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# **Quantitative Big Imaging - Introduction**

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CHAPTER  
ONE

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## TODAYS LECTURE

- About the course
- Motivating the use of quantitative methods in imaging
- What is an image?
- Where do images come from?
- Science and Reproducibility
- Workflows

### 1.1 We need some python modules

Python is a modular scripting language with limited functionality. Features are added through modules that are imported. These are the modules that are needed for this lecture. Please run this cell before you start using the notebook.

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

from skimage.io import imread
from scipy.ndimage import convolve
from skimage.morphology import disk
from skimage.transform import resize
from itertools import product
import os
from io import StringIO
```

```
-----
ModuleNotFoundError                                     Traceback (most recent call last)
<ipython-input-1-1b9e7bae5a49> in <module>
      1 import numpy as np
----> 2 import pandas as pd
      3 import matplotlib.pyplot as plt
      4 get_ipython().run_line_magic('matplotlib', 'inline')
      5

ModuleNotFoundError: No module named 'pandas'
```



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**CHAPTER  
TWO**

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## **ABOUT THE COURSE**

- Who are we?
- Who are you?
- What is expected?
- **Why does this class exist?**
- Collection
- Changing computing (Parallel / Cloud)
- Course outline

### **2.1 Who are we?**

#### **Anders Kaestner**

You will hear me a lot during this course. I am the lecturer and I will also support you with problems during the exercises.

- **Beamline scientist** at the ICON Beamline at the SINQ (Neutron Source) at Paul Scherrer Institute
  - **Lecturer** at ETH Zurich
- **Algorithm developer** Varian Medical Systems, Baden-Daettwil
- **Post Doc** at ETH Zurich, Inst for Terrestrial Ecology
- **PhD** at Chalmers Institute of Technology, Sweden, Signal processing

[anders.kaestner@psi.ch](mailto:anders.kaestner@psi.ch)

#### **Stefano van Gogh**

Will help you during the exercise sessions.

- **PhD Student** in the X-Ray Microscopy Group at ETH Zurich and Swiss Light Source at Paul Scherrer Institute
- Teaching assistant

[stefano.van-gogh@psi.ch](mailto:stefano.van-gogh@psi.ch)

## 2.2 Who are you?

This course is targeting a wide range of students with different levels of experience. In the table you'll see where students came from in previous years. Some have a technical background others are merely producing images in the line of their project and have never seen much more than photoshop and similar programs for processing image data. Using some kind of programming is necessary to perform quantitative image analysis on large data sets. A single or a few images can easily be handled with interactive software, but taking it beyond that is hard without writing some lines of code.

Now, some of you have little to no programming experience while others have been programming since they got their first computer in the hand.

A wide spectrum of backgrounds	A wide range of skills
Biomedical Engineers	
Physicists	I think I've heard of python before
Chemists	.
Art History Researchers	.
Mechanical Engineers and Computer Scientists	I write template C++ code and hand optimize it afterwards

## 2.3 So how will this ever work?

Now you maybe start to get worried! The purpose of this course is not to teach you programming but rather to provide you with a bag full recipes that you can use in your projects. Most of these recipes are just a list of the commands from different python modules that you need to perform your analysis. A side-effect will probably be that you learn one or two programming tricks on the way.

In the lectures, there will be small code pieces on the slides. Some of these are there to illustrate how an operation works, while other parts are there for the nice presentation of the results (this is mostly the second half of the code cell). Presenting the results is important. In the end, you want to show your results to the scientific community. So even though the plotting clutters the slide, there is something to learn there as well.

### Adaptive assignments

- Conceptual, graphical assignments with practical examples
  - Emphasis on choosing correct steps and understanding workflow
- Opportunities to create custom implementations, and perform more complicated analysis on larger datasets if interested
  - Emphasis on performance, customizing analysis, and scalability

## 2.4 Course Expectations

The practical part of the course has two parts. None of these are mandatory, but they will help you to better understand use the material you have learnt in the lectures.

Exercises
Usually 1 set per lecture
Optional (but recommended!)
Easy - jupyter notebooks are prepared for the exercises
Advanced - Writing Python, Java, C++, ...

The exercises are prepared in a way that you learn step by step what you need to do and guides you through the problems. We will be using jupyter notebooks for the lectures. This is a very common way to work with image data these days.

Science project
Optional (but strongly recommended)
Applying Techniques to answer scientific question!
Ideally use on a topic relevant for your current project, thesis, or personal activities
or choose from one of ours (will be online, soon)
Present approach, analysis, and results

In the optional science projects you will have the opportunity to test what you have learned during the course on real problems. This is the place for your creativity.

## 2.5 Projects

- A small image processing project
- Can be related to your Master or PhD project
- You will get input and ideas for your own projects
- You will get hands on experience on the techniques you learn here
- Can be used as discussion base for your exam

## 2.6 Course Overview

Topic	Date	Title	Description
<b>Introduc-tion</b>	25th February	Introduction and Workflows	Basic overview of the course, introduc-tion to ...
<b>Data</b>	4th March	Image Enhancement	Overview of what techniques are avail-able for ...
	11th March	Ground Truth: Building and Augmenting Datasets	Examples of large datasets, how they were buil... .
<b>Segmenta-tion</b>	18th March	Basic Segmentation, Discrete Binary Structures	How to convert images into structures, startin... .
	25th March	Advanced Segmentation	More advanced techniques for extracting struct... .
	1st April	Supervised Problems and Segmentation	More advanced techniques for extracting struct... .
	8th April	Easter break	Search for eggs
<b>Analysis</b>	15th April	Analyzing Single Objects, Shape, and Texture	The analysis and characterization of single st... .
	22th April	Analyzing Complex Objects and Distributions	What techniques are available to analyze more ... .
	29th April	Dynamic Experiments	Performing tracking and registration in dynami... .
<b>Big Imag-ing</b>	6th May	Imaging with multiple modalities	Combining information from different sources
	13th May	Ascension	Enjoy a lovely early summers day
	20th May	Scaling Up / Big Data	Performing large scale analyses on clus-ters
<b>Wrapping up</b>	27th May	Project Presentations	You present your projects

## 2.7 Today's Reading Material

- Some book on image processing with python (to be updated)
- Cloud Computing
- The Case for Energy-Proportional Computing *Luiz André Barroso, Urs Hözle, IEEE Computer, December 2007*
- Concurrency
- Reproducibility
- Trouble at the lab *Scientists like to think of science as self-correcting. To an alarming degree, it is not*
- Why is reproducible research important? *The Real Reason Reproducible Research is Important*
- Science Code Manifesto
- Reproducible Research Class @ Johns Hopkins University

## **2.8 Literature / Useful References**

These are books that are useful in many of the lectures. In particular the Image processing hand book by John Russ shows you an overview of typical image processing techniques.

- John C. Russ, “The Image Processing Handbook”,(Boca Raton, CRC Press)
  - Available [online](#) within domain [ethz.ch](#) (or [proxy.ethz.ch](#) / public VPN)
- Jean Claude, Morphometry with R
  - [Online](#) through ETHZ



## MOTIVATION - YOU HAVE DATA!

### 3.1 Imaging experiments produce a lot of data

Working with imaging techniques you will get a lot of images that shows the sample in the eye of the technique you are using. The experiment were you acquire these images is only a small fraction of the complete workflow from idea to the final scientific publication. The amount of data can also be overwhelming for many scientist with the consequence that the data is never analyzed properly, and then also not published in the way it really deserves.



Gigabytes....  
...or even  
terabytes of data

Fig. 3.1: A typical imaging experiment produces large amounts of data.

## 3.2 Motivation - how to proceed?

Now is the question how to proceed towards a working analysis workflow that results in repeatable analyses for your data.



Fig. 3.2: A crazy unstructured and unclear work flow to analyze images from your experiment.

- To understand what, why and how from the moment an image is produced until it is finished (published, used in a report, ...)
- To learn how to go from one analysis on one image to 10, 100, or 1000 images (without working 10, 100, or 1000X harder)

## 3.3 High acquisition rates

The trend in imaging is that experimentalist want to follow faster and faster processes. This wish can be supported the technical development of new detectors that provide very high acquisition rates. Here, we can also see that some cameras are able to produce more data than is uploaded per day on facebook and instagram!

- Detectors are getting bigger and faster constantly
- Todays detectors are really fast
  - $2560 \times 2160$  images @ 1500+ times a second = 8GB/s
- Matlab / Avizo / Python / ... are saturated after 60 seconds

Many of the analysis platforms are already overwhelmed with handling the data rates produced by typical detector systems at imaging instrument. This restriction is partly due to hardware limitations. The memory is to small, hard drives are not sufficiently fast. The other side of the problem is that these tools are not prepared to work with large data streams.

- A single camera
  - More information per day than Facebook
  - Three times as many images per second as Instagram

## 3.4 Different sources of images

Images are produced by many different detectors and in some cases they are even the output from simulations. In the next sections we see some different imaging modalities and the data rates they produce.

### 3.4.1 X-Ray

X-ray imaging at synchrotron light sources produces very high frame rates thanks to the high brilliance of the source. Here are some examples of data rates from some instruments.

- SRXTM images at (>1000fps) → 8GB/s
- cSAXS diffraction patterns at 30GB/s
- Nanoscopium Beamline, 10TB/day, 10-500GB file sizes

### 3.4.2 Optical

Optical imaging methods are more modest than the X-ray techniques, but still they produce data in the order of some hundred Mb per second.

- Light-sheet microscopy (see talk of Jeremy Freeman) produces images → 500MB/s
- High-speed confocal images at (>200fps) → 78Mb/s

### 3.4.3 Personal

Finally, we also take a look at cameras on the consumer market and see that these devices also produce relatively high data rates. This data must mostly be handled by normal household computers, which can be a challenging task...

- GoPro 4 Black - 60MB/s (3840 x 2160 x 30fps) for \$600
- [fps1000](#) - 400MB/s (640 x 480 x 840 fps) for \$400

## 3.5 The experiment life cycle

Now we have seen that there is a wish to obtain data at high rates and that there are technological solutions to provide this. The remaining part to develop is the post processing.

1. **Experimental Design** finding the right technique, picking the right dyes and samples has stayed relatively consistent, better techniques lead to more demanding scientists.
  2. **Measurements** the actual acquisition speed of the data has increased wildly due to better detectors, parallel measurement, and new higher intensity sources
  3. **Management** storing, backing up, setting up databases, these processes have become easier and more automated as data magnitudes have increased
  4. **Post Processing** this portion has been the most time-consuming and difficult and has seen minimal improvements over the last years
- 

The post processing is the least trivial part to generalize. The initial steps are often possible to generalize as these are operations that all types of imaging experiments need to go through. When it comes to the experiment specific

analysis the degree of generalization decreases and the scientists are left to develop their own procedure to extract the quantitative information from the images.

### 3.5.1 How is time used during the experiment life cycle?

With the development of faster acquisition systems there has been a change in the ratio between

- Experiment design and preparation
- Measurements
- Data management
- and post processing

over the years. This in particular the case for X-ray imaging where the flux is high and the acquisition is limited by the detector technology. In other modalities, where the measurement is flux limited we see a different distribution.

What also increases the post processing time is that the experiments have become more complicated over the years. Twenty years ago, it was sufficient to show qualitative information a beautify volume rendering or a movie of the sample. Meanwhile, it has become a requirement that you provide quantitative results from the images.

## 3.6 Handling masses of images

### 3.6.1 So... how much is a TB, really?

We have been talking about different data amounts of MB, GB, and TB. But, what does that really mean in reality? Let us explore what is a TB.

If you looked at one image with 1000 x 1000 pixels (1 Mpixels)

Here we create one image with 1000x1000 pixels with random values form a uniform distribution [0,1] and show it.

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

plt.matshow(np.random.uniform(size = (1000, 1000)),
            cmap = 'viridis');
```



Fig. 3.3: The ratio of how much time is spent on different tasks during the lifecycle of an imaging experiment.



every second, it would take you

```
# assuming 16 bit images and a 'metric' terabyte
OneTB      = 1e12
ImageSize  = 1000*1000*16/8
hour       = 60*60

time_per_tb = OneTB/ImageSize/hour
print("{0:.1f} hours to view a terabyte".format(time_per_tb))
```

```
138.9 hours to view a terabyte
```

### 3.6.2 Overwhelmed scientist

Providing quantitative statements about image data is often very hard. You can manage to do it on a single images like the bone image below.

You would like to know:

- Count how many cells are in the bone slice.
- Ignore the ones that are ‘too big’ or shaped ‘strangely’.
- Are there more on the right side or left side?
- Are the ones on the right or left bigger, top or bottom?



Fig. 3.4: A slice image show bone cells.

### 3.6.3 More overwhelmed scientist

Statistical analysis requires that you study many samples and not just a single one. The samples are also objects which requires 3D data instead of a single 2D slice.

Many samples are needed:

- Do it all over again for 96 more samples
- This time in 3D with 2000 slices instead of just one!

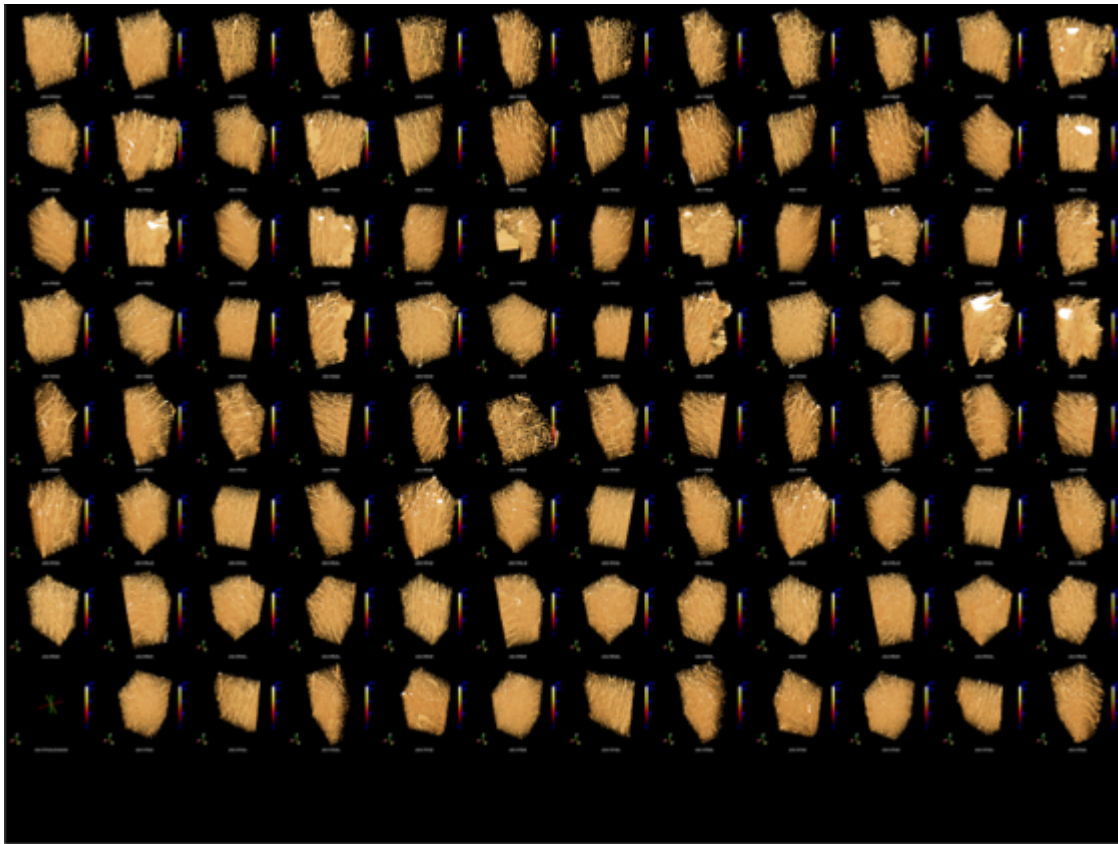


Fig. 3.5: A collection of 96 volume images from different bone samples.

Working with multiple 3D images is not feasible anymore to do manually. We need some kind of automated process to perform the analysis.

### 3.6.4 Bring on the pain

The 96 samples only represented one of our cases in the study. Now, if we want to study different ages, healthy/diseased, etc, we need to add a sample batch for each case. Maybe we even need to increase the number of samples in each test group. With all these variations, we can easily end up in a thousand samples to analyze.

Great variations in the population

- Now again with 1090 samples!
- How to measure?
- How to analyze?



Fig. 3.6: A collection of 1090 bone samples. This is a massive task to analyze!

With so many samples we stand in front of a logistic problem to measure the data and once the data is there we have to analyze it. As a first step, we have to specify how to analyze these images to obtain results that may or may not support a hypothesis.

### 3.6.5 It gets better

The metrics we specified in the previous example are easy to observe and also to measure. They are direct measurements of pixels and positions. What if we now want to make more complicated inquiries even. Now how do we categorize the images or collections of features using soft metrics?

- Those metrics were quantitative and could be easily visually extracted from the images
- What happens if you have *softer* metrics
  - How aligned are these cells?
  - Is the group on the left more or less aligned than the right?
  - errr?



Fig. 3.7: Close-up on different bone segments. How aligned are the cells in these images?

### **3.6.6 Dynamic Information**

Many experiments are on top of the spatial dimensions also studies over time. This brings us 4D data sets to analyze. How are we supposed to handle this? Looking at the movie we

- How many bubbles are here?
- How fast are they moving?
- Do they all move the same speed?
- Do bigger bubbles move faster?
- Do bubbles near the edge move slower?
- Are they rearranging?



## 4.1 An introduction to images

### 4.1.1 What is an image?

A very abstract definition:

- **A pairing between spatial information (position)**
- **and some other kind of information (value).**

In most cases this is a 2- or 3-dimensional position (x,y,z coordinates) and a numeric value (intensity)

### 4.1.2 Image sampling

The world is	The computer needs
Continuous	Discrete levels
No boundaries	Limited extent



Fig. 4.1: The real world is sampled into discrete images with limited extent.

### 4.1.3 What does sampling mean

```
img=np.load('.../common/data/wood.npy');
plt.figure(figsize=[15,7])
plt.subplot(2,3,1); plt.imshow(img); plt.title('Original')
downsize = 2; plt.subplot(2,3,2); plt.imshow(resize(img,(img.shape[0] // downsize,
.shape[1] // downsize), anti_aliasing=False)); plt.title('Downsize {0}x{0}'.
format(downsize))
downsize = 32; plt.subplot(2,3,3); plt.imshow(resize(img,(img.shape[0] // downsize,
.shape[1] // downsize),anti_aliasing=False)); plt.title('Downsize {0}x{0}'.
format(downsize))
levels = 16; plt.subplot(2,3,5); plt.imshow(np.floor(img*levels)); plt.title('{0}'.
format(levels));
levels = 4 ; plt.subplot(2,3,6); plt.imshow(np.floor(img*levels)); plt.title('{0}'.
format(levels));
```



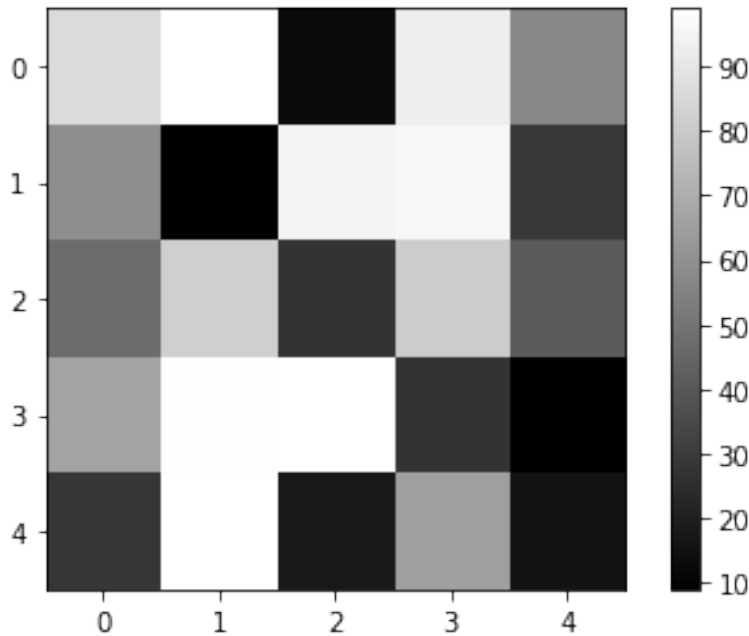
### 4.2 Let's create a small image

```
basic_image = np.random.choice(range(100), size = (5,5))

xx, yy = np.meshgrid(range(basic_image.shape[1]), range(basic_image.shape[0]))
image_df = pd.DataFrame(dict(x = xx.ravel(),
                             y = yy.ravel(),
                             Intensity = basic_image.ravel()))
image_df[['x', 'y', 'Intensity']].head(5)
```

x	y	Intensity
0	0	53
1	0	14
2	0	82
3	0	78
4	0	2

```
import matplotlib.pyplot as plt
plt.imshow(basic_image, cmap = 'gray')
plt.colorbar();
```



### 4.3 2D Intensity Images

The next step is to apply a color map (also called lookup table, LUT) to the image

- so it is a bit more exciting
- some features are easier to detect Rogowitz et al. 1996

```
fig, ax1 = plt.subplots(1,1)
plot_image = ax1.matshow(basic_image, cmap = 'Blues')
plt.colorbar(plot_image)

for _, c_row in image_df.iterrows():
    ax1.text(c_row['x'], c_row['y'], s = '%02d' % c_row['Intensity'], fontdict =
    dict(color = 'r'))
```



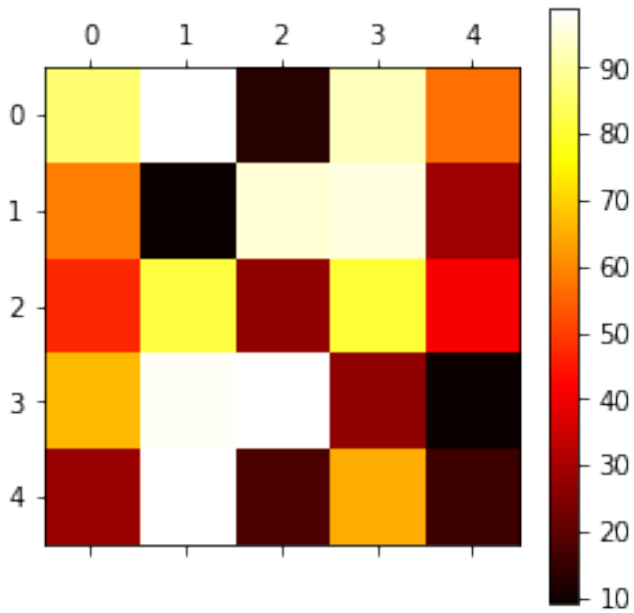
### 4.3.1 Different colormaps

Color maps can be arbitrarily defined based on how we would like to visualize the information in the image

```
plot_image = plt.matshow(basic_image, cmap = 'jet')
plt.colorbar(plot_image);
```



```
plot_image = plt.matshow(basic_image, cmap = 'hot')
plt.colorbar(plot_image);
```



### 4.3.2 Lookup Tables

Formally a color map is lookup table or a function which  $f(\text{Intensity}) \rightarrow \text{Color}$

#### Matplotlib's color maps

```
import numpy as np
import matplotlib as mpl
import matplotlib.pyplot as plt
from matplotlib import cm
#from colorspacious import cspace_converter
from collections import OrderedDict
cmmaps = OrderedDict()
cmmaps['Perceptually Uniform Sequential'] = [
    'viridis', 'plasma', 'inferno', 'magma', 'cividis']

cmmaps['Sequential'] = [
    'Greys', 'Purples', 'Blues', 'Greens', 'Oranges', 'Reds',
    'YlOrBr', 'YlOrRd', 'OrRd', 'PuRd', 'RdPu', 'BuPu',
    'GnBu', 'PuBu', 'YlGnBu', 'PuBuGn', 'BuGn', 'YlGn']

cmmaps['Sequential (2)'] = [
    'binary', 'gist_yarg', 'gist_gray', 'gray', 'bone', 'pink',
    'spring', 'summer', 'autumn', 'winter', 'cool', 'Wistia',
    'hot', 'afmhot', 'gist_heat', 'copper']

cmmaps['Diverging'] = [
    'PiYG', 'PRGn', 'BrBG', 'PuOr', 'RdGy', 'RdBu',
    'RdYlBu', 'RdYlGn', 'Spectral', 'coolwarm', 'bwr', 'seismic']

cmmaps['Cyclic'] = ['twilight', 'twilight_shifted', 'hsv']
```

(continues on next page)

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```
cmaps['Qualitative'] = ['Pastel1', 'Pastel2', 'Paired', 'Accent',
                        'Dark2', 'Set1', 'Set2', 'Set3',
                        'tab10', 'tab20', 'tab20b', 'tab20c']

cmaps['Miscellaneous'] = [
    'flag', 'prism', 'ocean', 'gist_earth', 'terrain', 'gist_stern',
    'gnuplot', 'gnuplot2', 'CMRmap', 'cubeHelix', 'brg',
    'gist_rainbow', 'rainbow', 'jet', 'nipy_spectral', 'gist_ncar']

nrows = max(len(cmap_list) for cmap_category, cmap_list in cmaps.items())

gradient = np.linspace(0, 1, 256)
gradient = np.vstack((gradient, gradient))

def plot_color_gradients(cmap_category, cmap_list, nrows):
    fig, axes = plt.subplots(nrows=nrows)
    fig.subplots_adjust(top=0.95, bottom=0.01, left=0.2, right=0.99)
    axes[0].set_title(cmap_category + ' colormaps', fontsize=14)

    for ax, name in zip(axes, cmap_list):
        ax.imshow(gradient, aspect='auto', cmap=plt.get_cmap(name))
        pos = list(ax.get_position().bounds)
        x_text = pos[0] - 0.01
        y_text = pos[1] + pos[3]/2.
        fig.text(x_text, y_text, name, va='center', ha='right', fontsize=10)

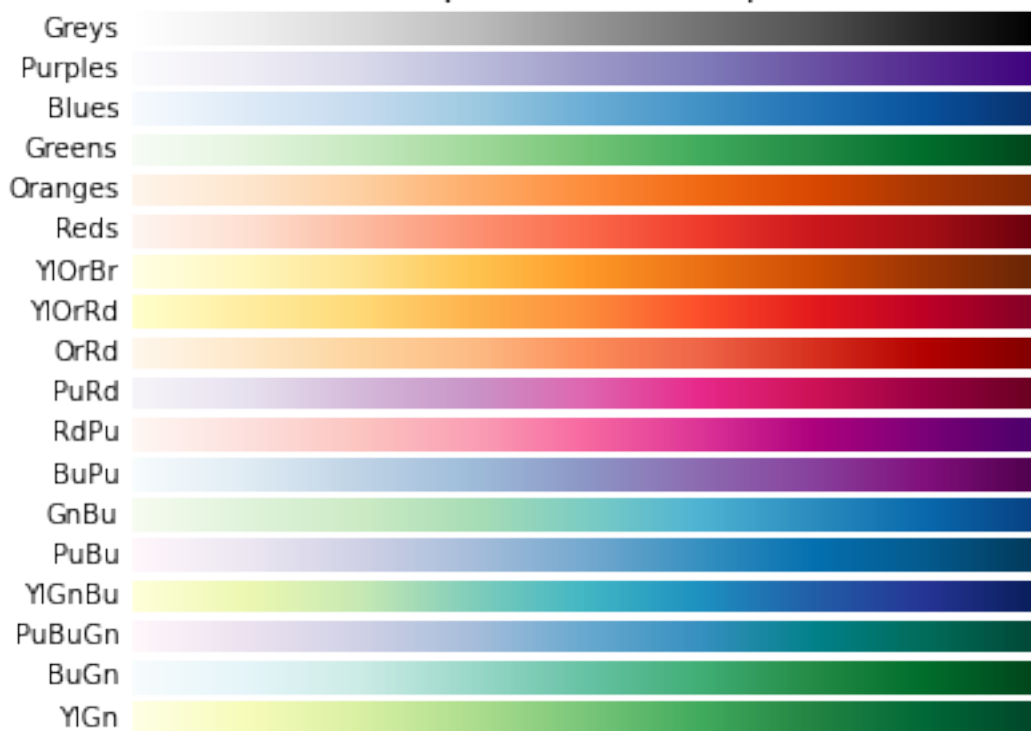
    # Turn off *all* ticks & spines, not just the ones with colormaps.
    for ax in axes:
        ax.set_axis_off()

for cmap_category, cmap_list in cmaps.items():
    plot_color_gradients(cmap_category, cmap_list, nrows)
```

### Perceptually Uniform Sequential colormaps



### Sequential colormaps



### Sequential (2) colormaps



### Diverging colormaps



### Cyclic colormaps



### Qualitative colormaps





### 4.3.3 How are the colors combined

```
%matplotlib inline
import matplotlib.pyplot as plt
import numpy as np
xlin = np.linspace(0, 1, 100)
colors = ['Red', 'Green', 'Blue']
plt.figure(figsize=[15, 4])
for i in np.arange(0,3) :
    plt.subplot(1,3,i+1)

    plt.scatter(xlin,
                plt.cm.hot(xlin) [:,i],
                c = plt.cm.hot(xlin),label="hot")
    plt.scatter(xlin,
                plt.cm.Blues(xlin) [:,i],
                c = plt.cm.Blues(xlin),label="blues")

    plt.scatter(xlin,
                plt.cm.jet(xlin) [:,i],
                c = plt.cm.jet(xlin),label='jet')

plt.xlabel('Intensity');
plt.ylabel('{0} Component'.format(colors[i]));
```



#### 4.3.4 Applied LUTs

These transformations can also be non-linear as is the case of the graph below where the mapping between the intensity and the color is a log relationship meaning the the difference between the lower values is much clearer than the higher ones

```
%matplotlib inline
import matplotlib.pyplot as plt
import numpy as np
xlin = np.logspace(-2, 5, 500)
log_xlin = np.log10(xlin)
norm_xlin = (log_xlin-log_xlin.min())/(log_xlin.max()-log_xlin.min())
fig, ax1 = plt.subplots(1,1)

ax1.scatter(xlin, plt.cm.hot(norm_xlin)[:,0],
            c = plt.cm.hot(norm_xlin))

ax1.scatter(xlin, plt.cm.hot(xlin/xlin.max())[:,0],
            c = plt.cm.hot(norm_xlin))
ax1.set_xscale('log');ax1.set_xlabel('Intensity');ax1.set_ylabel('Red Component');
```



### 4.3.5 LUTs on real images

On a real image the difference is even clearer

```
%matplotlib inline
import matplotlib.pyplot as plt
from skimage.io import imread
fig, (ax1, ax2, ax3) = plt.subplots(1,3, figsize = (12, 4))
in_img = imread('figures/bone-section.png')[:, :, 0].astype(np.float32)
ax1.imshow(in_img, cmap = 'gray');
ax1.set_title('grayscale LUT');

ax2.imshow(in_img, cmap = 'hot');
ax2.set_title('hot LUT');

ax3.imshow(np.log2(in_img+1), cmap = 'gray');
ax3.set_title('grayscale-log LUT');
```



## 4.4 3D Images

For a 3D image, the position or spatial component has a 3rd dimension (z if it is a spatial, or t if it is a movie)

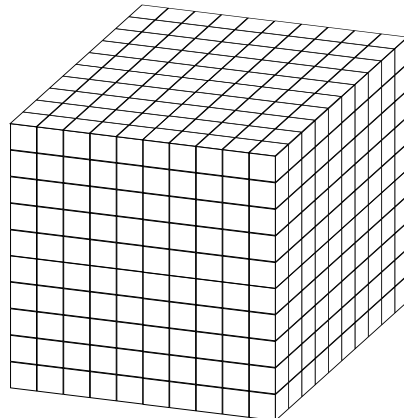


Fig. 4.2: Three-dimensional data can be a volume in space.



Fig. 4.3: A movie can also be seen as a three-dimensional image.

### 4.4.1 A 3D image as array

```
import numpy as np
vol_image = np.arange(27).reshape((3, 3, 3))
print(vol_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]]]]
```

### 4.4.2 Showing 2D slices from volume

This can then be rearranged from a table form into an array form and displayed as a series of slices

```
%matplotlib inline
import matplotlib.pyplot as plt
from skimage.util import montage as montage2d
print(montage2d(vol_image, fill = 0))
plt.matshow(montage2d(vol_image, fill = 0), cmap = 'jet');
```

```
[[ 0  1  2   9 10 11]
 [ 3  4  5 12 13 14]
 [ 6  7  8 15 16 17]
 [18 19 20  0  0  0]
 [21 22 23  0  0  0]
 [24 25 26  0  0  0]]
```



## 4.5 Multiple Values per pixel

In the images thus far, we have had one value per position, but there is no reason there cannot be multiple values. In fact this is what color images are (red, green, and blue) values and even 4 channels with transparency (alpha) as a different. For clarity we call the **dimensionality** of the image the number of dimensions in the spatial position, and the **depth** the number in the value.

```
base_df = pd.DataFrame([dict(x = x, y = y) for x,y in product(range(5), range(5))])
base_df['Intensity'] = np.random.uniform(0, 1, 25)
base_df['Transparency'] = np.random.uniform(0, 1, 25)
base_df.head(5)
```

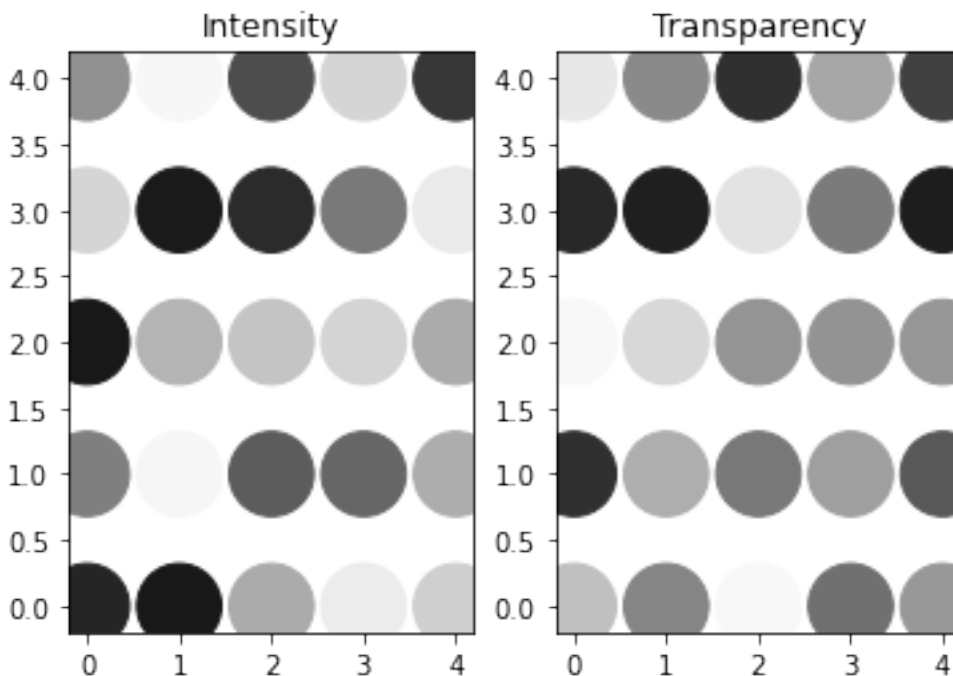
	x	y	Intensity	Transparency
0	0	0	0.144754	0.750454
1	0	1	0.496202	0.189002
2	0	2	0.094933	0.968853
3	0	3	0.833134	0.158436
4	0	4	0.566797	0.909068

This can then be rearranged from a table form into an array form and displayed as a series of slices

#### 4.5.1 Display multi-valued pixels separately

The most straight forward way to display multiple pixel values is to display each value separately. This method is, however, mostly not very suitable as the values often are related in some sense. Therefore it is recommended to combine the values in the same plot.

```
fig, (ax1, ax2) = plt.subplots(1, 2)
ax1.scatter(base_df['x'], base_df['y'], c = plt.cm.gray(base_df['Intensity']), s =_
↪1000)
ax1.set_title('Intensity')
ax2.scatter(base_df['x'], base_df['y'], c = plt.cm.gray(base_df['Transparency']), s =_
↪1000)
ax2.set_title('Transparency');
```



In this example we combined two values use one value to control the colormap and the other to control the size of the dots. How you combine the data is related to the type of data you want to combine. If the values are components of a vector it makes more sense to show arrows of different length and direction, etc.

```
fig, (ax1) = plt.subplots(1, 1)
ax1.scatter(base_df['x'], base_df['y'], c = plt.cm.jet(base_df['Intensity']), s =_
↪1000*base_df['Transparency'])
ax1.set_title('Intensity');
```



## 4.6 Hyperspectral Imaging

At each point in the image (black dot), instead of having just a single value, there is an entire spectrum. A selected group of these (red dots) are shown to illustrate the variations inside the sample. While certainly much more complicated, this still constitutes an image and requires the same sort of techniques to process correctly.

```
%matplotlib inline
import matplotlib.pyplot as plt
import pandas as pd
from skimage.io import imread
import os
raw_img = imread('.../common/data/raw.jpg')
im_pos = pd.read_csv('.../common/data/impos.csv', header = None)
im_pos.columns = ['x', 'y']
fig, ax1 = plt.subplots(1,1, figsize = (8, 8));
ax1.imshow(raw_img);
ax1.scatter(im_pos['x'], im_pos['y'], s = 1, c = 'blue');
```



### 4.6.1 Looking at the pixel values

Each pixel of this data set is represented by a full spectrum. This means that we have both wave numbers and intensity values. In many cases, the wavenumbers are the same for all pixels, which makes it possible to reduce the redundancy of the wave number vector using a single 1D array to represent this information.

```
full_df = pd.read_csv('../common/data/full_img.csv').query('wavenum<1200')
print(full_df.shape[0], 'rows')
full_df.head(5)
```

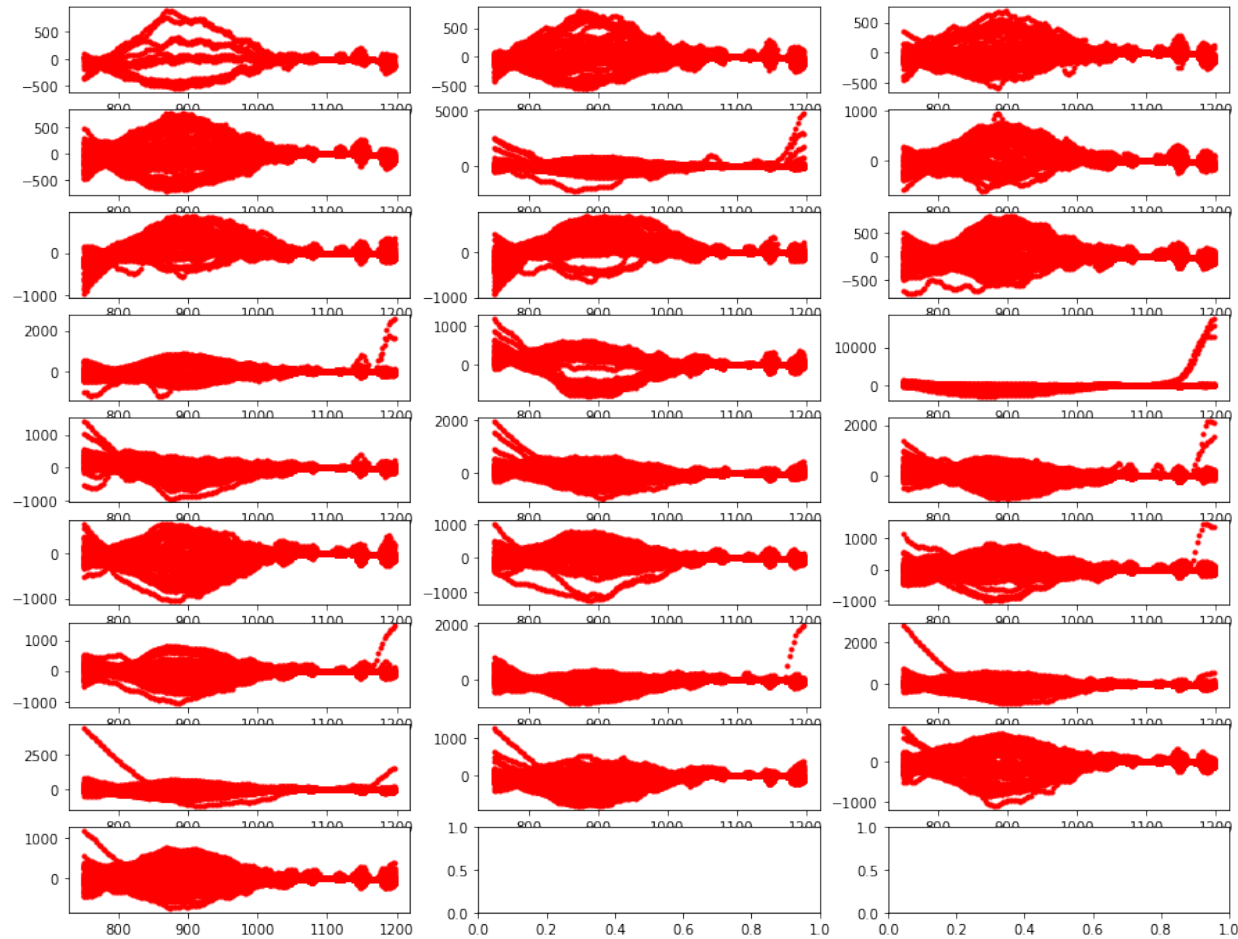
```
210750 rows
```

	x	y	wavenum	val
0	168.95	358.8	750	527.571102
1	168.95	358.8	753	459.778584
2	168.95	358.8	756	406.337255
3	168.95	358.8	759	341.858123
4	168.95	358.8	762	246.645673

```

full_df['g_x'] = pd.cut(full_df['x'], 5)
full_df['g_y'] = pd.cut(full_df['y'], 5)
fig, m_axs = plt.subplots(9, 3, figsize = (15, 12)); m_axs=m_axs.ravel()
for ((g_x, g_y), c_rows), c_ax in zip(full_df.sort_values(['x','y']).groupby(['g_x', 'g_y']), m_axs):
    c_ax.plot(c_rows['wavenum'], c_rows['val'], 'r.')

```



## 4.7 Image Formation

The image formation process is the process to use some kind of excitation or impulse probe a sample. This requires the interaction of four parts.

- **Impulses** Light, X-Rays, Electrons, A sharp point, Magnetic field, Sound wave
- **Characteristics** Electron Shell Levels, Electron Density, Phonons energy levels, Electronic, Spins, Molecular mobility
- **Response** Absorption, Reflection, Phase Shift, Scattering, Emission
- **Detection** Your eye, Light sensitive film, CCD / CMOS, Scintillator, Transducer



Fig. 4.4: The parts involved in the image formation process probing a sample.

#### 4.7.1 Where do images come from?

Modality	Impulse	Characteristic	Response	Detection
Light Microscopy	White Light	Electronic interactions	Absorption	Film, Camera
Phase Contrast	Coherent light	Electron Density (Index of Refraction)	Phase Shift	Phase stepping, holography, Zernike
Confocal Microscopy	Laser Light	Electronic Transition in Fluorescence Molecule	Absorption and reemission	Pinhole in focal plane, scanning detection
X-Ray Radiography	X-Ray light	Photo effect and Compton scattering	Absorption and scattering	Scintillator, microscope, camera
Neutron Radiography	Neutrons	Interaction with nucleus	Scattering and absorption	Scintillator, optics, camera
Ultrasound	High frequency sound waves	Molecular mobility	Reflection and Scattering	Transducer
MRI	Radio-frequency EM	Unmatched Hydrogen spins	Absorption and reemission	RF coils to detect
Atomic Force Microscopy	Sharp Point	Surface Contact	Contact, Repulsion	Deflection of a tiny mirror

## 4.8 Acquiring Images

### 4.8.1 Traditional / Direct imaging

- Visible images produced or can be easily made visible
- Optical imaging, microscopy

```
bone_img = imread('figures/tiny-bone.png').astype(np.float32)
# simulate measured image
conv_kern = np.pad(disk(2), 1, 'constant', constant_values = 0)
meas_img = convolve(bone_img[:-1], conv_kern)
# run deconvolution
dekern = np.fft.ifft2(1/np.fft.fft2(conv_kern))
```

(continues on next page)

(continued from previous page)

```

rec_img = convolve(meas_img, dekern) [::-1]
# show result
fig, (ax_orig, ax1, ax2) = plt.subplots(1,3, figsize = (15, 5))
ax_orig.imshow(bone_img, cmap = 'bone'); ax_orig.set_title('Original Object')
ax1.imshow(np.real(meas_img), cmap = 'bone'); ax1.set_title('Measurement')
ax2.imshow(np.real(rec_img), cmap = 'bone', vmin = 0, vmax = 255); ax2.set_title(
    'Reconstructed');

```



## 4.8.2 Indirect / Computational imaging

- Recorded information does not resemble object
- Response must be transformed (usually computationally) to produce an image

```

bone_img = imread('figures/tiny-bone.png').astype(np.float32)
# simulate measured image
meas_img = np.log10(np.abs(np.fft.fftshift(np.fft.fft2(bone_img))))
print(meas_img.min(), meas_img.max(), meas_img.mean())
fig, (ax1, ax_orig) = plt.subplots(1,2,
                                    figsize = (12, 6))
ax_orig.imshow(bone_img, cmap = 'bone')
ax_orig.set_title('Original Object')

ax1.imshow(meas_img, cmap = 'hot')
ax1.set_title('Measurement');

```

```
1.146423838816545 6.61125552595089 3.3563935033662253
```



#### 4.8.3 Traditional Imaging

### 4.9 Traditional Imaging: Model



$s_{ab}$  is the only information you are really interested in, so it is important to remove or correct for the other components. For color (non-monochromatic) images the problem becomes even more complicated.  
 $\int_0^\infty [[(b(x, y, \lambda) * s_{ab}(x, y, \lambda)) \otimes h_{fs}(x, y, \lambda)] * h_{op}(x, y, \lambda)] * h_{det}(x, y, \lambda) d\lambda + d_{dark}(x, y)$

### 4.9.1 Indirect Imaging (Computational Imaging)

With indirect imaging you make acquisitions in a form that don't represent the information you want to have. It is needed to perform a numeric transformation to obtain images in observation space.

Some examples are:

- Tomography through projections
- Microlenses [Light-field photography](#)
- Diffraction patterns
- Hyperspectral imaging with Raman, IR, CARS
- Surface Topography with cantilevers (AFM)
- Tomography through projections
- Microlenses [Light-field photography](#)
- Diffraction patterns
- Hyperspectral imaging with Raman, IR, CARS
- Surface Topography with cantilevers (AFM)



## DIFFERENT VIEWS ON IMAGE ANALYSIS

Image analysis is a complex task and there are many ways to reach the quantitative results from the images.



Fig. 5.1: Your background often decides how you approach an image analysis problem.

We can make two initial statements:

- An image is a bucket of pixels.
- How you choose to turn it into useful information is strongly dependent on your background

## **5.1 Image Analysis: Experimentalist**

The experimentalist looks with a problem driven concept on the analysis task. It is often a top down approach aiming at solving the specific problem at hand. The solution is often reality driven and aims at finding models explaining the information presented in the images.

Typical task the experimentalist tries to solve are very practical and specific like counting cells in the image or to measure the porosity of the sample.

### **5.1.1 Characteristics**

- Problem-driven
- Top-down
- *Reality* Model-based

### **5.1.2 Examples**

- cell counting
- porosity

## **5.2 Image Analysis: Computer Vision Approaches**

The computer vision/signal processing scientist works to develop methods to solve a class of image processing problem. The approach is based on abstract features found in the image. The models are based on features and noise found in the images. The systematic approach is even based on engineered image features to better test and evaluate the developed methods.

The computer vision approach is typical looking to detect features like edges, structures, and also complicated features like faces.

### **5.2.1 Characteristics**

- Method-driven
- Feature-based
- *Image* Model-based
- Engineer features for solving problems

## 5.2.2 Examples

- Edge detection
- Face detection

## 5.3 Image Analysis: Deep Learning Approach

Finally, the deep learning approach is data driven and inspired by the way nature solves the image analysis problem. This approach rebuilds the way image processing is done from scratch, but at the same time it also based on concepts developed in computer vision. The deep learning approaches doesn't require a specific model describing the images it is meant to analyze, but rather make conclusions based on the previous images it has been exposed to.

The deep learning approach is good handling rare events in the data and when it trained correctly it is also capable of generalizing to detect new features. This may sound like magic, but this is also a well founded and structured approach to analyzing images. Care must however be taken not to over fit or to generalize too much. The models are never better than the data they have been exposed to.

Examples where deep learning is frequently used are to detect anomalies in the data or to label images based on their contents.

### 5.3.1 Characteristics

- Results-driven
- Biology 'inspired'
- Build both image processing and analysis from scratch

### 5.3.2 Examples

- Captioning images
- Identifying unusual events

## 5.4 Summary analysis approaches

These three approaches have their own advantages and disadvantages, therefore it is good to know them all to be able to select the adequate method for the task you have to solve. It is not unusual that you will have to use a mix of the approaches. It is important to be open minded and think outside the box. In the end, what matters is that you can provide a reliable analysis of your data.



---

CHAPTER  
SIX

---

ON SCIENCE

## 6.1 What is the purpose?

- Discover and validate new knowledge

### 6.1.1 How?

- Use the scientific method as an approach to convince other people
- Build on the results of others so we don't start from the beginning

### 6.1.2 Important Points

- While **qualitative** assessment is important, it is difficult to reliably produce and scale
- **Quantitative** analysis is far from perfect, but provides metrics which can be compared and regenerated by anyone

Inspired by: [imagej-pres](#)

## 6.2 Science and Imaging

Images are great for qualitative analyses since our brains can quickly interpret them without large *programming* investments.

### 6.2.1 Proper processing and quantitative analysis is however much more difficult with images.

- If you measure a temperature, quantitative analysis is easy,  $50K$ .
- If you measure an image it is much more difficult and much more prone to mistakes, subtle setup variations, and confusing analyses

## 6.2.2 Furthermore in image processing there is a plethora of tools available

- Thousands of algorithms available
- Thousands of tools
- Many images require multi-step processing
- Experimenting is time-consuming

## 6.3 Important questions



Fig. 6.1: A realistic rendering of an ancient gold pendant.

- It is important to define how you want to use your images
- Nice renderings is mostly not sufficient for a publication

### 6.3.1 Initial questions

You need to ask yourself a number of questions before you start to plan an experiment that involves images. It is good to at least have a plan about how you are going to analyze your images once you have them. It is not always obvious how you get from the experiment idea, to performing the experiment and finally how you perform the analysis to obtain the information you are really interested in for your research project.

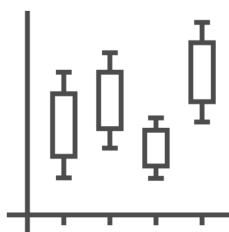
- What am I looking for?
- Is my original question specific or too general?
- How can I plan my experiment to make the analysis easier?
- How can I transfer the original question into the vocabulary of image analysis?
- How much a priori information about the sample do I have? Can it be used for the analysis?



Fig. 6.2: Material analysis of a rusty roman nail.



Compare treatments



Understand and model processes



Inspect components

Fig. 6.3: Things a scientist wants to know from an experiment.

### 6.3.2 What is the purpose of the experiment

- 3D visualization
- Sample characterization
- Model process parameters
- etc.

### 6.3.3 What can we measure?

There is a leap between what you really want to know and what we actually can measure in the images.

Quantification from gray values	Quantification from shapes
Material composition	Identify and count items
Material transport	Measure volume
	Characterize shape



Find hidden objects



Dimensions



Counting objects



Mixing ratios



Track changes

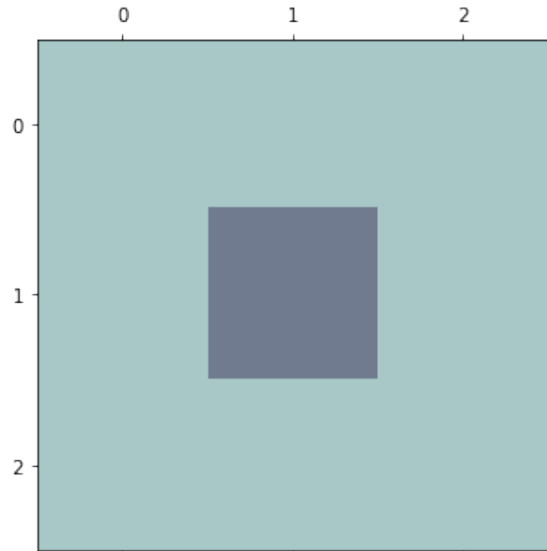
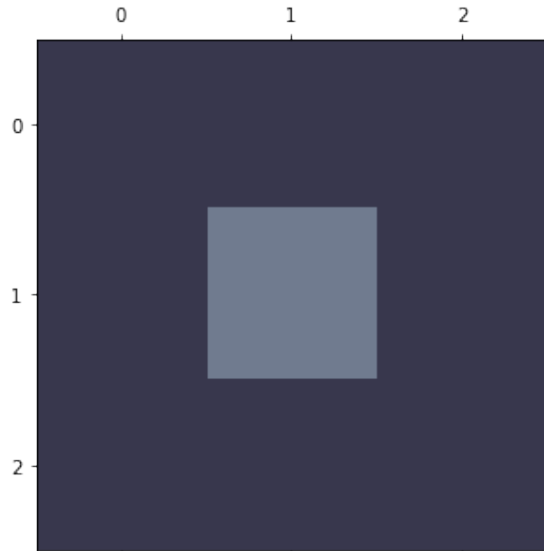
Fig. 6.4: These are things we can measure in images.

## 6.4 Why quantitative?

### 6.4.1 Human eyes have issues

Which center square seems brighter?

```
%matplotlib inline
import matplotlib.pyplot as plt
import numpy as np
xlin = np.linspace(-1,1, 3)
xx, yy = np.meshgrid(xlin, xlin)
img_a = 25*np.ones((3,3))
img_b = np.ones((3,3))*75
img_a[1,1] = 50
img_b[1,1] = 50
fig, (ax1, ax2) = plt.subplots(1,2, figsize = (12, 5));
ax1.matshow(img_a, vmin = 0, vmax = 100, cmap = 'bone');
ax2.matshow(img_b, vmin = 0, vmax = 100, cmap = 'bone');
```



## 6.5 Intensity gradients

Are the intensities constant in the image?

```
%matplotlib inline
import matplotlib.pyplot as plt
import numpy as np
xlin = np.linspace(-1,1, 10)
xx, yy = np.meshgrid(xlin, xlin)

fig, ax1 = plt.subplots(1,1, figsize = (6, 6))
ax1.matshow(xx, vmin = -1, vmax = 1, cmap = 'bone');
```



## 6.6 Selecting the right tool for your task

### 6.6.1 Criteria

There are many questions you have to ask about the data and analysis task you are facing. Here are some relevant questions that will guide you towards the choice of tool(s) you need to use when you analyses the data.

- How many samples?
- How complex is the sample?
- Is human interpretation needed?
- What is the end product?
- Are there methods tools available?
- Will there be more similar data sets?

### 6.6.2 The choice

You have different options to choose the tool for your analysis.

- Interactive tools
- Scripting using existing toolboxes
- Development of new algorithms Which one you actually select for your analysis workflow depends on many factors like the criteria listed in the previous section, but also on your experience, preferences in the group you are working with and so on.

In general, there is **no golden recipe!** You have to be flexible from task to task.

**"I SPEND A LOT OF TIME ON THIS TASK.  
I SHOULD WRITE A PROGRAM AUTOMATING IT!"**

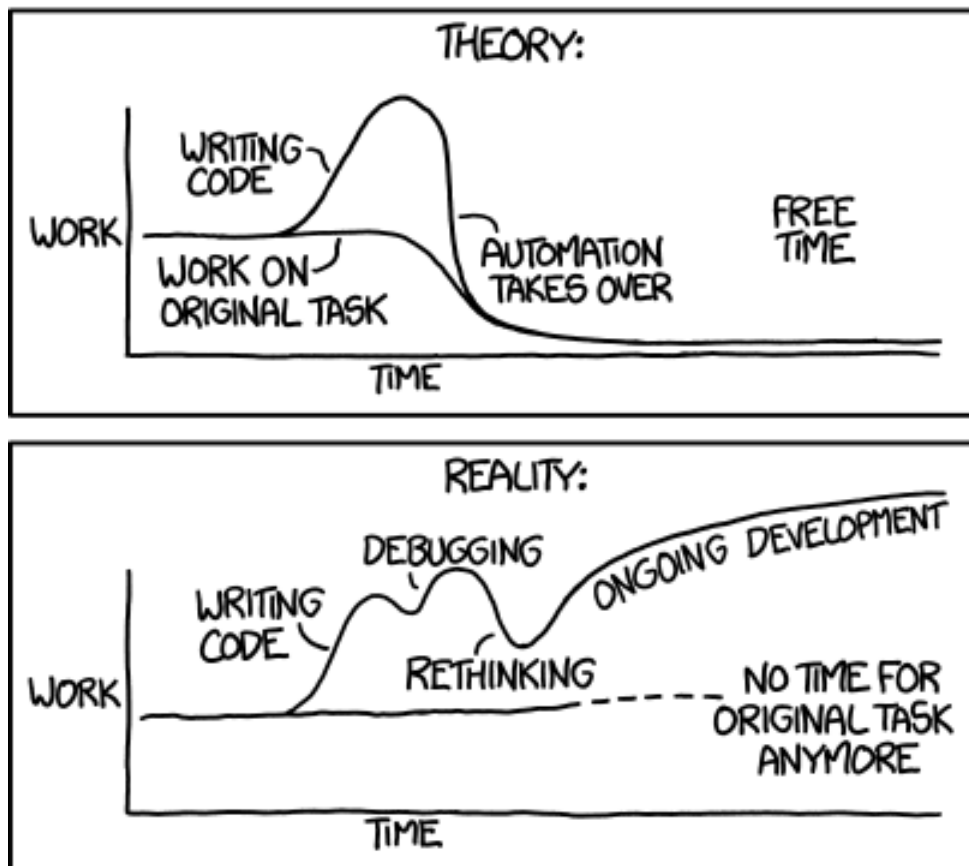


Fig. 6.5: It sometimes feels deievingly simple to implement a script that automates your task... be careful!

### 6.6.3 Criteria

- How many samples?
- How complex is the sample?
- Is human interpretation needed?
- What is the end product?
- Are there methods/tools available?
- Will there be more similar data sets?

### 6.6.4 The choice

There is **no golden recipe!**

- Interactive tools
- Scripting using existing toolboxes
- Development of new algorithms

## 6.7 Reproducibility vs. Repeatability

### 6.7.1 Reproducibility

To reproduce an experiment means that someone else can repeat your entire workflow from sample preparation and experiment to the analysis to obtain the same results you got in your report. This should be possible completely independently for an experiment to be called reproducible.

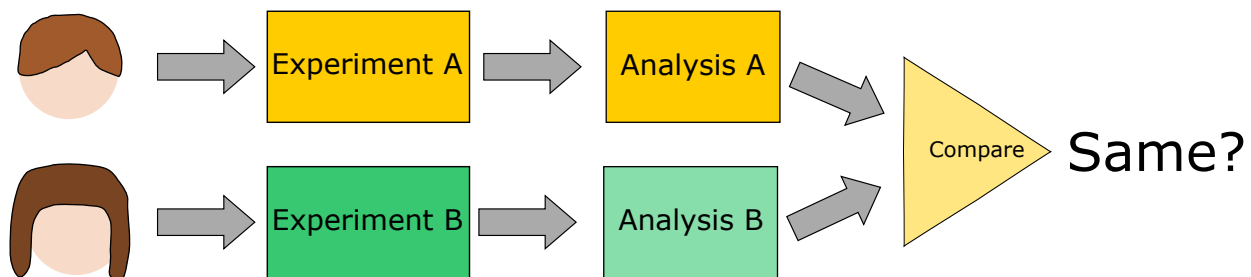


Fig. 6.6: A workflow describing the concept of reproducibility.

### 6.7.2 Repeatability

Repeatability is a more relaxed concept. It requires that someone else (or even yourself) can obtain the same results multiple time using the same data and analysis workflow. This means that there must very little room left for individual decisions that may have impact on the outcome.



Fig. 6.7: A workflow describing the concept of reproducibility.

## 6.8 Reproducibility vs. Repeatability

Science demands **repeatability!** and really wants **reproducibility**

- Experimental conditions can change rapidly and are difficult to make consistent
  - Animal and human studies are prohibitively time consuming and expensive to reproduce
  - Terabyte datasets cannot be easily passed around many different groups
  - Privacy concerns can also limit sharing and access to data
- 
- *Science* is already difficult enough
  - Image processing makes it even more complicated
  - Many image processing tasks are multistep, have many parameters, use a variety of tools, and consume a very long time

## 6.9 How can we keep track of everything for ourselves and others?

We can make the data analysis easy to repeat by an independent 3rd party

- Document the analysis steps
- Write clear and understandable code

## WORKFLOWS FOR IMAGE ANALYSIS

### 7.1 Computing has changed: Parallel

#### 7.1.1 Moores Law

$$\text{Transistors} \propto 2^{T/(18 \text{ months})}$$

```
# Borrowed from https://gist.github.com/humberto-ortiz/de4b3a621602b78bf90d
moores_txt=[ "Id Name Year Count(1000s) Clock(MHz)\n",
    "0      MOS65XX 1975      3.51      14\n",
    "1      Intel8086 1978     29.00      10\n",
    "2      MIPSR3000 1988     120.00     33\n",
    "3      AMDAm486 1993    1200.00    40\n",
    "4      NexGenNx586 1994   3500.00   111\n",
    "5      AMDAthlon 1999   37000.00  1400\n",
    "6      IntelPentiumIII 1999  44000.00 1400\n",
    "7      PowerPC970 2002   58000.00  2500\n",
    "8      AMDAthlon64 2003  243000.00 2800\n",
    "9      IntelCore2Duo 2006 410000.00 3330\n",
    "10     AMDPhenom 2007  450000.00 2600\n",
    "11     IntelCorei7 2008  1170000.00 3460\n",
    "12     IntelCorei5 2009  995000.00 3600"]

sio_table = StringIO(''.join(moorees_txt)); moore_df = pd.read_table(sio_table, sep =
    '\s+', index_col = 0);
fig, ax1 = plt.subplots(1,1, figsize = (8, 4)); ax1.semilogy(moore_df['Year'], moore_
    df['Count(1000s)'], 'b.-', label = '1000s of transitions'); ax1.semilogy(moore_df[
    'Year'], moore_df['Clock(MHz)'], 'r.-', label = 'Clockspeed (MHz)') ;ax1.legend(loc_
    = 2);
```



Based on data from <https://gist.github.com/humberto-ortiz/de4b3a621602b78bf90d>

### 7.1.2 Why doesn't the clock rate follow Moore's law?

There are now many more transistors inside a single computer but the CPU clock rate hasn't increased. It is actually stagnating since the early 2000s. The reason for this is that the number of processing cores has started to increase. Each core is a CPU is an autonomous CPU which makes it possible to perform many individual tasks in parallel.

- Multiple Core
- Multiple CPUs
- New modalities
  - GPUs provide many cores which operate at slow speed

The conclusion is that we need to make the processing on multiple Cores/CPUs possible. This can be done by choosing multithreaded modules for the processing. In the extreme case we need to implement new multi-process code for the analysis. How this is done depends in the target hardware. You will use different approaches for a CPU than you use in a cluster or in cloud computing. In general, we can conclude:

**Parallel Code is important**

## 7.2 Computing has changed: Cloud

Cloud computing is a type of service that has evolved from the big data era. Large companies have built up their analysis resources to match the need at peak demand. These resources were often underused which resulted in a new business model. Providing computing resources to external customers. In this way the customers only pay for the computing time they need and let someone else handle investments and system maintenance.

- Computer, servers, workstations are *wildly underused* (majority are <50%)
- Buying a big computer that sits *idle most of the time* is a waste of money
  - <http://www-inst.eecs.berkeley.edu/~cs61c/sp14/>



Fig. 7.1: Different cloud services where it is possible to buy CPU time.

– “The Case for Energy-Proportional Computing,” Luiz André Barroso, Urs Hözle, IEEE Computer, December 2007

- Traditionally the most important performance criteria was time, how fast can it be done
- With Platform as a service servers can be *rented instead of bought*
- Speed is still important but using cloud computing \$ / Sample is the real metric
- In Switzerland a PhD student is 400x as expensive per hour as an Amazon EC2 Machine
- Many competitors keep prices low and offer flexibility

## 7.3 Workflow analysis

It sounds like a great idea to perform tasks in parallel, but how do we bring the computer to do so. The following example from real life is used to demonstrate how a greater task can be broken down into smaller pieces.

### 7.3.1 Simple Soup

We want to cook a simple soup. There is already a recipe telling us what to do to produce a soup. It is easy to follow the list, anyone with the right steps can execute and repeat (if not reproduce) the soup

1. Buy {carrots, peas, tomatoes} at market
2. *then* Buy meat at butcher
3. *then* Chop carrots into pieces
4. *then* Chop potatoes into pieces
5. *then* Heat water
6. *then* Wait until boiling then add chopped vegetables
7. *then* Wait 5 minutes and add meat

All steps are done in a sequence to produce this soup. This would correspond to performing the task on a single CPU. Now, how can we change this?

### 7.3.2 More complicated soup

The following recipe is harder to follow and you need to carefully keep track of what is being performed. If you look at it in detail, you will see that some of the steps can be performed independently of the others. This is our opportunity to dispatch parallel tasks that would be running on different CPUs. In the kitchen, you could delegate tasks to other people and thus shorten the time until the soup is ready.

**Steps 1-4** 4. *then* Mix carrots with potatos →  $mix_1$  4. *then* add egg to  $mix_1$  and fry for 20 minutes 4. *then* Tenderize meat for 20 minutes 4. *then* add tomatoes to meat and cook for 10 minutes →  $mix_2$  5. *then* Wait until boiling then add  $mix_1$  6. *then* Wait 5 minutes and add  $mix_2$

## 7.4 Using flow charts / workflows

The processing task can be visualize using flow charts to better get an overview of the steps to be done. A sequence has a trivial shape, but it hard to get the big picture as soon as you start creating more complicated schemes .

### 7.4.1 Workflow of the Simple Soup

In this example we use the graph plotting module graphviz to draw the flow chart of the soup example. Graphviz is only a plotting tool with no direct processing capacity.

```
from IPython.display import Image, display
import pydot
graph = pydot.Dot(graph_type='digraph', rankdir="LR")
node_names = ["Buy\nvegetables", "Buy meat", "Chop\nvegetables", "Heat water", "Add_\nVegetables",
              "Wait for\nboiling", "Wait 5\nadd meat"]
nodes = [pydot.Node(name = '%04d' % i, label = c_n)
         for i, c_n in enumerate(node_names)]
for c_n in nodes:
    graph.add_node(c_n)

for (c_n, d_n) in zip(nodes, nodes[1:]):
    graph.add_edge(pydot.Edge(c_n, d_n))

plt = Image(graph.create_png()); display(plt);
```



### 7.4.2 Workflows - the complicated soup

Clearly a linear set of instructions is ill-suited for even a fairly easy soup, it is then even more difficult when there are dozens of steps and different pathways.

Furthermore a clean workflow allows you to better parallelize the task since it is clear which tasks can be performed independently.

```
from IPython.display import Image, display
import pydot
graph = pydot.Dot(graph_type='digraph', rankdir="LR")
node_names = ["Buy\nvegetables", "Buy meat", "Chop\nvegetables", "Heat water", "Add_\nVegetables",
```

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```

    "Wait for\nboiling", "Wait 5\nadd meat"]
nodes = [pydot.Node(name = '%04d' % i, label = c_n, style = 'filled')
         for i, c_n in enumerate(node_names)]
for c_n in nodes:
    graph.add_node(c_n)

def e(i, j, col = None):
    if col is not None:
        for c in [i, j]:
            if nodes[c].get_fillcolor() is None:
                nodes[c].set_fillcolor(col)
    graph.add_edge(pydot.Edge(nodes[i], nodes[j]))

e(0, 2, 'gold'); e(2, 4); e(3, -2, 'springgreen'); e(-2, 4, 'orange'); e(4, -1) ; e(1,
→ -1, 'dodgerblue')
plt = Image(graph.create_png()); display(plt);

```



## 7.5 Directed Acyclical Graphs (DAG)

- We can represent almost any computation without loops as DAG.
- This allows us to break down a computation into pieces which can be carried out independently.

There are a number of tools which let us handle this issue.

- PyData Dask - <https://dask.pydata.org/en/latest/>
- Apache Spark - <https://spark.apache.org/>
- Spotify Luigi - <https://github.com/spotify/luigi>
- Airflow - <https://airflow.apache.org/>
- KNIME - <https://www.knime.com/>
- Google Tensorflow - <https://www.tensorflow.org/>
- Pytorch / Torch - <http://pytorch.org/>

### 7.5.1 Concrete example - Creating a DAG

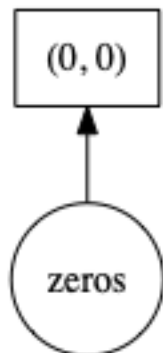
What is a DAG good for?

#### Create two variables

Let's create a DAG to demonstrate how it works. This first piece creates two small images (5x5 pixels) each containing either '0's or '1's. We still haven't started any calculations using this DAG.

```
import dask.array as da

from dask.dot import dot_graph
image_1 = da.zeros((5,5), chunks = (5,5))
image_2 = da.ones((5,5), chunks = (5,5))
dot_graph(image_1.dask)
```



#### Add two variables

Adding two variables in a DAG looks like the code to add any variables. Now if we look at the graph created by this operation, we see that there is a directed flow of the data in this graph.

$$Image_3 = Image_1 + Image_2$$

```
image_3 = image_1 + image_2
dot_graph(image_3.dask)
```



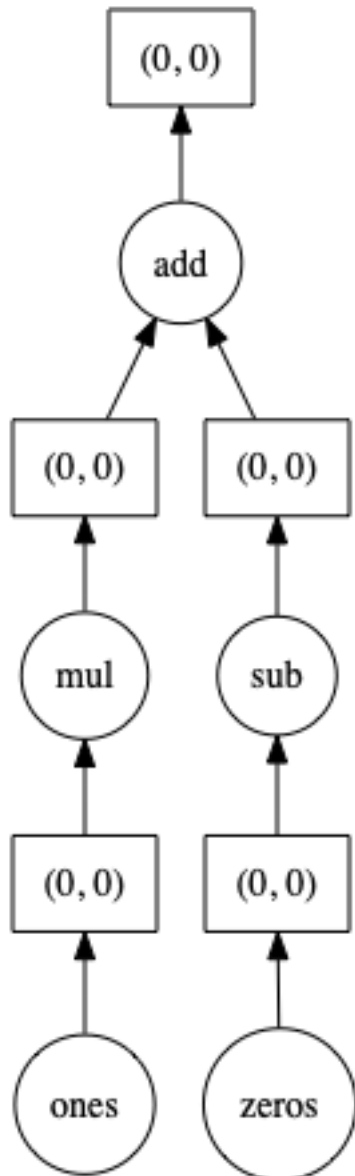
### A more complicated operation

Now lets make a more complicated calculation like:

$$\text{Image}_4 = (\text{image}_1 - 10) + (\text{image}_2 * 50)$$

```

image_4 = (image_1-10) + (image_2*50)
dot_graph(image_4.dask)
  
```



We can here see that there are two clear branches in the flow chart. This indicates that here is an opportunity to perform these tasks in parallel.

## 7.5.2 Let's go big

Now let's see where this can be really useful. Dask allows you to split the data into smaller pieces (chunks). In the example below we have created a 1024x1024 pixels image and also told dask that this image should be divided into chunks of 512x512 pixels.

### Creating a large image with chunks

```
import dask.array as da
from dask.dot import dot_graph
image_1 = da.zeros((1024, 1024), chunks = (512, 512))
image_2 = da.ones((1024, 1024), chunks = (512, 512))
dot_graph(image_1.dask)
```



You see now that there are four graph created. The boxes on the top indicates the location of the chunk in the large image.

### Computing something with a larger image

If we now apply the “complicated” calculation

$$Image_4 = (image_1 - 10) + (image_2 * 50)$$

on the large image with chunks, we see that the code hasn't changed at all. When we look at the graph generated with this data, we see that there are four graphs repeating the same operations. One graph for each chunk we defined in the data.

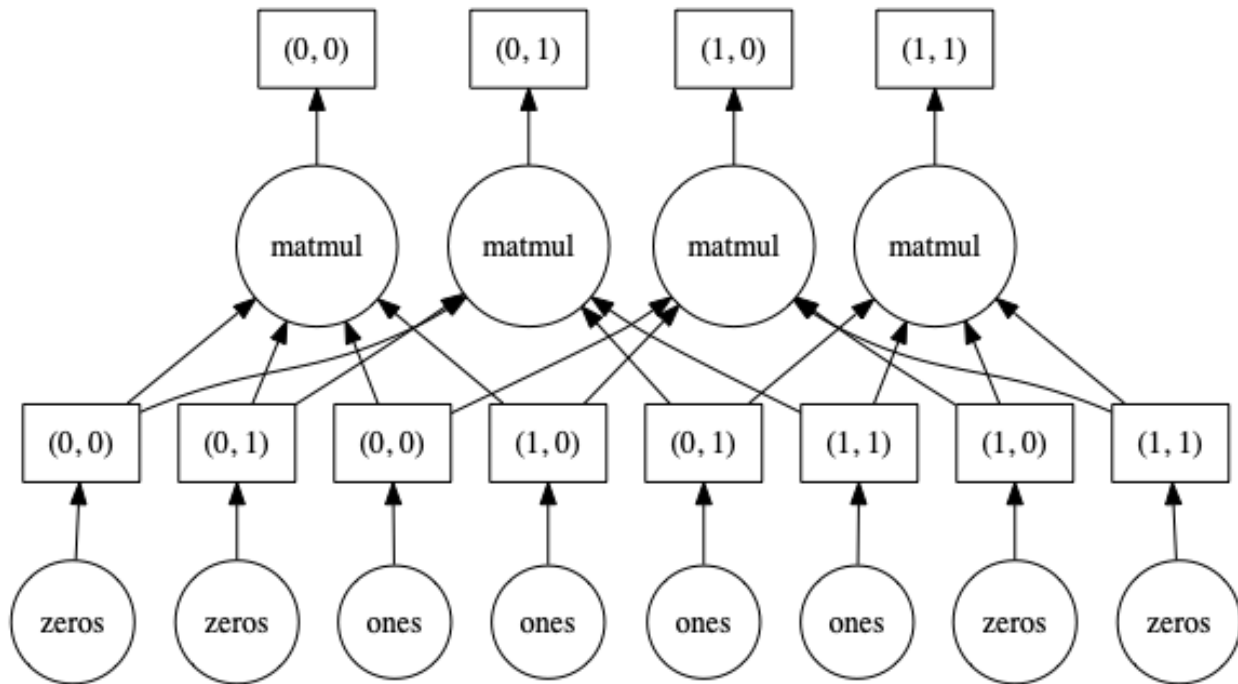
```
image_4 = (image_1-10) + (image_2*50)
dot_graph(image_4.dask)
```



### 7.5.3 Matrix multiplication in a DAG

A matrix multiplication is a more complicated operation than the previous examples. This operation needs information from all the other chunks to calculate the current chunk. This can also be seen in the wiring of the graph below.

```
image_5 = da.matmul(image_1, image_2)
dot_graph(image_5.dask)
```



#### 7.5.4 An even more complicated computation

Combining different operations into a final calculation like

$$Image_6 = (Image_1 \cdot Image_2 + Image_1) \circ Image_2$$

```
image_6 = (da.matmul(image_1, image_2)+image_1)*image_2
dot_graph(image_6.dask)
```



We see here that the convolution part connects all branches of the graph which mean all processing must be synchronized here. Afterwards there are no connections and each sub graph can work independently.

### 7.5.5 Convolution using a DAG

Convolution makes life harder for a DAG. There is a need for an overlap between the chunks to avoid boundary effects. These boundary effects are something we will look into in next week's lecture about filters.

$$Image_7 = Image_6 * Image_1$$

```
import dask_ndfilters as da_ndfilt
image_7 = da_ndfilt.convolve(image_6, image_1)
dot_graph(image_7.dask)
```



## 7.6 Deep Learning

We won't talk too much about deep learning now, but it certainly shows why DAGs are so important.



The steps above are simple toys compared to what tools are already in use for machine learning

[https://keras.io/api/utils/model\\_plotting\\_utils/](https://keras.io/api/utils/model_plotting_utils/)

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**CHAPTER  
EIGHT**

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**SUMMARY**

In this lecture we saw that:

- Images reveal information about different samples
- Images are signals that need to be quantitatively analyzed
- Science with images is a non-trivial task
- Proper workflows are required for efficient analysis repeatable analysis.

## **8.1 Next weeks lecture**

- Noise
- Filters
- Filter evaluation