

Digital Image Processing in Python

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<https://github.com/alonyan/DIP>

Workshop overview

- Day 1
 - Core concepts and motivation
 - What is a digital image
 - Pixel-level operations
- Day 2
 - Pixel-level operations (cont.)
 - Variance balancing (thresholding)
 - Image segmentation
- Day 3
 - Image segmentation (cont.)
 - Feature extraction
- Day 4
 - Cell tracking
 - Intro to modern computer vision

Workshop goals

The workshop is an introduction to digital image processing, designed to give you a taste of what's possible with a specific emphasis on microscopy data.

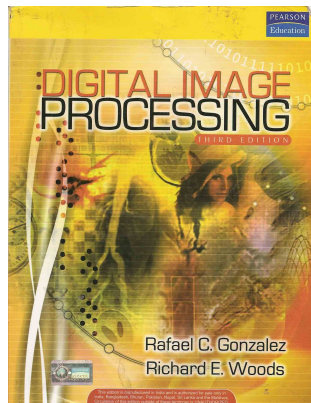
By the end of the workshop, you should:

- Acquire basic understanding and familiarity with computer vision.
- Appreciate the importance of rigorous and systematic image analysis for reproducible and quantitative science.

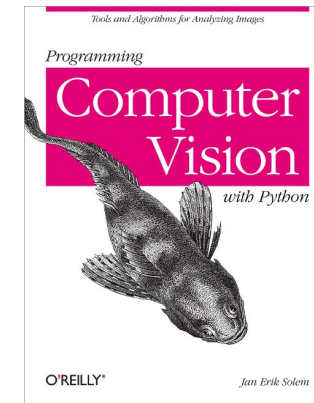
Other resources

MOOC: <https://www.mooc-list.com/course/image-analysis-methods-biologists-futurelearn>

<https://www.youtube.com/watch?v=1xo4vi6Ub4I>



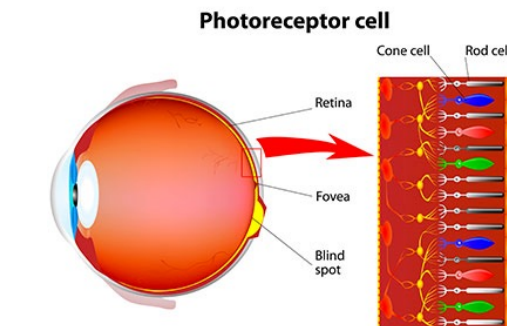
http://web.ipac.caltech.edu/staff/fmasci/home/astro_refs/Digital_Image_Processing_2ndEd.pdf



<http://programmingcomputervision.com/>

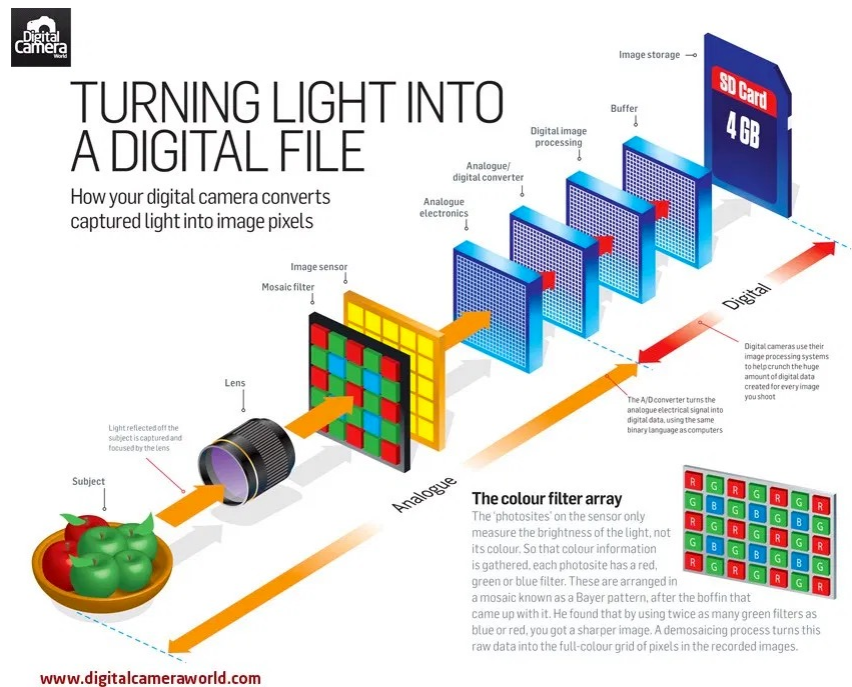
What is an Image?

- Photons are focused by lens (and cornea) and form an image on the retina.
- Activate a 2D array of photoreceptor cells that are sensitive to different colors.
- Information is passed to brain via optic nerves.



What is an Image?

- Photons are focused by a lens and form an image on an image sensor.
- Activate (charge) a 2D array of pixels on an image sensor (CCD/CMOS/PMT).
- (Sometimes there are color filters)
- Computers store information in bits , so the information must be digitized (ADC).
- Information is stored as a matrix of numbers.



How is pixel data stored in the computer?

Data type	Range
uint8 – unsigned 8 bit integer	0 to 255
uint16 – unsigned 16 bit integer	0 to 65535
uint32 – unsigned 32 bit integer	0 to $2^{32} - 1$
float – floating point, single (32 bit) or double (64 bit) precision	-1 to 1 or 0 to 1
int8 - signed 8 bit integer	-128 to 127
int16 - signed 16 bit integer	-32768 to 32767
int32 - signed 32 bit integer	- 2^{31} to $2^{31} - 1$

- The cost of higher precision is higher memory use.
- Higher precision is sometimes pointless, depending on the data and what you need to get from it.

What is an Image?

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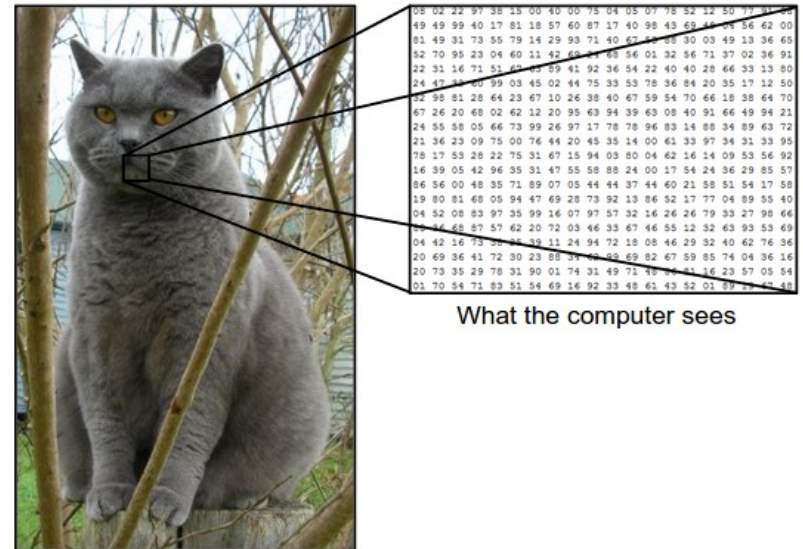


Image Resolution

- Light propagates as a wave.
- Waves can have constructive or destructive interference.
- Lenses are designed to maximize constructive interference at the focal plane.

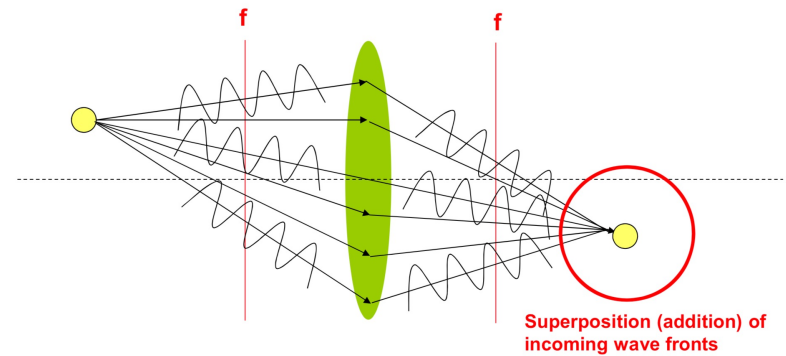


Image Resolution – Point Spread Function

- Image of point source - PSF

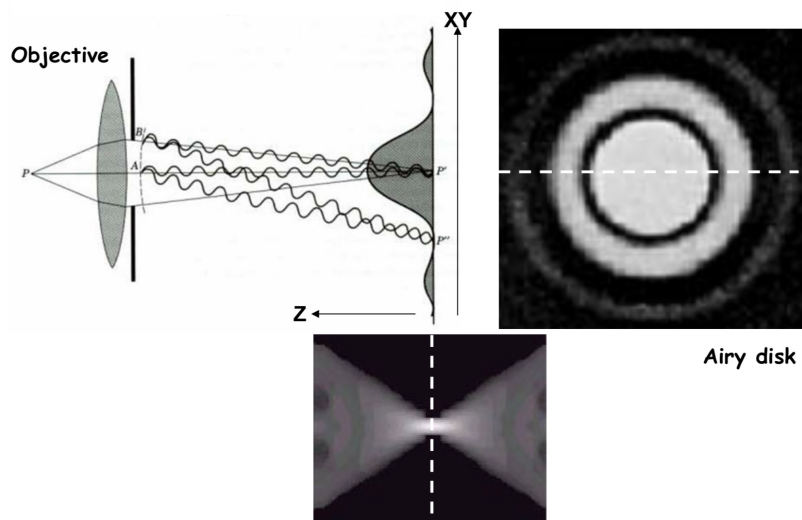


Image Resolution – Effect of Numerical Aperture

- $NA = n \cdot \sin \theta$
- Larger NA = Thinner PSF

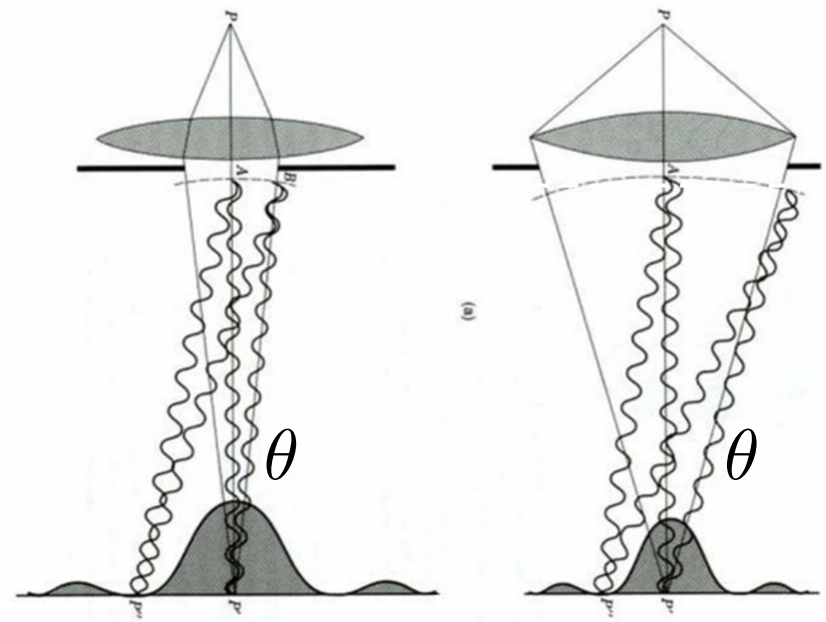
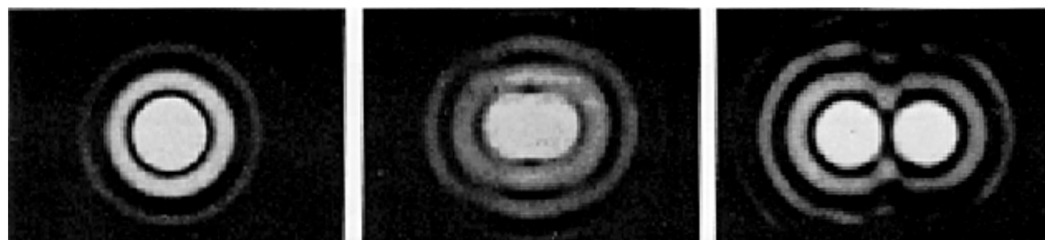


Image Resolution – How close two point can be and still be separable



$$\text{Lord Rayleigh's criterion: } \delta^R = 0.61 \frac{\lambda}{\text{NA}} \qquad \delta_z^R = 2 \frac{\lambda n}{\text{NA}^2}$$

Physiologically motivated !!!

Aka Abbe
diffraction
limit

Image Resolution and magnification

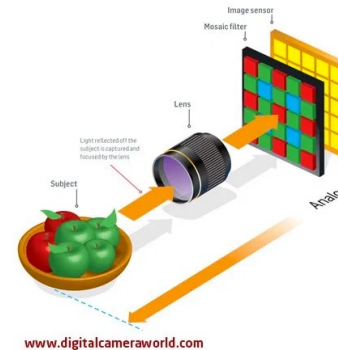
- Notice that the objective magnification **does not affect** Abbe's resolution formulae:

$$\delta^R = 0.61 \frac{\lambda}{NA} \quad \delta_z^R = 2 \frac{\lambda n}{NA^2}$$



- Caveat: $\frac{\text{pixel size}}{\text{Mag}}$ is the area captured by each pixel in the camera.

If that is larger than Abbes limit than the resolution is set by that. In that case the system is not diffraction limited.



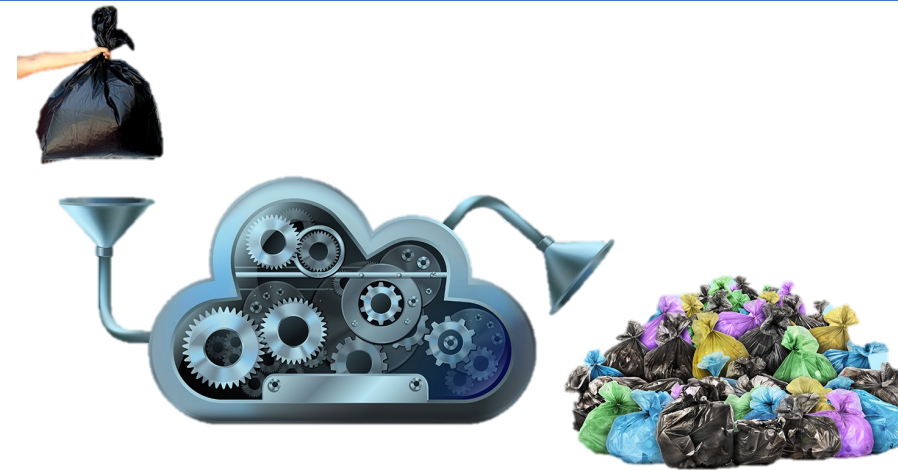
What is Image Processing?



Bunch of monkeys with
different attributes

- Our brain is really really good at this!
- Computer vision - teach **computers** to interpret and understand the visual world

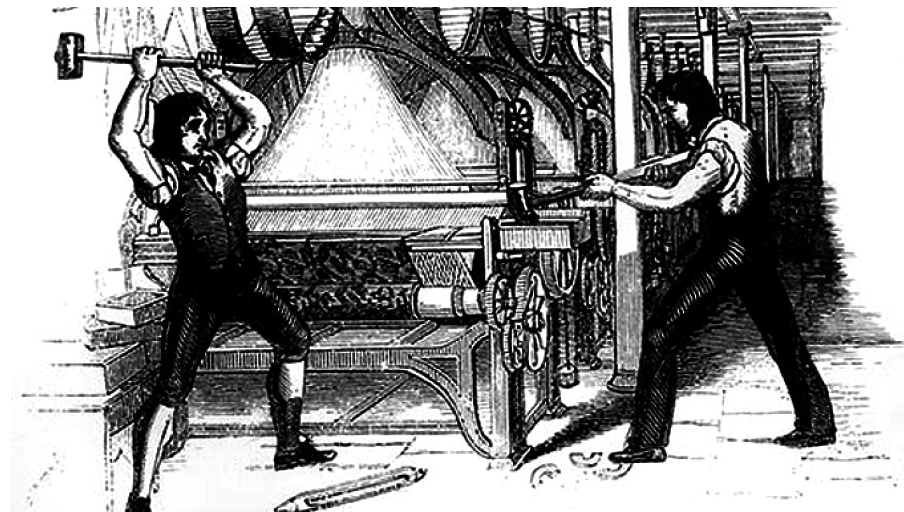
What is not Image Processing?



- Image processing is not magic
- Image processing doesn't create new data – that's called image manipulation/forgery
- It's always better to fix problems on the acquisition side than on the processing side
- **Good data is key!**

Why do we need image processing?

- If we're so good at this, why do we need computers?



(Optical) Microscopy circa y<2000 (b.f.p)

- Primarily antibody based
- Mostly fixed samples
- Low (throughput) and slow
- MB scale data,
typically ~10 images

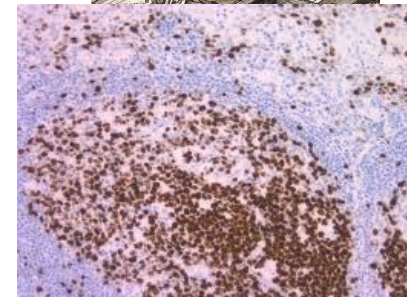
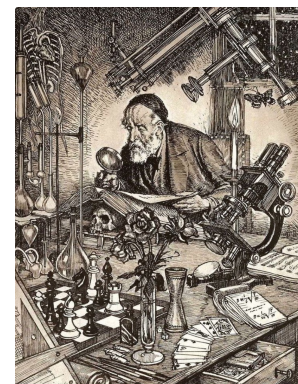


Image analysis was often performed manually,
usually on a subset of the data (“cherry picking”)
And heuristic methods that “make sense”

Microscopy circa y>2000 (a.f.p)

- FPs allow multichannel, live, long time imaging.
- New technologies:
 - Imaging beyond Abbe limit (STORM, PALM, STED, SIM...).
 - Imaging of whole organisms in 5D using LSFM (SPIM, Scanning beam, OPM,...).
 - Multiplexed *in situ* RNA FISH (MERFISH, SeqFISH).
 - Multiplexed IF (CyCIF, MIBI)
 - Cryo-EM

The scale and type of data for the modern microscopist has exploded in the last 2 decades.

GB-TB scale data. In some cases $\sim 10^6$ Images/experiment. Manual image analysis becomes impractical.

Imaging has become a big data + high throughput problem.

Reproducibility

- Our goal is to extract meaningful, quantifiable information in a non-biased manner from imaging data.
- Manual, non-systematic, image analysis is prone to bias.



Computational Image processing

- Image processing is about *extracting* the relevant data from your measurements, but not about analyzing them.



Something
something
computers
...

Monkey ID	M/F	size	Coat Color	Attr 1	Attr 2	Attr 3	...
1	F	24	...				
2	F	40					
3	M	87					
4	M	21					
5	F	31					
6	F	24					
7	F	54					

What might an image processing pipeline look like?

Image acquisition



What might an image processing pipeline look like?

Image acquisition



Image Preprocessing

- Aligning
- Tiling
- Denoising

What might an image processing pipeline look like?

Image acquisition



Image Preprocessing

- Aligning
- Tiling
- Denoising

Image Processing

- Background / Foreground
- Segmentation
- Feature extraction
- Conversion to data

What might an image processing pipeline look like?

Image acquisition



Image Preprocessing

- Aligning
- Tiling
- Denoising

Image Processing

- Background / Foreground
- Segmentation
- Feature extraction
- Conversion to data

Data Analysis

- Anything that can be learned from the data

What kinds of images might we look at?

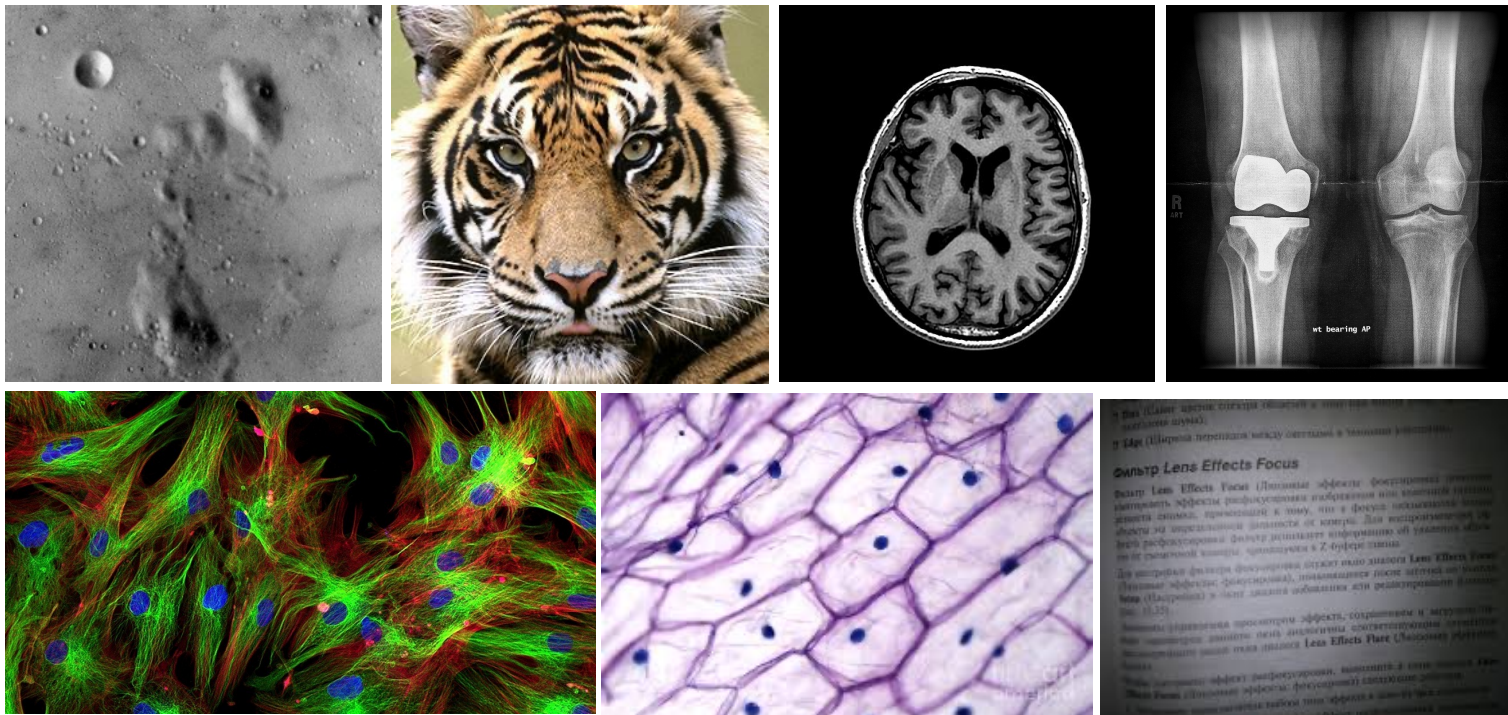


Image formats and compression

- It is important to have your data in the right format for the analysis you would like to perform.
- Some formats are considered **lossy** due to some compression when converting and saving in this format.
- Others are **lossless**, and uncompressed. This is more important when you would like to extract quantitative data from your image. They also tend to have larger file sizes.
- Many microscope manufacturers have their own proprietary image types and these will require specific tools to extract data from.

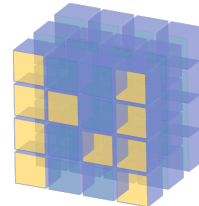
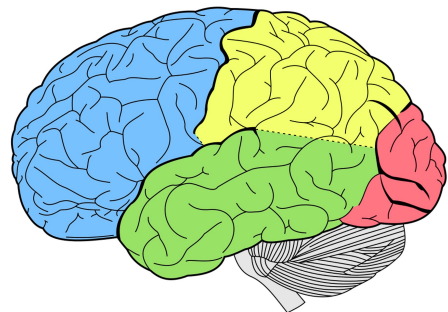
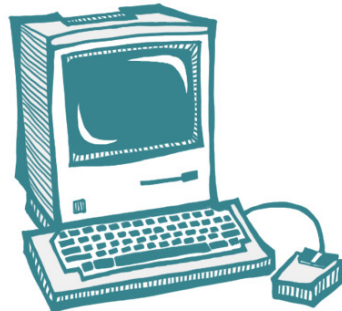
Some common data formats

- JPG or JPEG – Typically lossy and small bit depth (values range from 0 to 255). Can be grayscale or RGB, most common on standard digital cameras.
- PNG – Lossless image format but not high bit depth (0 to 255). Can also contain a transparency channel (alpha). One of the most common image formats.
- TIFF or TIF – Most common format in scientific imaging. Generally lossless and can have a bit depth up to 16 bit (0 to 65,535) in both grayscale and color. TIFF files can also be image stacks.
- CZI (zeiss) and LIF (Leica) – Proprietary image formats that can contain extra information about imaging and microscopy conditions.
- HDF5 / N5 - Hierarchical Data Format, designed to store very large amounts of structured data.

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Tools used in this workshop



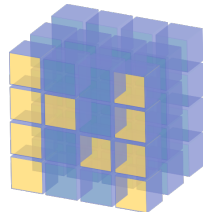
NumPy



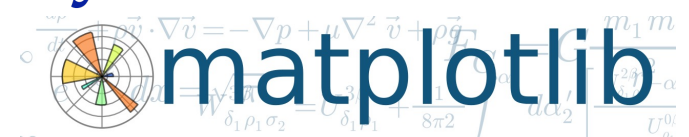
Libraries we will be using

One of python's major advantages is it's extensive libraries that have been developed for a variety of different applications.

We'll be focusing this workshop on the libraries shown here.



NumPy



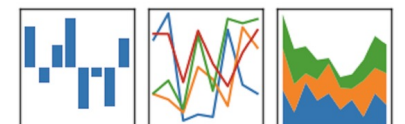
matplotlib

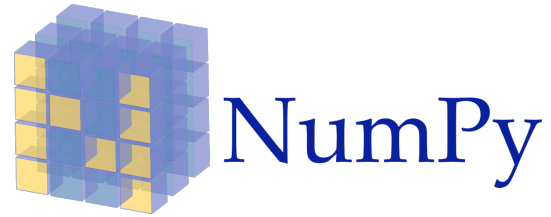


scikit-image
image processing in python

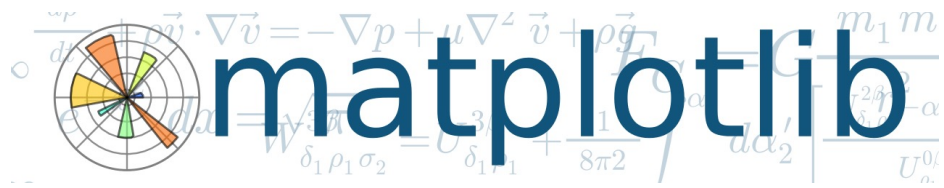
pandas

$$y_{it} = \beta' x_{it} + \mu_i + \epsilon_{it}$$





Advanced matrix calculator and
high-level math



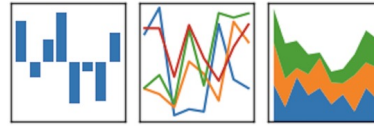
Graphing, plotting, and viewing images



High-level image processing

pandas

$$y_{it} = \beta' x_{it} + \mu_i + \epsilon_{it}$$



Data management, manipulation and
analysis

What we'll be doing

The course has two tracks

1. Fully completed notebooks
2. Fill in the blanks notebooks

- You're strongly encouraged to work in pairs/groups, and to use google to look for answers. This is how this works in the real world!
- If you have basic experience programing, you should be able to fill in the blanks.
- If you don't, you can either tag along with someone who's more experienced, or just follow along with a completed notebook.

We'll be going over everything together, leaving time for you to try and figure stuff out on your own.

Working on your own data

- Test data is included as part of the workshop.
- Please only attempt to try your own data **after** you completed the task with the test data.
- My ability to debug your code will be limited

Setup

Let's get going!

We need to download/clone the relevant course material and access it using jupyter notebook

Cloning/Downloading the course repository

- Go to <https://github.com/alonyan/DIP>

The screenshot shows the GitHub repository page for 'alonyan / DIP'. The page has a dark theme. At the top, there's a navigation bar with links for Pull requests, Issues, Marketplace, and Explore. Below that is a header with the repository name 'alonyan / DIP', a star icon, a 'Unwatch' button, a star count of 1, a 'Star' button, a fork count of 0, and a 'Fork' button. A yellow callout box with the text 'Download the repo' and an arrow points to the 'Clone or download' button, which is highlighted in green. The main content area shows repository statistics: 2 commits, 1 branch, 0 releases, 1 contributor, and an MIT license. It also shows a dropdown for the branch ('master') and a 'New pull request' button. Below these are commit details for 'Alon Oyler-Yaniv' and three files: '20190311', 'LICENSE', and 'README.md', each with their respective file icons, commit descriptions, and timestamps.

UCLA QCBio Collaboratory workshop 21: Digital Image Analysis with Python

Manage topics

Branch: master ▾ New pull request

2 commits 1 branch 0 releases 1 contributor MIT

Alon Oyler-Yaniv and Alon Oyler-Yaniv first commit

Latest commit f312cdb 29 seconds ago

20190311 first commit 29 seconds ago

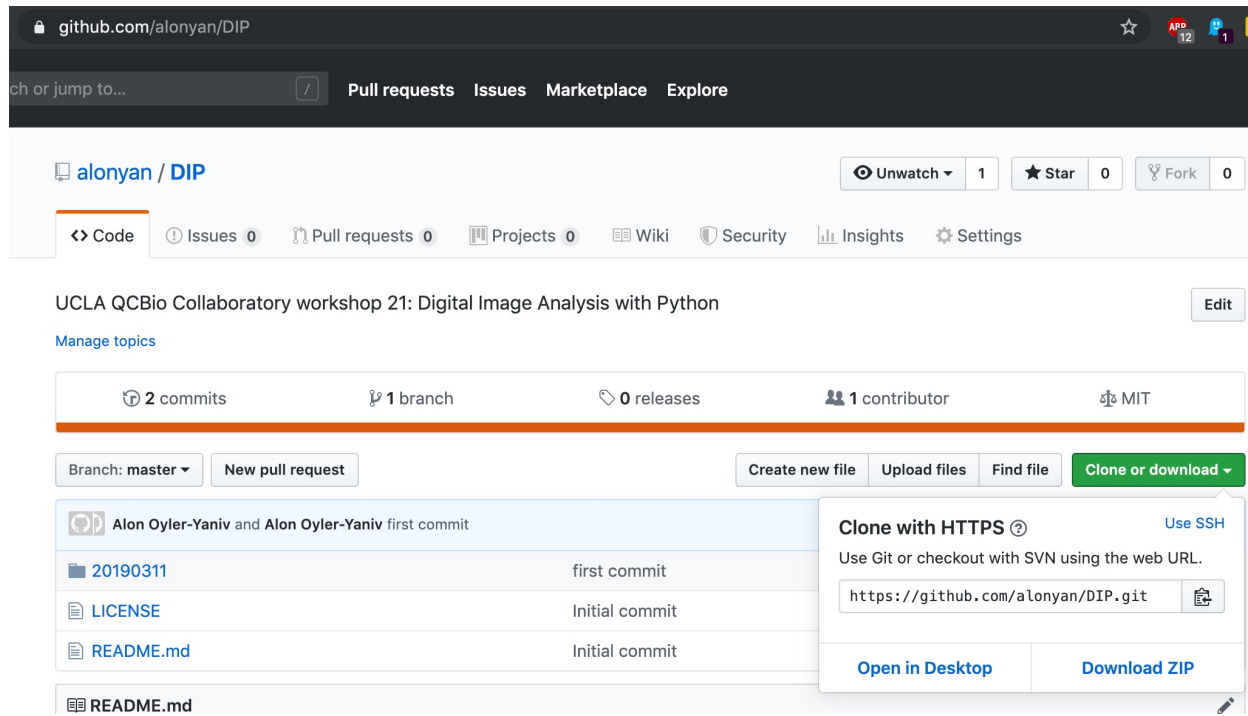
LICENSE Initial commit 8 minutes ago

README.md Initial commit 8 minutes ago

Download the repo

Cloning/Downloading the course repository

- Go to <https://github.com/alonyan/DIP>



Cloning/Downloading the course repository

- Move the .ZIP file to wherever you want it (/Documents is a good spot) and unzip
- You can also navigate (cd) in the terminal/command line to wherever you want it and type:

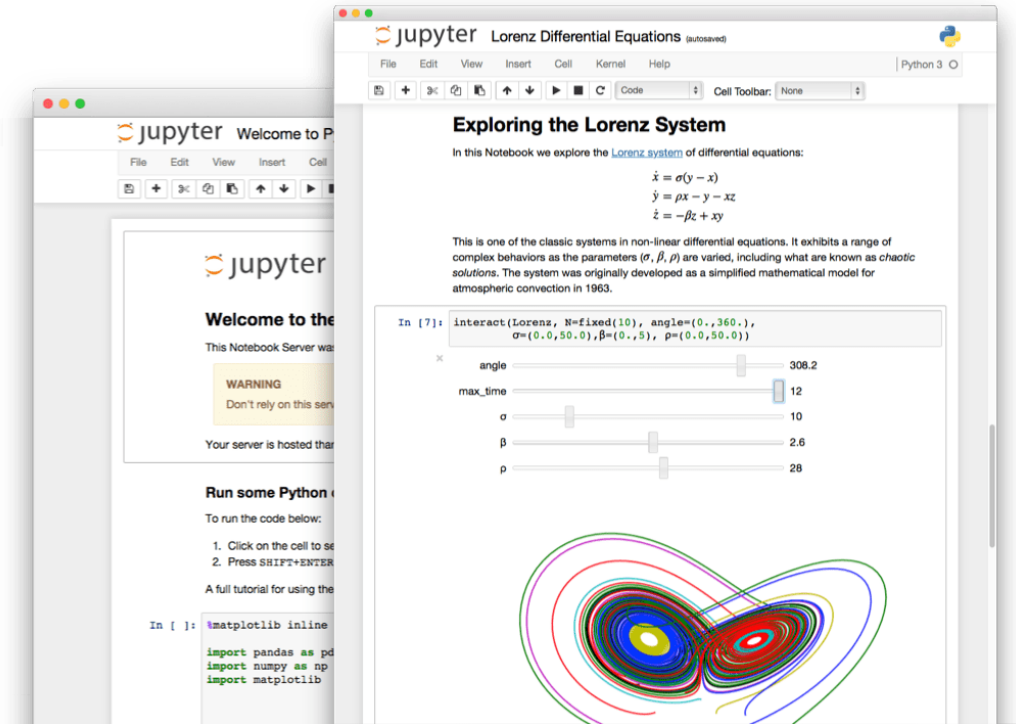
git clone <https://github.com/alonyan/DIP>

Getting started with Jupyter Notebooks

Jupyter notebooks

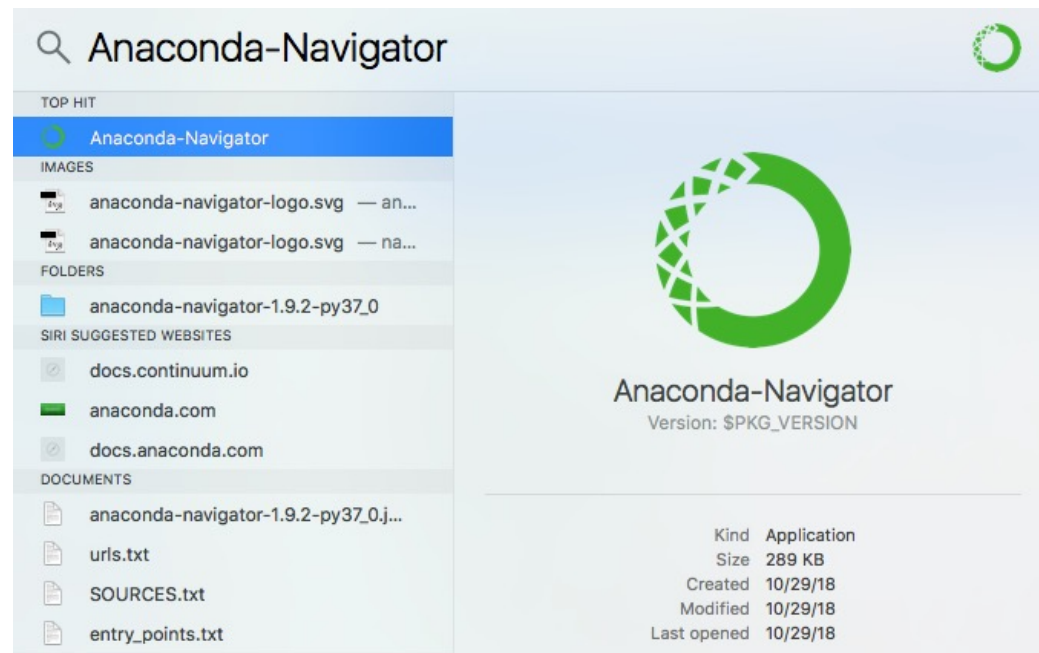
Jupyter notebooks are special documents that allow you to create and share documents **live code**.

They are a great way to keep your coding endeavors organized and share useful analysis with others.



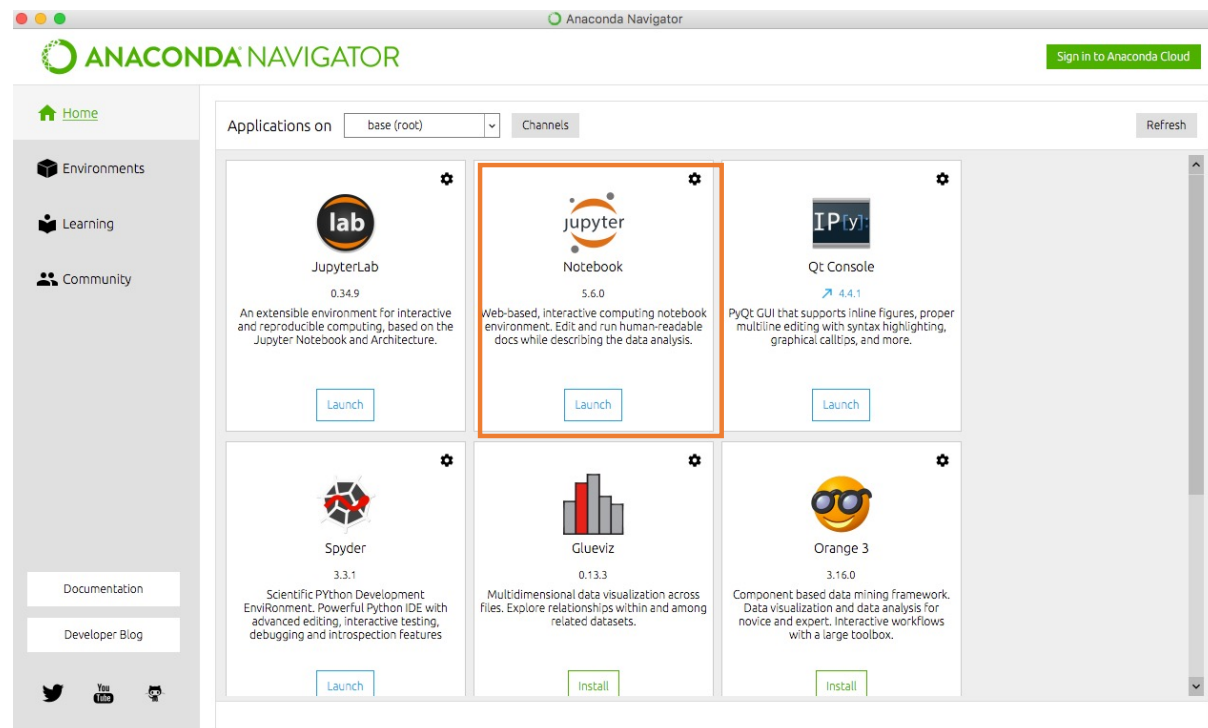
Getting started in Anaconda

- Launch the Anaconda application.
- MAC: Hit command+<space bar> and type “Anaconda”
- WINDOWS: Look for Anaconda on your start menu
- This should start you in the home directory.

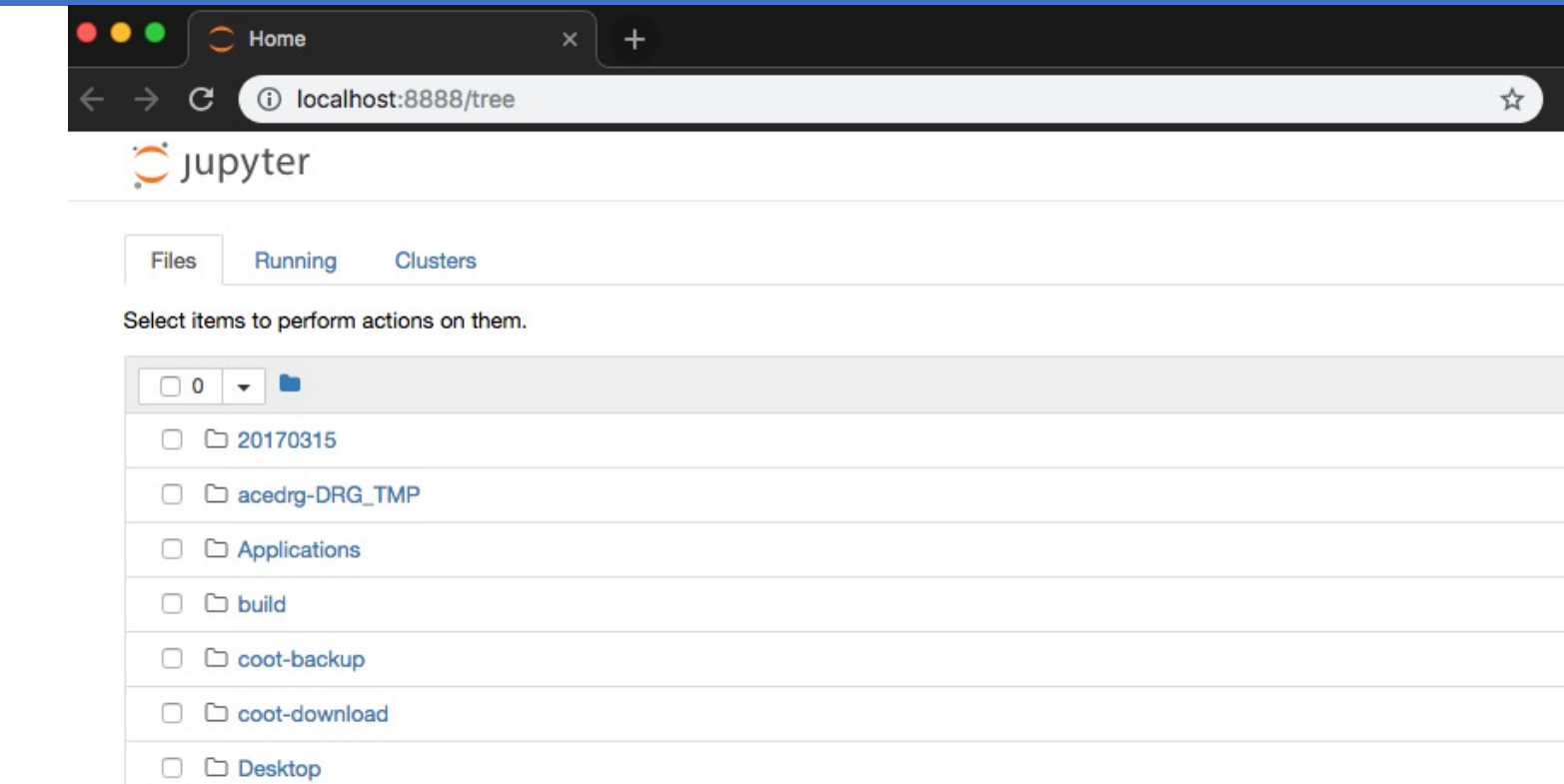


Getting started from Anaconda

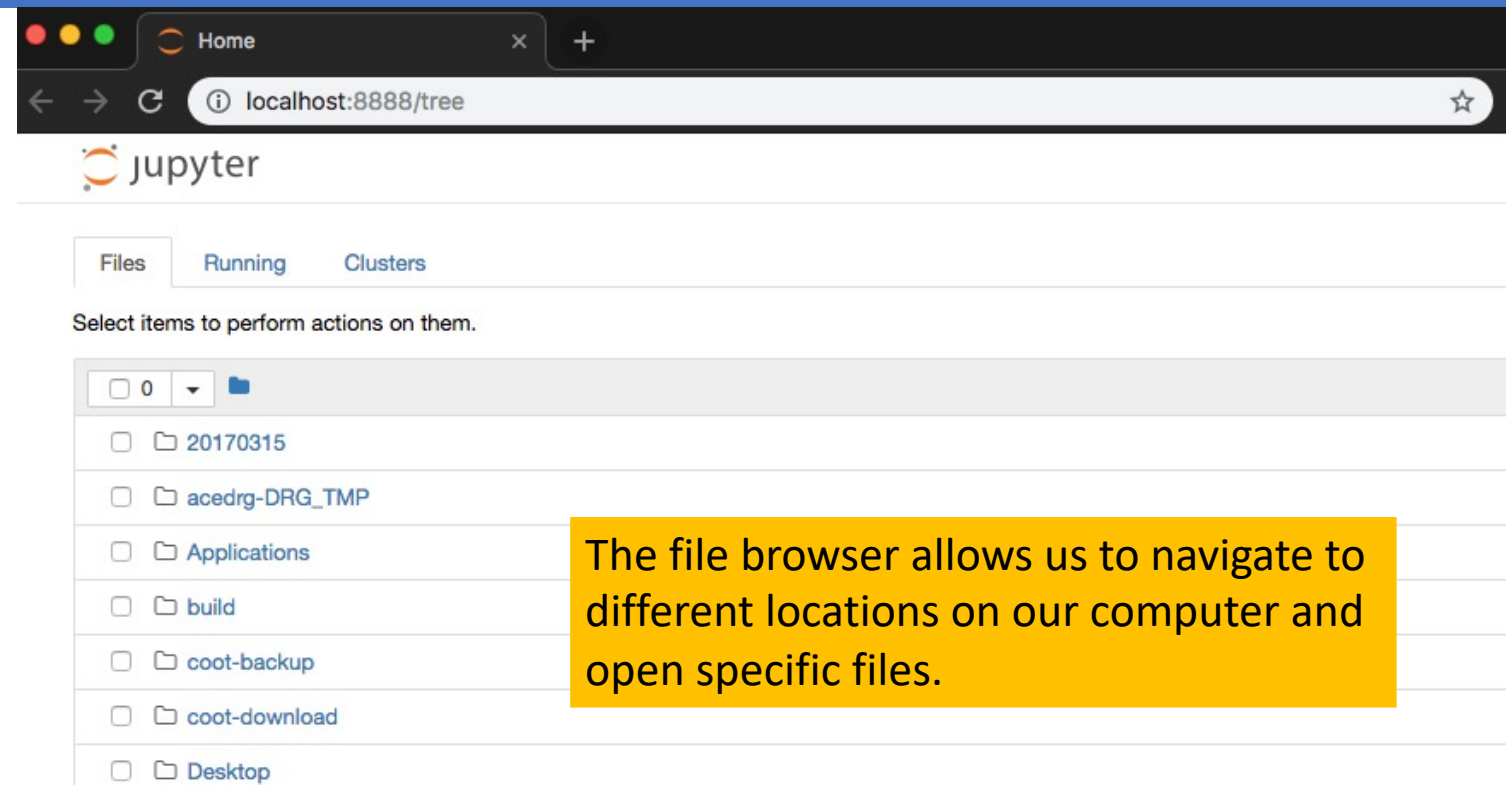
Click on the launch button under the **jupyter notebook** icon. After a few moments the dashboard should open in your default web browser.



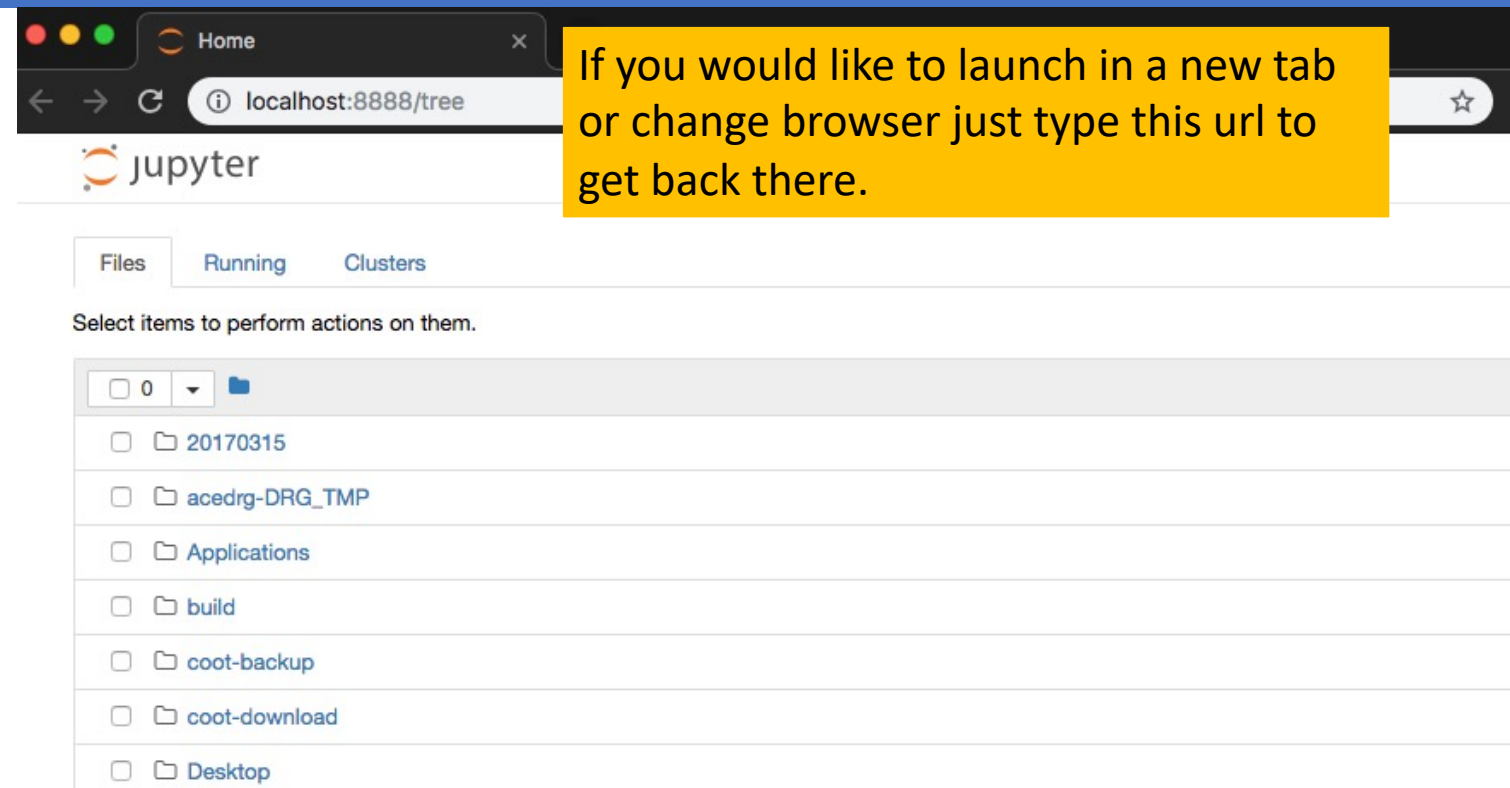
The jupyter dashboard



The jupyter dashboard



The jupyter dashboard



The jupyter dashboard

Navigate to wherever you saved the repository

The screenshot shows the Jupyter Notebook dashboard interface. At the top, there is a blue header bar with the text "The jupyter dashboard". Below this is a yellow navigation bar with the text "Navigate to wherever you saved the repository". The main content area has a dark header bar with a back arrow, a refresh icon, and the URL "localhost:8888/tree/Documents/Repos/DIP/20190311/JupyterNote...". To the right of the URL are icons for a star, a red circle with "ABP", a blue user icon with "0", a yellow folder icon, a "F" icon, and a circular profile picture. Below the header is a logo for "jupyter" with a circular orange icon. On the right side of the header are "Quit" and "Logout" buttons. The main body of the dashboard has three tabs: "Files" (selected), "Running", and "Clusters". Below the tabs, there is a message "Select items to perform actions on them." followed by "Upload", "New ▾", and a refresh icon. A file list table is shown, with columns for checkboxes, file/folder icons, names, last modified times, and file sizes. The table contains the following data:

	Name	Last Modified	File size
<input type="checkbox"/>	..	seconds ago	
<input type="checkbox"/>	images	10 minutes ago	
<input type="checkbox"/>	DIP_AOY_Student.ipynb	2 days ago	53.3 kB
<input type="checkbox"/>	DIP_AOY_Teacher.ipynb	2 days ago	7.8 MB

The jupyter dashboard

* .ipynb is the suffix of a notebook. Open the Student version (preferably) or the Teacher version of the nb.

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<input type="checkbox"/>	0	<input type="button"/>	Documents / Repos / DIP / 20190311 / JupyterNotebooks	Name	Last Modified	File size
			..		seconds ago	
			<input type="checkbox"/> images		10 minutes ago	
			<input type="checkbox"/> DIP_AOY_Student.ipynb		2 days ago	53.3 kB
			<input type="checkbox"/> DIP_AOY_Teacher.ipynb		2 days ago	7.8 MB

jupyter DIP_AOY_Student Last Checkpoint: 01/14/2020 (autosaved)



Logout

File Edit View Insert Cell Kernel Widgets Help

Trusted

Python 3



disclaimer: To ensure that the notebook can be run from (more or less) any point, I try to load the relevant functions or modules whenever I use them in a cell. This is generally not good practice as it adds unnecessary overhead

0. Image representation as numerical arrays

You should see something like this

We start by importing numpy and c
array

```
In [1]: %matplotlib inline
%load_ext autoreload
%autoreload
import numpy as np

#make a 9x9 checkerboard
checkBoard = #

print(checkBoard)

File "<ipython-input-1-149394df68b1>", line 7
```

Notebooks are divided into code blocks called “cells”.
To run all the code in a specific cell hit Shift+Enter

Python package structure – Modular programming

Package . Module . Function

Collection of modules Collection of functions Does a thing

Directory containing *.py files (and an __init__.py) Single *.py file Text within a *.py file

Import package **as pkg**

From package import module

From package.module import function

From module import *

pkg.module.function()

module.function()

function()

any_function_in_module()

**Shall we
begin?**

quickmeme.com