape=new_shape, strides=new_strides, writeable=False)
```

```

# Apply the mask with the proper broadcasting
kernel_reshaped = kernel[np.newaxis, np.newaxis, :, :, np.newaxis]
result = np.sum(blocked_image*kernel_reshaped, axis=(2, 3))

# Cleaning
result[result<0] = 0
result[result>255] = 255

# Return
return result.astype(np.uint8)

```

It is worth to mention that all this code is already written in some of the tools mentioned at the begining of this chapter, but the goal here is to learn how the tools are made (and possible make new ones!). To finalize the process, we can first compare we get the same result as the explicit loop function, and then compare the timing for those two functions:

```

# Check compatibility with the previous function
im = image_full[500:1700, 1500:2700]
kernel = np.ones(shape=(12, 12))/(12*12)
im12x12_loop = np.array([[apply_filter_one_block(im, kernel, i, j) for j in range(0, 100)]
    for i in range(0, 100)])
im12x12_strides = apply_filter_strides(im, kernel)

# Check that all pixels are the same in both images
print('Are the two results equal? {}'.format(np.all(im12x12_strides==im12x12_loop)))

```

Are the two results equal? True

```

# Execution time
kernel = np.ones(shape=(3, 3))/(3*3)
%timeit apply_filter_strides(image_full, kernel)

```

866 ms ± 6.46 ms per loop (mean ± std. dev. of 7 runs, 1 loop each)

The fully vectorized approach based on stride manipulations is *more than 20 times faster* than the explicit loop.

5.4 Few typical filters

5.4.1 Few utility functions

In order to perfom image processing in a systematic way, we need to build up few helper functions which are written below. They are all based on the above developpements, except the sliding windows application using `scipy.signal.convolve2d()` function. The full documentation of this function, together with some examples, can be found [here](#) and will not be discussed in this lecture.

Note: there is slight difference in the definition of convolve2d() function and the weighted sum used in the blocks approach. The convolve2d() perform the weighted sum using the transposed kernel, which explain the the considered kernel in the below function is kernel.T.

Clean image (value and data type):

```
def clean_image(image):
    image[image < 0] = 0
    image[image > 255] = 255
    return image.astype(np.uint8)
```

Normalise the kernel values:

```
def normalize_filter(kernel):
    if np.sum(kernel) != 0:
        return kernel/np.sum(kernel)
    else:
        return kernel
```

Apply filter with sliding windows (based on scipy.signal 2D convolution function)

```
def apply_filter_windows(image, kernel):
    from scipy.signal import convolve2d

    # Performing 2D convolution on every color
    args = {'mode': 'same', 'boundary': 'fill', 'fillvalue': 0}
    r, g, b = image[:, :, 0], image[:, :, 1], image[:, :, 2]
    filtered_colors = [convolve2d(ch, kernel.T, **args) for ch in [r, g, b]]

    # Stacking all colors together
    filtered_image = np.stack(filtered_colors, axis=2)

    # Result
    return clean_image(filtered_image)
```

Applying filter with blocks (based in stride-based manipulation)

```
def apply_filter_blocks(image, kernel):
    from numpy.lib.stride_tricks import as_strided

    # Get the new shape
    m_shape, image_shape = np.array(kernel.shape), np.array(image.shape)
    new_shape = tuple(image_shape[:2] // m_shape) + tuple(m_shape) + (image_shape[-1],)

    # Get the new strides
    new_strides = tuple(image.strides[:2] * m_shape) + image.strides

    # Get the new blocked image (Nx_new, Ny_new, Nx_mask, Ny_mask, 3)
```

```

blocked_image = as_strided(image, shape=new_shape, strides=new_strides, writeable=False)

# Filtered image
kernel_reshaped = kernel[np.newaxis, np.newaxis, :, :, np.newaxis]
filtered_image = np.sum(blocked_image*kernel_reshaped, axis=(2, 3))

# Result
return clean_image(filtered_image)

```

Test image definition:

```
image = image_full[500:1500, 1500:3000]
```

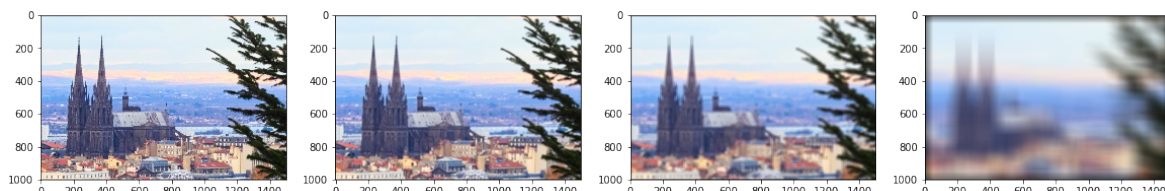
5.4.2 Blurry filter

The blurry filter just perform an average of all surrounding pixels. The corresponding kernel is a constant value in at position, with a sum of 1.0. Below, the blurry filter is applied for both sliding window and bloc approaches. *Caution:* the sliding window for a (100×100) kernel takes several minutes.

```

# Windows application
plt.figure(figsize=(20, 5))
for i, n in enumerate([3, 10, 20, 100]):
    kernel = normalize_filter(np.ones(shape=(n, n)))
    result = apply_filter_windows(image, kernel)
    plt.subplot(1, 4, i+1)
    plt.imshow(result)

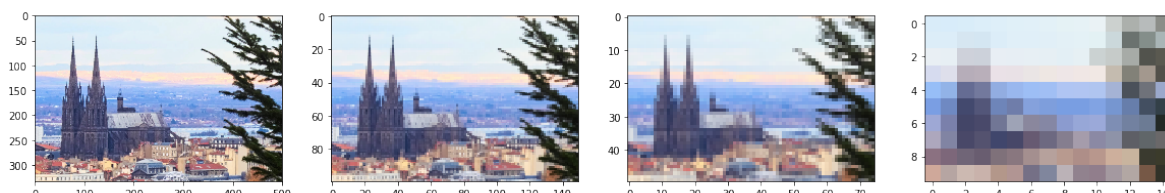
```



```

# Block application
plt.figure(figsize=(20, 5))
for i, n in enumerate([3, 10, 20, 100]):
    kernel = normalize_filter(np.ones(shape=(n, n)))
    result = apply_filter_blocks(image, kernel)
    plt.subplot(1, 4, i+1)
    plt.imshow(result)

```



5.4.3 Edge detection

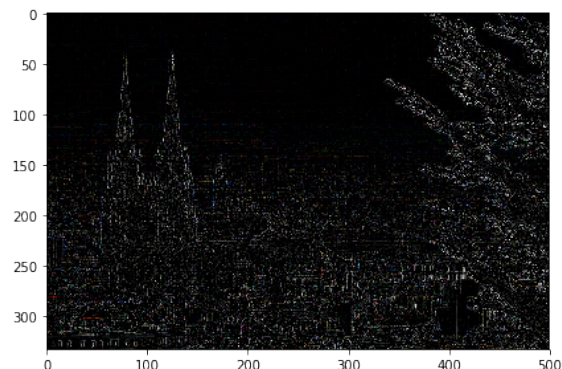
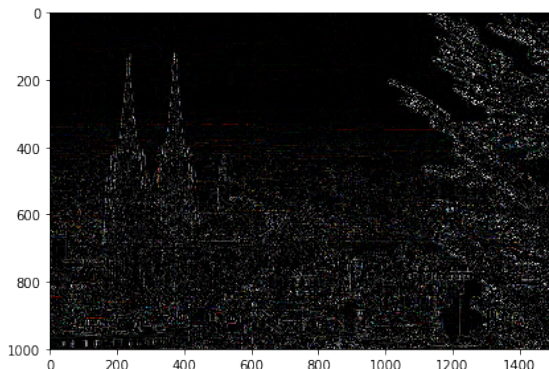
```

filter_border = np.array([
    [-1, -1, -1],
    [-1, 8, -1],
    [-1, -1, -1]
])

filter_border = normalize_filter(filter_border)
border_windows = apply_filter_windows(image, filter_border)
border_blocks = apply_filter_blocks(image, filter_border)

# Plotting the result
plt.figure(figsize=(15, 5))
plt.subplot(1, 2, 1)
plt.imshow(border_windows)
plt.subplot(1, 2, 2)
plt.imshow(border_blocks);

```



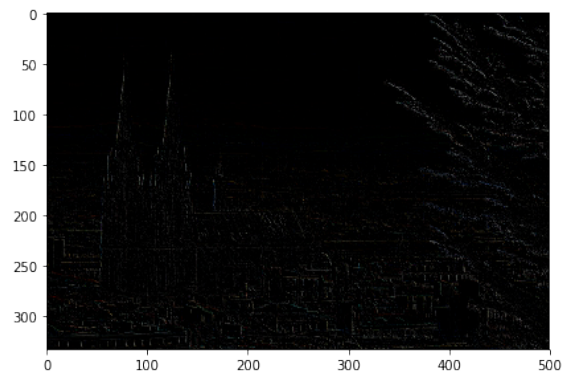
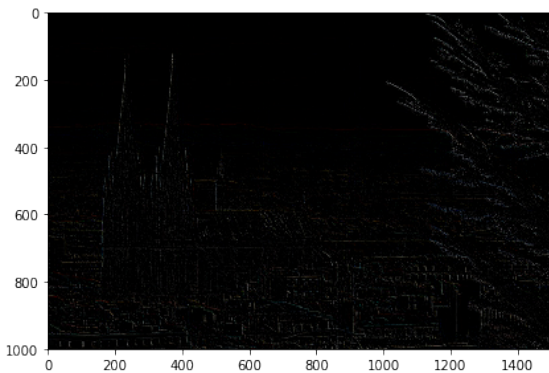
```

# Border which are direction-dependent: top-left
filter_topleft = np.array([
    [0, 0, -1],
    [0, 1, 0],
    [0, 0, 0]
])

filter_topright = normalize_filter(filter_topleft)
border_windows = apply_filter_windows(image, filter_topleft)
border_blocks = apply_filter_blocks(image, filter_topleft)

# Plotting the result
plt.figure(figsize=(15, 5))
plt.subplot(1, 2, 1)
plt.imshow(border_windows)
plt.subplot(1, 2, 2)
plt.imshow(border_blocks);

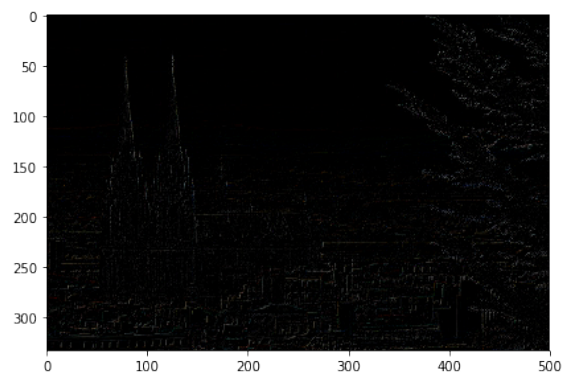
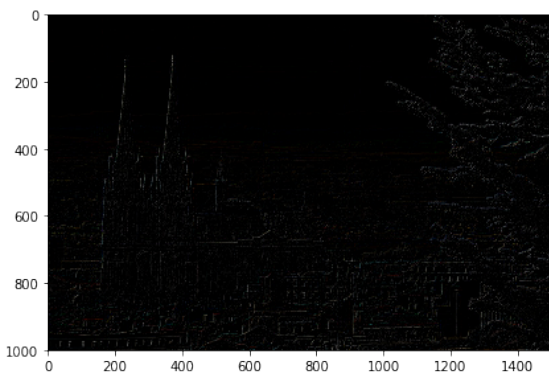
```



```
# Border which are direction-dependent: top-right
filter_topright = np.array([
    [ -1,  0,  0],
    [  0,  1,  0],
    [  0,  0,  0]
])

filter_topleft = normalize_filter(filter_topright)
border_windows = apply_filter_windows(image, filter_topright)
border_blocks = apply_filter_blocks(image, filter_topright) # Still a difference between
→ block/window-approaches

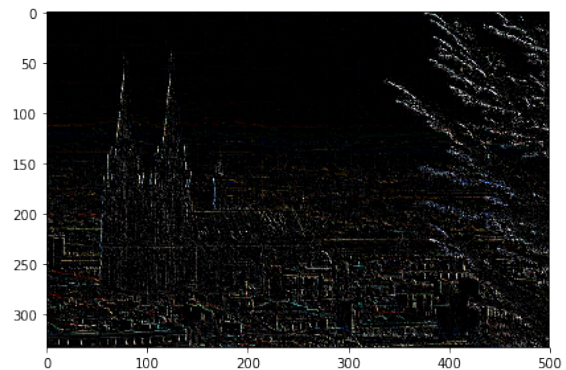
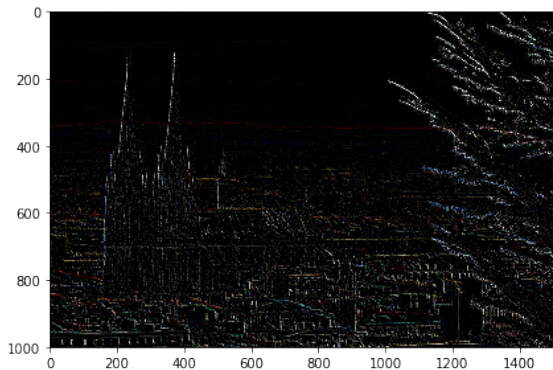
# Plotting the result
plt.figure(figsize=(15, 5))
plt.subplot(1, 2, 1)
plt.imshow(border_windows)
plt.subplot(1, 2, 2)
plt.imshow(border_blocks);
```



```
# Playing with intensity of the border
filter_topleft = np.array([
    [  0,  0, -3],
    [  0,  3,  0],
    [  0,  0,  0]
])
```

```
filter_topleft = normalize_filter(filter_topleft)
border_windows = apply_filter_windows(image, filter_topleft)
border_blocks = apply_filter_blocks(image, filter_topleft)

# Plotting the result
plt.figure(figsize=(15, 5))
plt.subplot(1, 2, 1)
plt.imshow(border_windows)
plt.subplot(1, 2, 2)
plt.imshow(border_blocks);
```



5.4.4 Sharpen filter

```
filter_sharpen = np.array([
    [ 0, -1,  0],
    [-1,  5, -1],
    [ 0, -1,  0]
])

filter_sharpen = normalize_filter(filter_sharpen)
sharpen_windows = apply_filter_windows(image, filter_sharpen)
sharpen_blocks = apply_filter_blocks(image, filter_sharpen)

# Plotting the result
plt.figure(figsize=(15, 5))
plt.subplot(1, 2, 1)
plt.imshow(sharpen_windows)
plt.subplot(1, 2, 2)
plt.imshow(sharpen_blocks);
```

