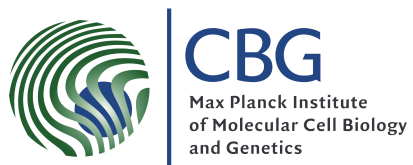
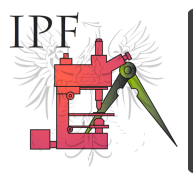
Basics of Quantitative Image Analysis

What you need to know about Image Processing ...

... but never thought to ask









Before you start writing...

See these slides at: https://ifn.mpi-cbg.de

under: Teaching

Also available on the Fiji Wiki

- ✓ Fiji is just ImageJ batteries included http://pacific.mpi-cbg.de
- √ Fiji tutorials
- ✓ DetectInfoLoss, ColocalisationAnalysis and more...
- ✓ Practicals etc. are included in online version...

Topics:

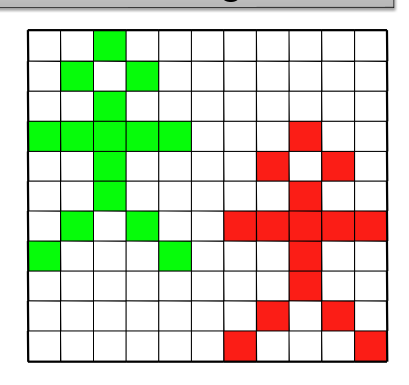
- ✓ Imaging Workflow
- ✓ Images = "Information" (Digital Images)
- ✓ What is a pixel?
- ✓ Info "about" the image = Meta Data
- ✓ Different ways to visualise / display image info

Quantitative Imaging? ...what does that mean?

Art or Science? Photography or Spectroscopy?

Science means to measure something!

- ✓ Numerical Results
- ✓ Statistics!
- ✓ Computers become useful...



What is Image Analysis / Quantification?

		l						
255	255	255	255	255	255	255	255	255
255	255	255	50	50	50	50	255	255
255	255	50	50	50	50	50	255	255
255	255	50	50	50	50	50	255	255
255	255	72	50	50	50	50	255	255
255	255	255	50	50	50	255	255	255
50	50	50				50	50	255
								255
200	233	255	255	30	200	200	200	200
255	255	255	50	255	255	255	255	255
255	255	255	50	50	50	50	51	168
255	255	255	50	255	255	255	255	255
255	255	50	255	255	255	255	255	255
255	255	50	255	255	255	255	255	255
255	50	255	255	255	255	255	255	255
	255 255 255 255 255 255 255 255 255	255 255 255 255 256 255 257 255 258 255 258 255 258 255 258 255 258 255	255 255 50 255 255 72 255 255 255 50 50 50 255 255 255 255 255 255 255 255 255 255 255 255 255 255 255 255 255 50 255 255 50 255 255 50	255 255 50 50 255 255 50 50 255 255 72 50 255 255 255 50 50 50 50 50 255 255 255 255 255 255 255 50 255 255 255 50 255 255 50 255 255 255 50 255 255 255 50 255	255 255 50 50 50 255 255 50 50 50 255 255 72 50 50 255 255 255 50 50 50 50 50 50 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255 255 255 50 255 255	255 255 50 50 50 50 255 255 50 50 50 50 255 255 72 50 50 50 255 255 255 50 50 50 50 50 50 50 50 255 255 255 255 50 255 255 255 255 255 50 255 255 255 255 255 50 50 50 255 255 255 50 255 255 255 255 255 50 255 255 255 255 50 255 255 255 255 255 50 255 255 255 255 255 50 255 255 255 255 255 50 255 255 255 255 255	255 255 50 255 255 50 50 50 255 255 255 250 50 50 50 50 50 50 50 50 50 50 255 2	255 255 50 50 50 50 50 255 255 255 50 50 50 50 255 255 255 72 50 50 50 50 255 255 255 255 50 50 50 255 255 50 50 50 50 50 50 50 50 255 255 255 255 50 255 255 255 255 255 255 255 50 255 255 255 255 255 255 255 255 50 255 255 255 255 255 255 255 50 255 255 255 255 255 255 255 50 255 255 255 255 255 255 255 50 255 255 255 255

Minimum: 50

Maximum: 255

Mean: 94.5

Std.Dev.: 93.2

Area: 10x14

Pixels: 140

Pix <255: 42

Object: Stick man

Body: 1

Head: 1

Legs: 2 (1 lifted)

Arms: 2 (2 lifted)

Walking left to

right...

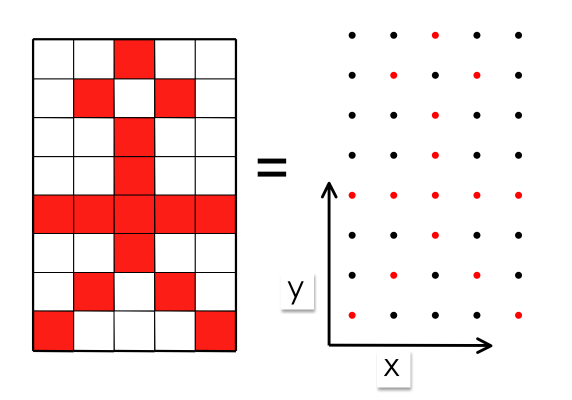




What is a (Digital) Image anyway..?

- ✓ it's a digital "representation" of reality!
- ✓ it's an artifact that contains less info than the object!
- ✓ it's just numbers! NOT analogue art!

The Image of a point is NOT a point!!! (Point Spread Function – PSF)



A digital image of ???

Image Analysis (Brain or Computer)

A stick man?
How do I know?
How can computer know
- algorithm?

Image = Information

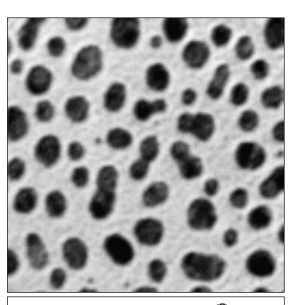
Images contain information!!!

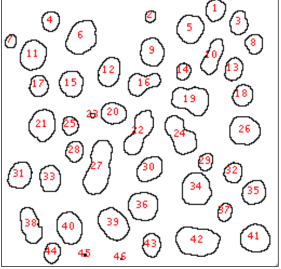
- ✓ Quantify / Measure / Analyse
- ✓ Meta data (what, where, when, how)
- ✓ Noise / Background

Manipulate Image = Changed Info!!!

Lost Info is lost forever

		1		1	1	T		1	
	Area	Mean	StdDev	Min	Max	IntDen	Median	XStart	YStart
1	285	255	0	255	255	72675	255	197	6
2	81	255	0	255	255	20655	255	136	17
3	278	255	0	255	255	70890	255	218	17
4	231	255	0	255	255	58905	255	42	18
5	501	255	0	255	255	127755	255	170	21
6	660	255	0	255	255	168300	255	75	26
7	99	255	0	255	255	25245	255	7	39
8	228	255	0	255	255	58140	255	231	39
9	448	255	0	255	255	114240	255	137	42
10	401	255	0	255	255	102255	255	198	43
11	520	255	0	255	255	132600	255	27	44
12	425	255	0	255	255	108375	255	99	60
13	271	255	0	255	255	69105	255	215	60
14	159	255	0	255	255	40545	255	168	65
15	412	255	0	255	255	105060	255	60	73
16	426	255	0	255	255	108630	255	123	75
17	260	255	0	255	255	66300	255	31	77
18	289	255	0	255	255	73695	255	222	85
19	676	255	0	255	255	172380	255	178	87





Slice	Count	Total Area	Average Size	Area Fraction
blobs.gif	46	17686.000000	384.478261	27.2

Image Data? What is it?

Intensity – Dye concentration??

Comparison of 2 colours / dyes / proteins??

Noisy Images?

Averaging?

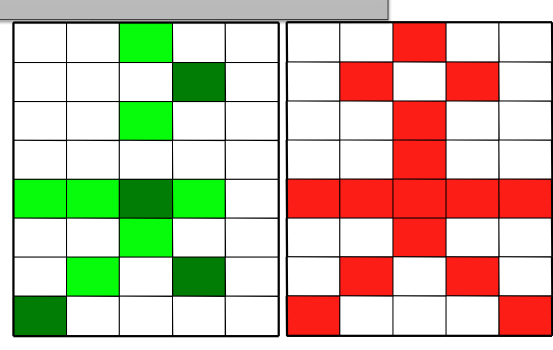
Pixel Time?

Shapes, Movement, Structure?

Internal controls!!!

A digital image with 2 channels / colours

What can you see here?

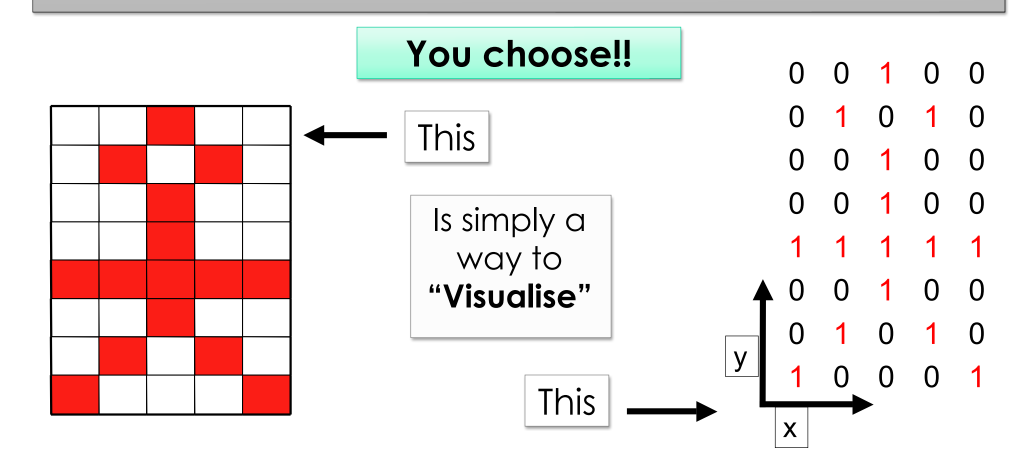


Photographer or Spectroscopist?

We can show you how to take pretty pictures (Art)

or

We can teach you how to get useful information (Science)



Photographer or Spectroscopist?

Science vs. Art

Objectivity vs. Subjectivity

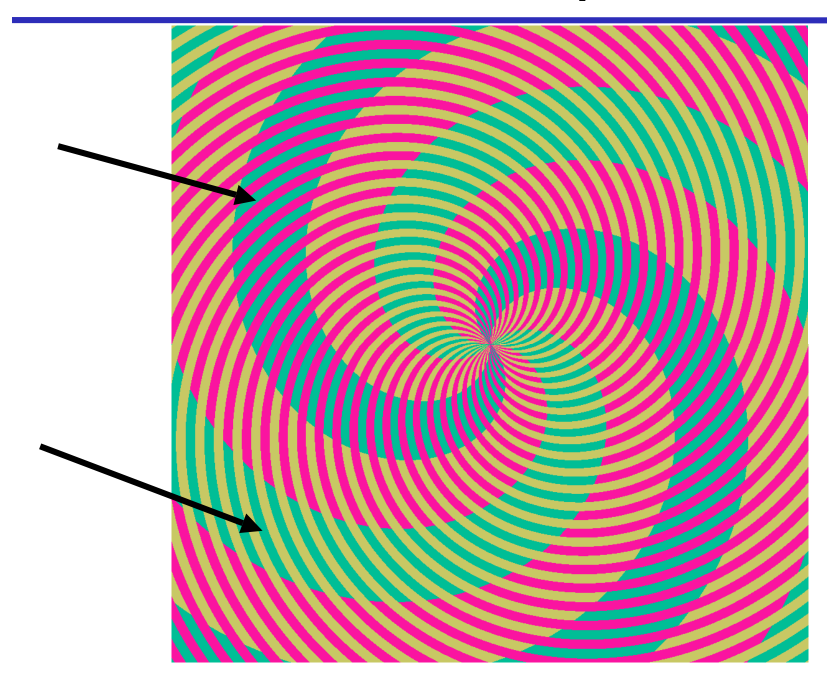
What I "think" I see vs. What is really there

Morphology can also be quantified!



249	244	240	230	209	233	227	251	255
248	245	210	93	81	120	97	193	254
250	170	133	94	137	120	104	145	253
241	116	118	107	134	138	96	92	163
277	142	121	113	124	115	107	71	179
234	106	84	125	97	108	125	106	204
241	202	102	132	75	73	141	246	252
253	252	244	239	178	199	242	250	245
255	249	244	250	226	231	240	251	253

Which colours can you see???



"Colour Merge" images could ruin your life

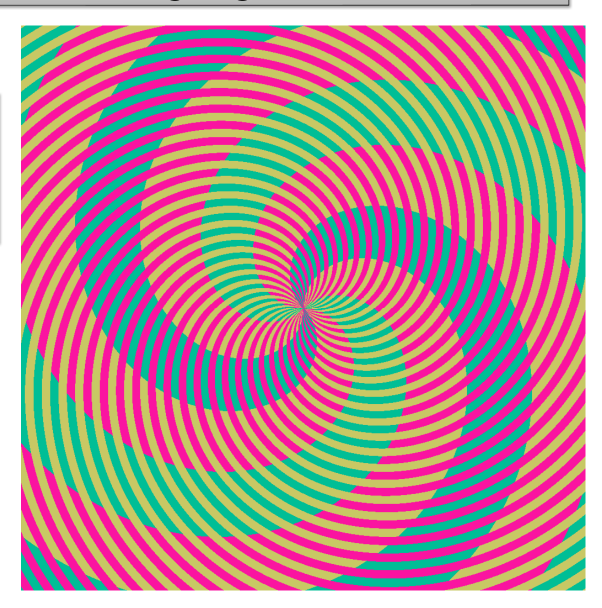
You see spirals, of pink, orange, green and blue?

Actually, the green and blue... are the same color!

Moral of the story:

Don't Trust Your

Eyes!

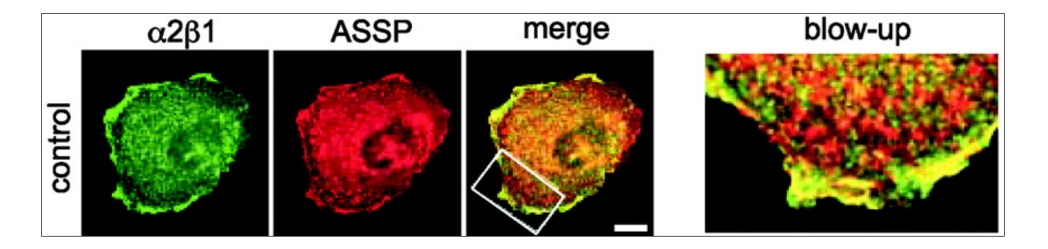


Colocalisation/Correlation

The past: "I see yellow - therefore there is colocalisation"

It is NOT possible to objectively decide about colocalisation by eye in a red-green merge image!

No colocalisation definition + No statistics = No Science



From Now On: 3D. Quantification. Correlation. Statistics.

Complementary methods: BioChemical, Optical (FRET, FLIM)

Colour Merge Images?

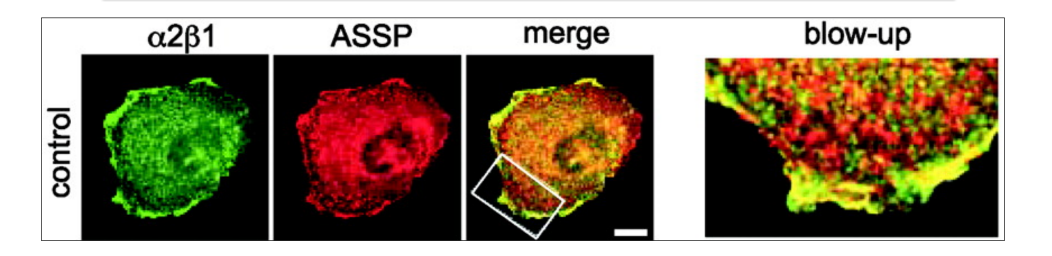
What are they good for?

Apart from looking pretty... not much.

Scientific conclusions from the image below? - No!

Colour blind people can't distinguish green and red!

So use Magenta and Green!



Publishing Images

or "how Photoshop can ruin your career"

CCD/PMT sees intensities differently than your eye/brain

LUT? Gamma correction?

Calibrate monitors

Journal Images ≠ Screen Images

Screen = RGB = Visualise

Inks = CMYK = Print

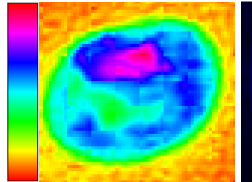
Compression

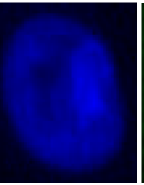
Lossless – Yes Lossy (JPEG) - NO

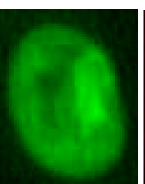
Image = data

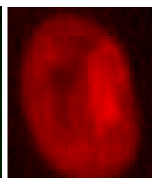
Don't corrupt information!

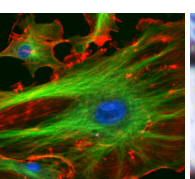
Always state the exact image processing done!







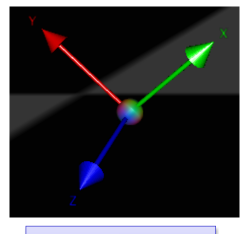




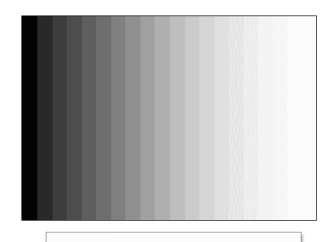


What can you digitise?

Dimensions!



SPACE

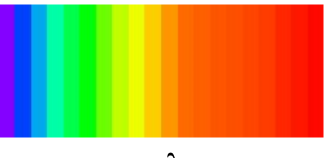


INTENSITY



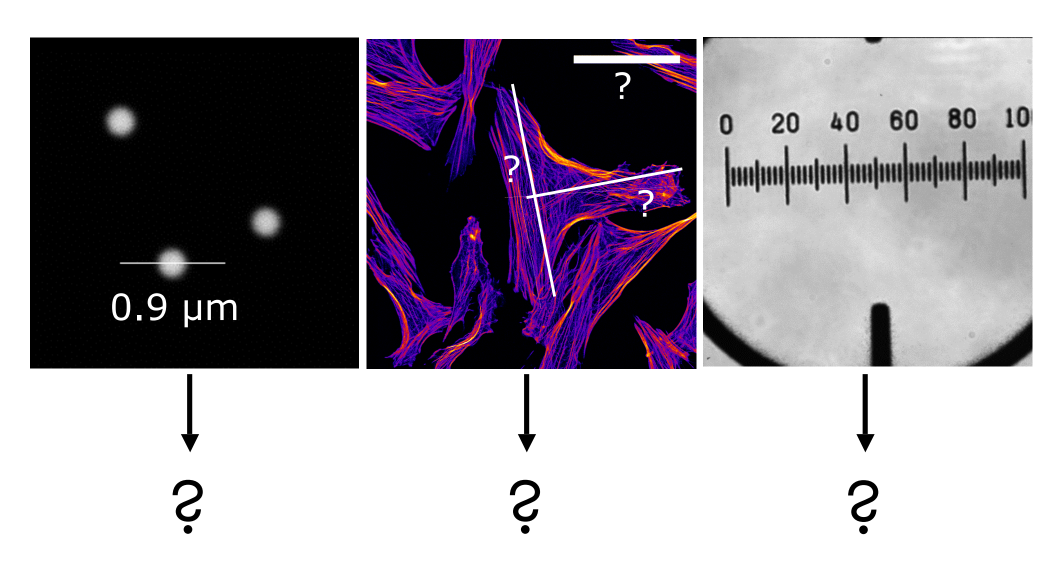
TIME

Wavelength

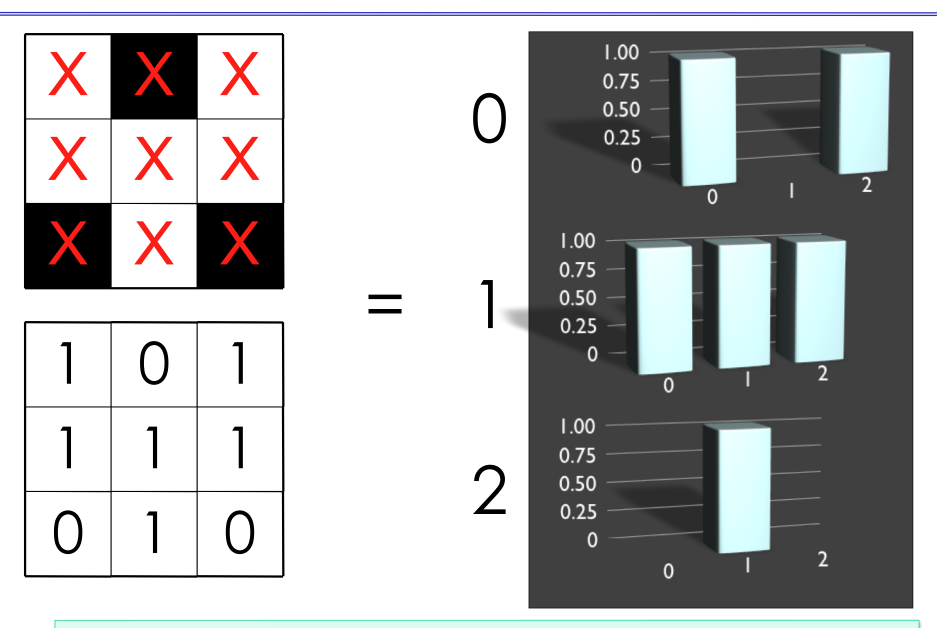


Colour

Pixel Size / Spatial Calibration



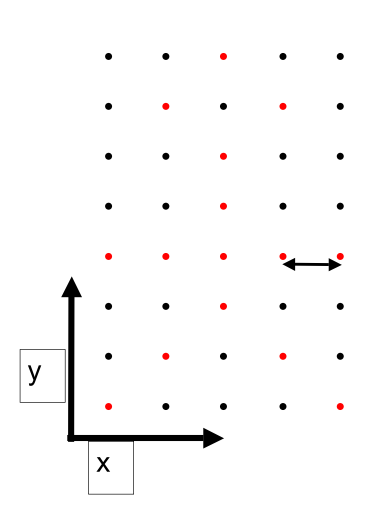
A pixel is NOT a little square!!!



A pixel is a point sample. It exists only at a point.

Digital spatial resolution

Projected pixel "size" at the sample/object is the point sample spacing



A pixel is not a "little square"

Point sample

=

Picture Element

_

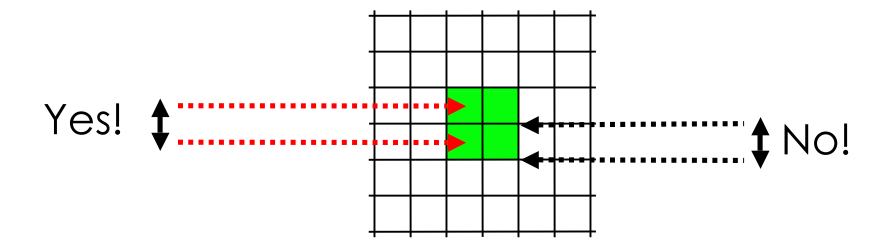
PixEl

Pixel Size

How big is a structure that is represented in my image?

How big is one pixel?

A pixel is NOT a little square!!!



- ✓ A pixel is a sample of "intensity" from a POINT in space
- ✓ "pixel size" is pixel spacing distance, not the imaginary pixel edge length!

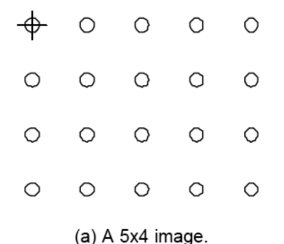
A pixel is NOT a little square,
A pixel is NOT a little square,
A pixel is NOT a little square!
(And a voxel is NOT a little cube)

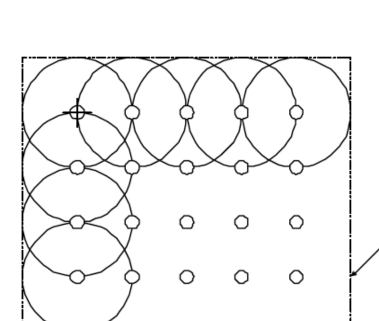
ftp://ftp.alvyray.com/Acrobat/6_Pixel.pdf

Alvy Ray Smith, July 17, 1995

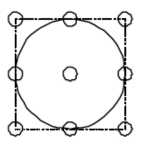
A pixel is a *point* sample. It exists only at a point.

Maybe it lies on a grid pattern...
...but that's accidental!





(c) Footprint of image under reconstruction.



(b) The footprint of a reconstruction filter. A truncated Gaussian, for example.

Or in our case the PSF
(Point spread function =
 image of a point)
 of the microscope
 system!

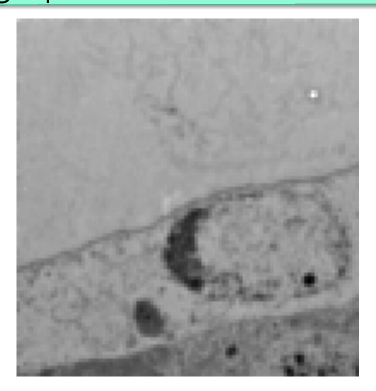
Dotted line is minimally enclosing rectangle

A pixel is not a little square ... So what?

Example – image shrinking 2048 x 2048 pixel electron micrograph – resized to 100 x 100



Wrong dumb interpolation of square pixels (aliased)



Correct
Gaussian smooth,
then down sample

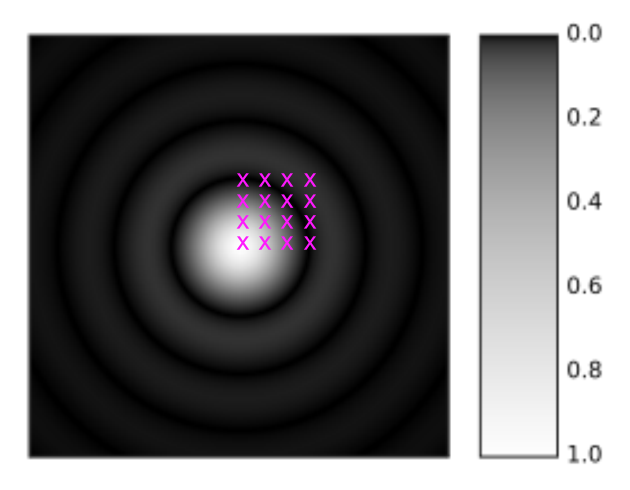
http://pacific.mpi-cbg.de/wiki/index.php/Downsample Compare plugins-examples-downsample with Image-scale

What does a point sample from a microscope detector contain?

Image of a point light source = Point Spread Function (PSF)

In the diffraction limited, high resolution case:

The PSF is **bigger** than the pixel / sample Nyquist spacing.



So what does a point sample from a confocal microscope detector contain?

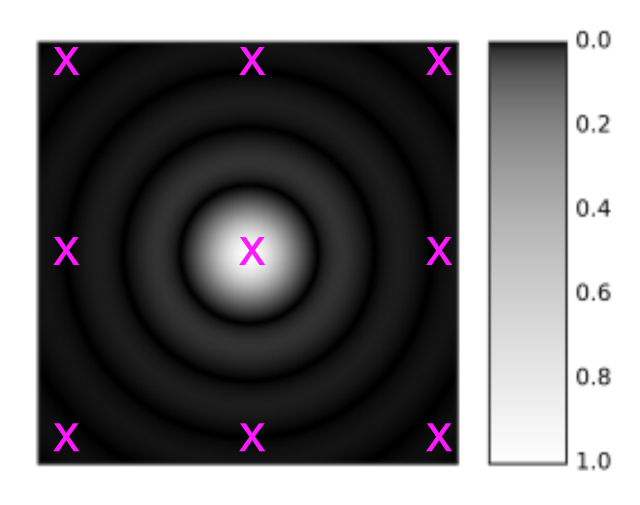
In the low resolution, big pixel case:

The PSF is

much smaller

than the pixel or
sample Nyquist
spacing.

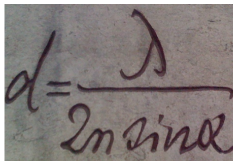
We miss spatial information = lower resolution



Abbe's diffraction limit / Rayleigh criterion



Limit the resolution of light microscopy



Airy Patterns and the Rayleigh Criterion online tutorial:

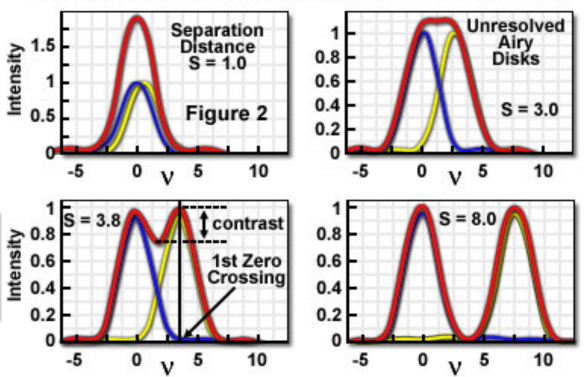
http://www.microscopy.fsu.edu/primer/java/imageformation/rayleighdisks/index.html

2 point light sources:

$$d = \frac{0.61 \times \lambda}{lens N.A.}$$

$$d = \frac{0.61 \times 550 \text{nm}}{1.4} = 240 \text{ nm}$$

Contrast and Resolution in Fluorescence Microscopy

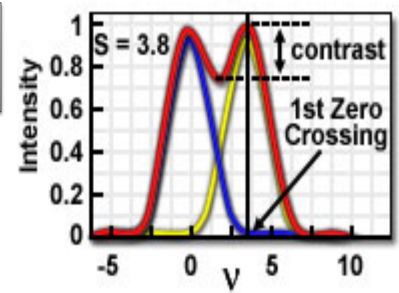


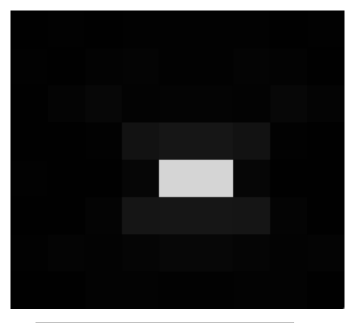
Digital spatial resolution

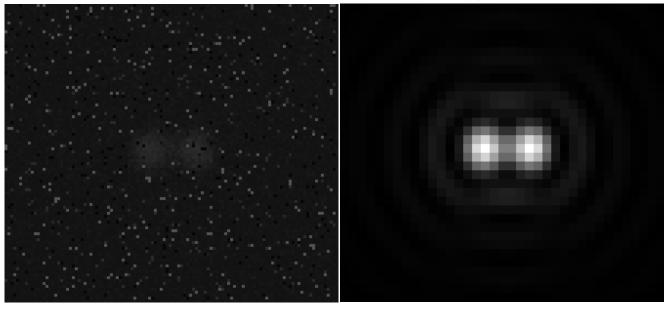
Projected pixel "size" at the sample/object

The point sample spacing

But what "should" it be?







under sampled

over sampled

good sampling

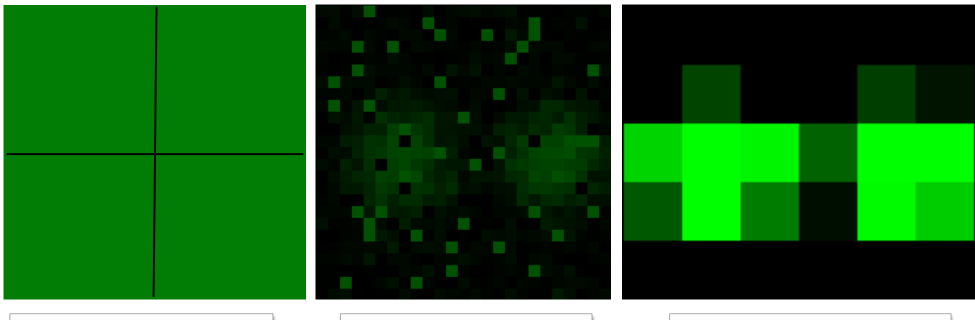
Pixel Size / Image Resolution

"Correct" image size? 64x64, 512x512, 2048x2048, ...

Nyquist - Shannon sampling theory: Proper spatial sampling

2.3 – 3 times smaller than optical resolution (x, y, AND z)

Adjust zoom, binning, and image size (no of pixels)



under sampled

over sampled

correct sampling

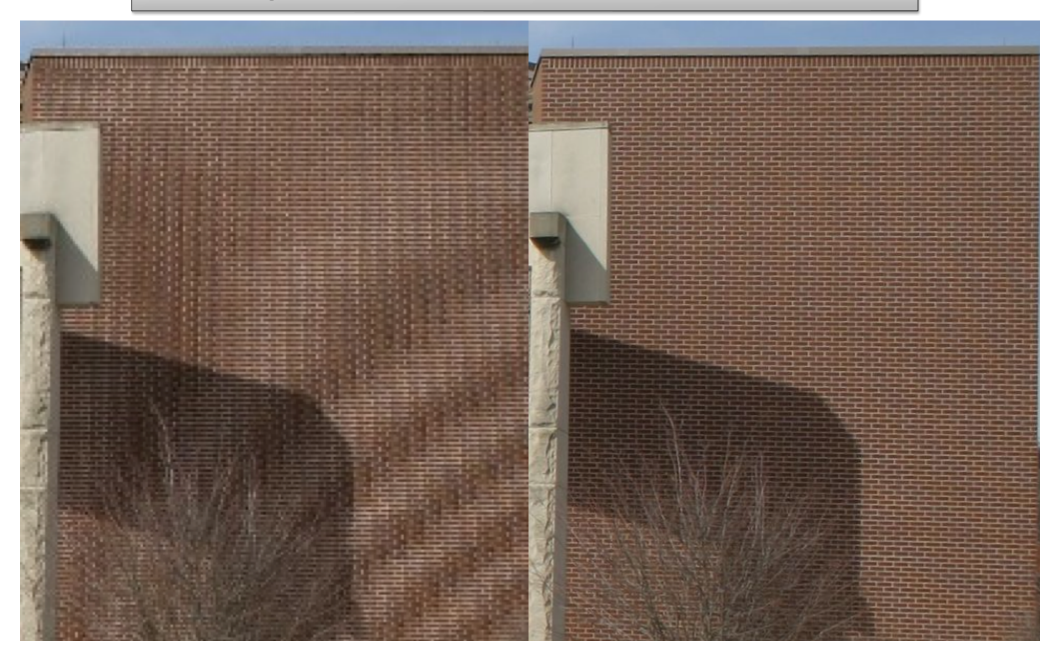
1 Airy unit

Harry Nyquist, 1889 - 1976

- ✓ Swedish American
- ✓ engineer in telecommunications
- ✓ worked at Bell labs
- ✓ 138 US patents



Aliasing: Moiré patterns / loss of information



Aliasing: Moiré patterns / loss of information



General form

Digital sampling frequency > analogue frequency x 2

Spatial representation

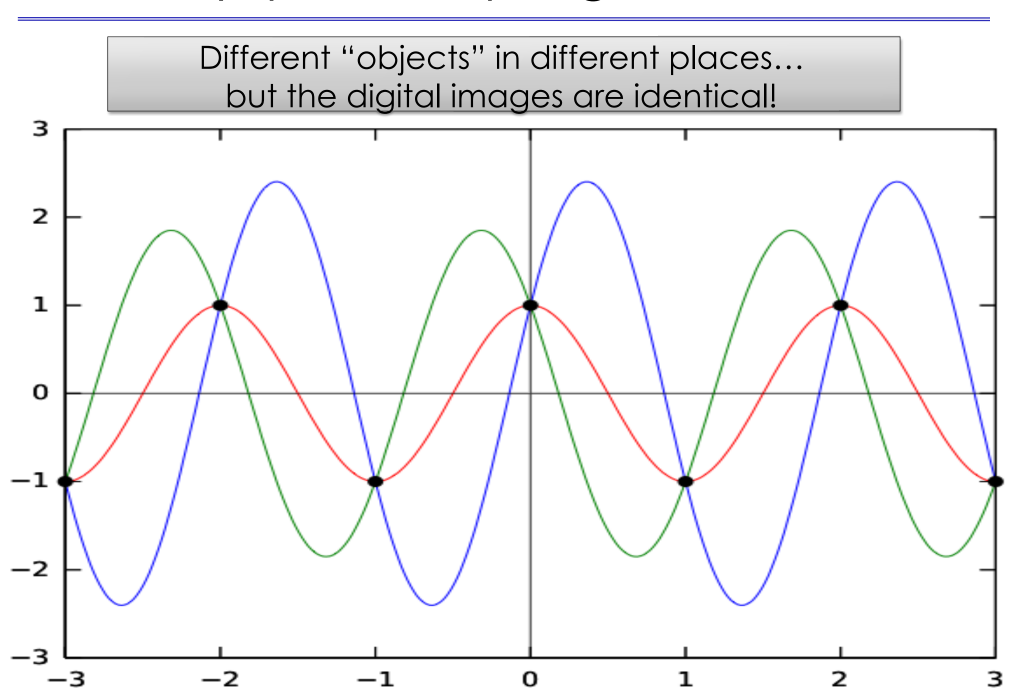
Image pixel size x 2.3 = smallest resolvable distance

Microscopy

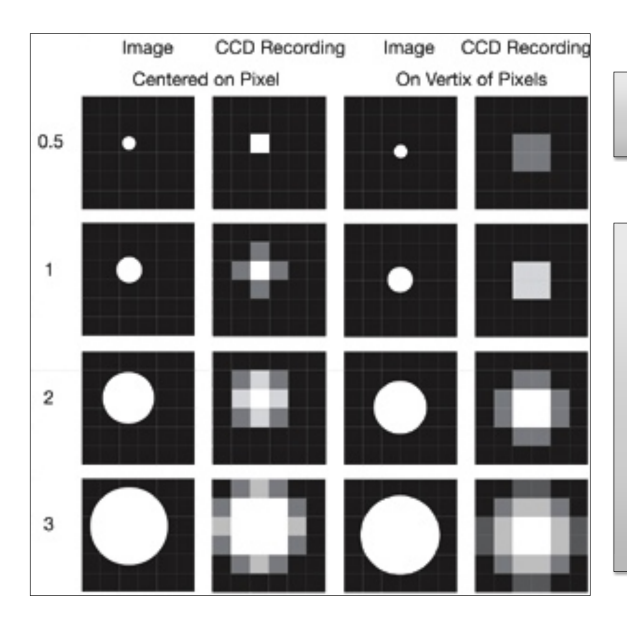
Image pixel size x 2.3 = optical resolution (d)

Aliasing

Moiré interference patterns = loss of information



More aliasing problems...



Pixel size relative to projected image

Image of object varies, depending on where it falls on detector

v for sm

Especially for small objects close to pixel size

Resolution - pixel size calculations:

Resolution, d = lambda / 2 NA

Required Pixel Spacing = d / 3

Optomistic pixel size calculations:

550 nm light; d=lambda/2NA; pix=d/3:

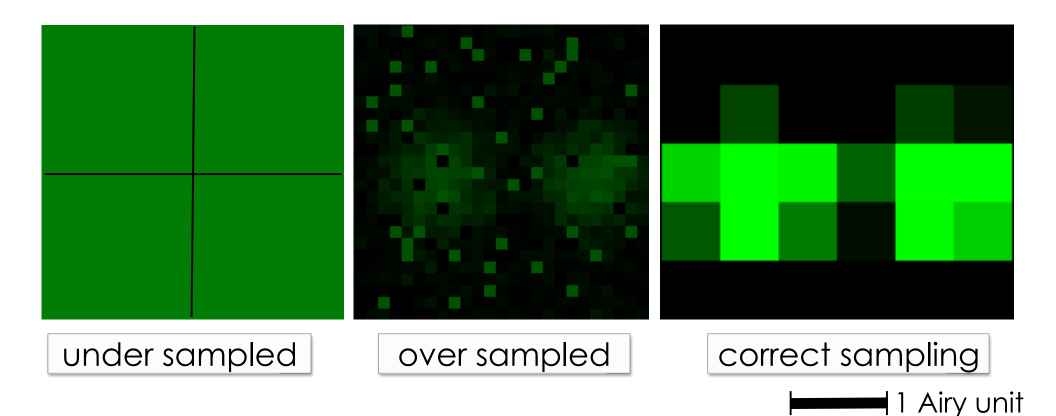
Objective (N.A.)	Optical Resolution limit (nm)	Projected size on CCD (um)	Required CCD pixel spacing (um)	
4x (0.2)	1400	5.5	2	
10x (0.4)	690	7	2	
40x (0.75)	366	14.5	5	
40x (1.3)	210	8.5	3	
63x (1.4)	200	12.5	4	
100x (1.4)	200	20	6.5	

Think about your digital spatial resolution carefully!

Pixel Size / Resolution

Remember !!!

Nyquist told us how to do digital sampling:
~1/3 x smallest feature.



Pixel size / Spatial Calibration

Pixel size is determined by the microscope system!

- ✓ CCD photodiode "pixel" size Magnification X
- ✓ Point scanner settings zoom and image size
- ✓ Field of View Size No. of Samples or "pixels"

It might be changed / lost during processing

It is stored in the "Meta Data"

So .. a dataset for image processing

=

Image data + Meta Data!

Practical Session 1a



Getting to know "Fiji" better –
Fiji is just ImageJ
http://pacific.mpi-cbg.de

File - Open Samples - Embryos or Bridge

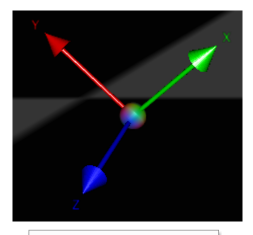
Spatial Scaling:

Can you measure the length and area of objects?

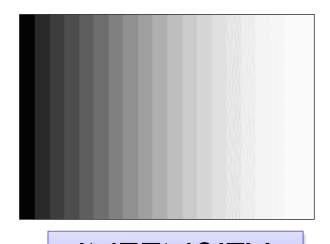
- → See Fiji Tutorial SpatialCalibration (search Wiki)
- ✓ Analyze Set Scale, Analyze-Tools-Scale Bar
- ✓ Line and ROI selection ctrl M (cmd M)
- ✓ Rectangle, Oval, Polygon, Freehand, Angle, Point, Wand.
- ✓ Analyze Set Measurements (Results Edit summarize)

What can you digitise?

Dimensions!



SPACE

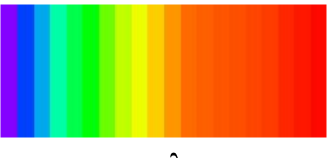


INTENSITY



TIME

Wavelength



Colour

Remember: Bit Depth

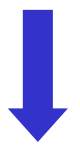
Measured intensity by detector



Corresponding level in image

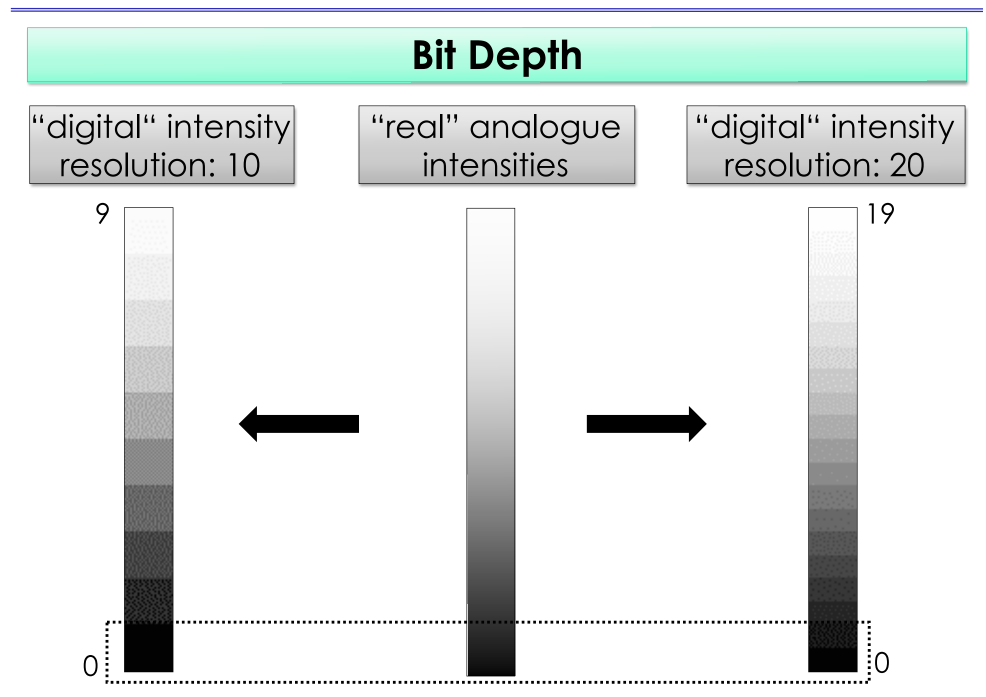
"Bucket" holds 0-9 electrons

5 electrons counted

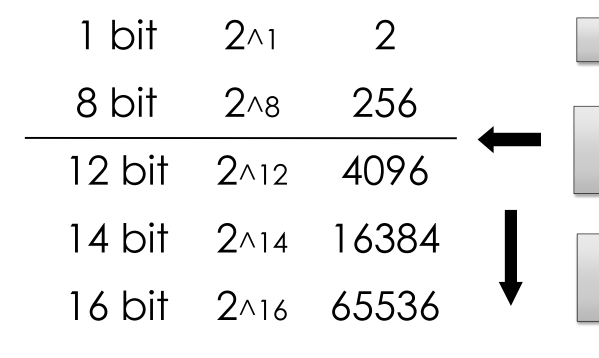


Bit depth: 10 (0 to 9) levels

Level 5 selected for RAW data "image"



Bit Depth

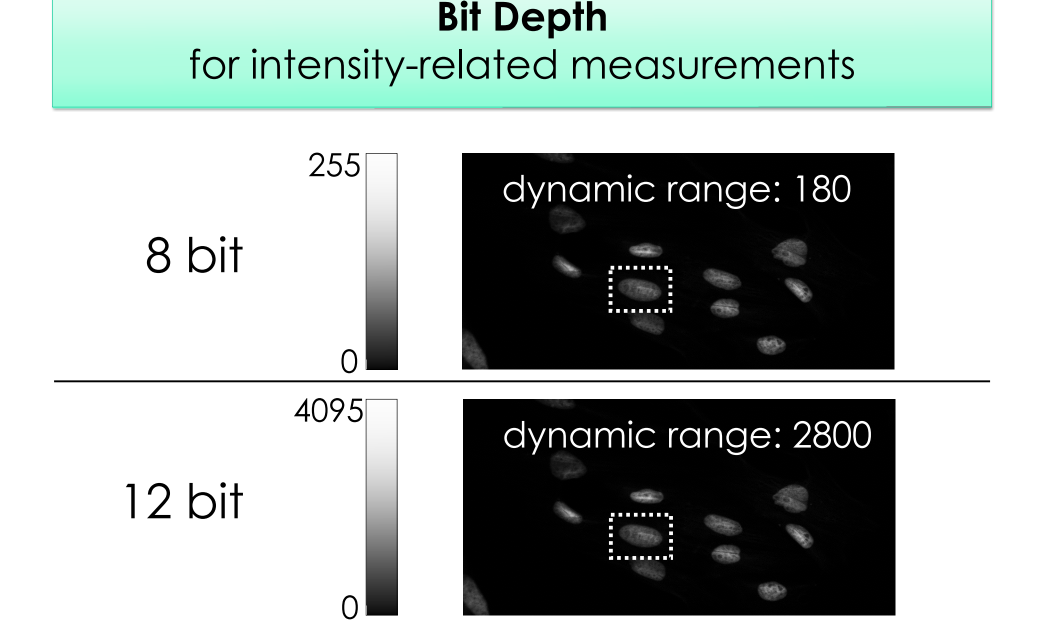


segmentation

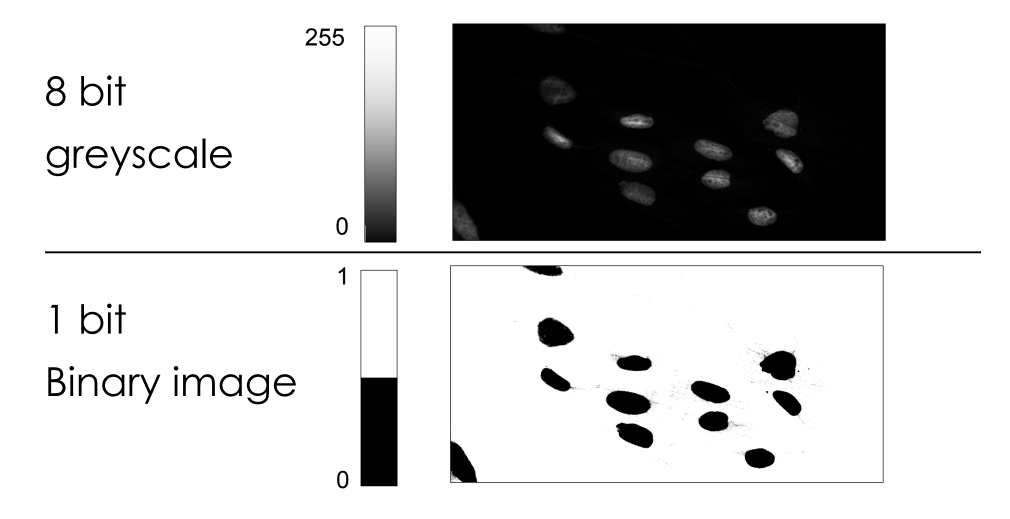
~ limit of human eye, displays...

Intensity-related measurements

• • •







Remember: Intensity / Exposure / Saturation

Do NOT over expose / saturate your image!!!

Why not? → Lost Information!

Use "Look Up Tables (LUT) / palettes to check the saturation

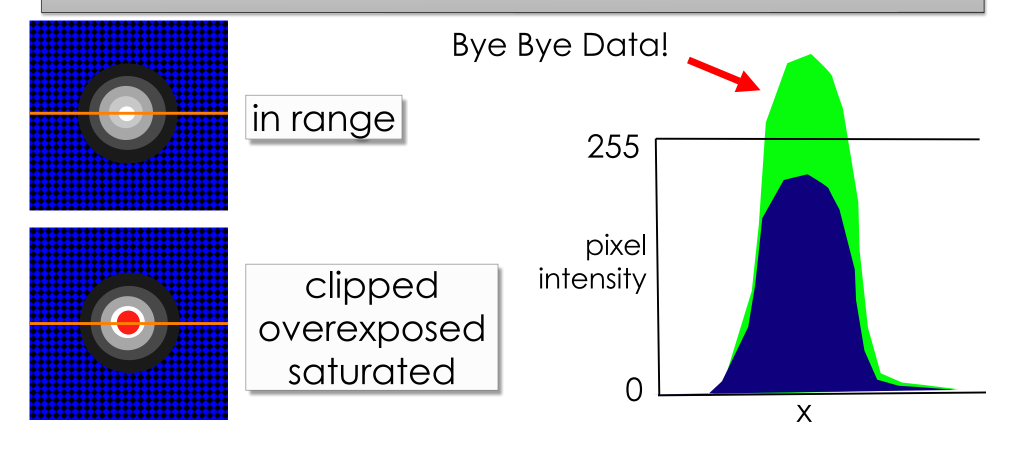
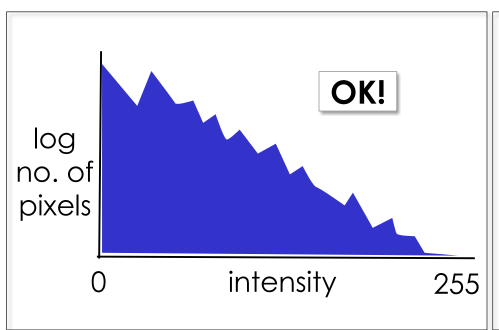
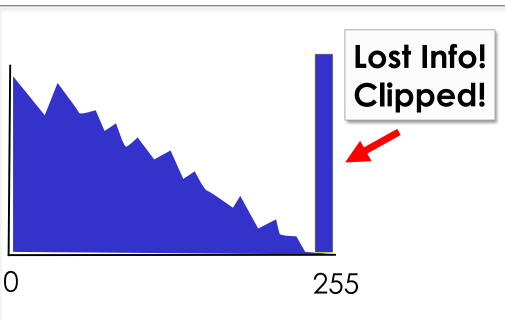
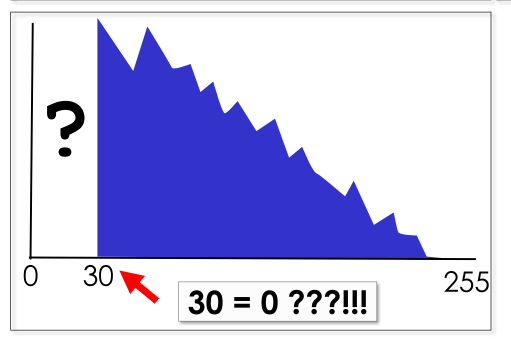


Image Intensity Histograms - Use them!

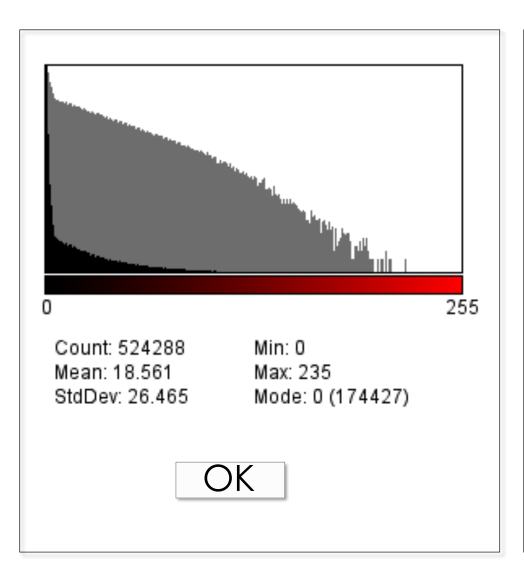


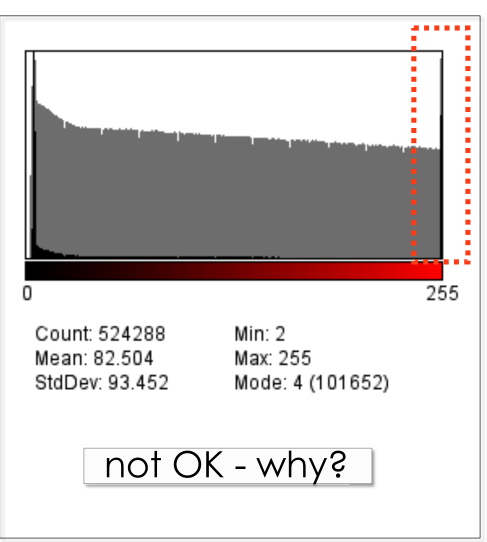




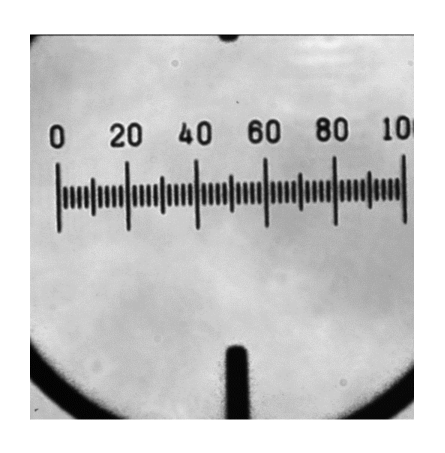
In Histograms:
easily see problems
for image
quantification!

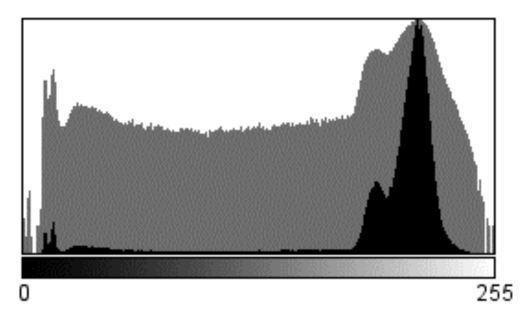
Fluorescence Microscopy





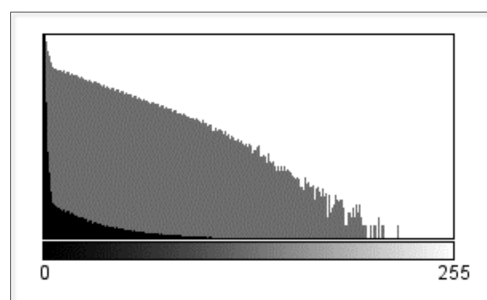
Brightfield Microscopy





Count: 262144 Min: 0 Mean: 191.793 Max: 255

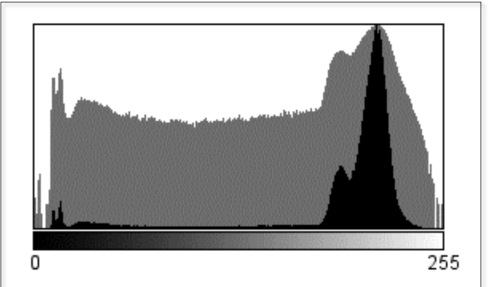
StdDev: 50.337 Mode: 214 (10291)



Count: 524288 Min: 0 Mean: 18.561 Max: 235

StdDev: 26.465 Mode: 0 (174427)

fluorescence



Count: 262144 Min: 0 Mean: 191.793 Max: 255

StdDev: 50.337 Mode: 214 (10291)

brightfield

Practical Session 1b



Getting to know "Fiji" better –
Fiji is just ImageJ
http://pacific.mpi-cbg.de

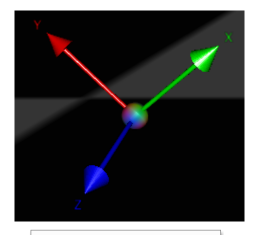
File - Open Samples - Neuron

Intensity clipping/saturation and offset:

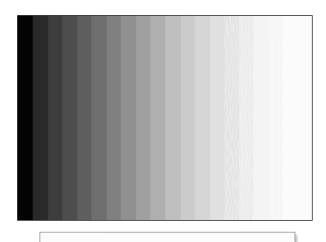
- ✓ <u>Bit Depth</u> change from 16 to 8. What happens to the numbers?
- ✓ <u>Brightness/Contrast</u>: Image-Adjust-Brightness/Contrast. Realize: you can loose data using "Apply"!
- ✓ <u>Intensity Histograms</u>: log scale for fluorescence

What can you digitise?

Dimensions!



SPACE

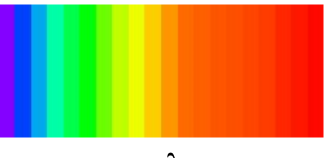


INTENSITY



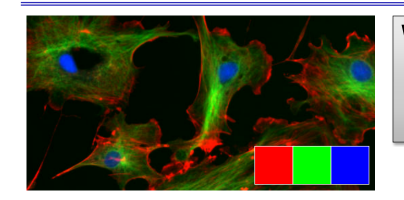
TIME

Wavelength

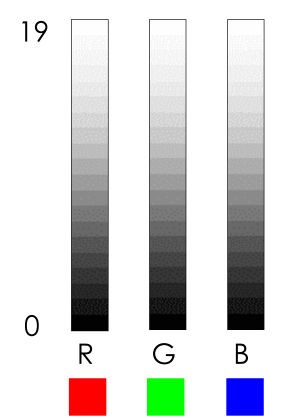


Colour

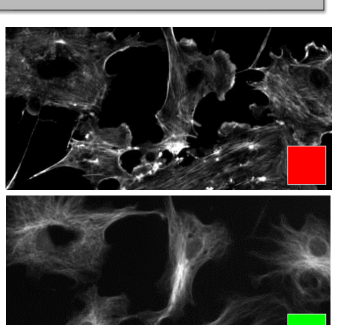
RGB Color Space

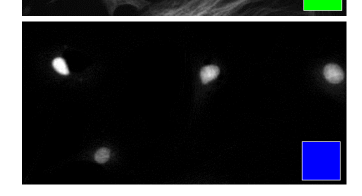


Why RGB? ... because we have red, green and blue sensitive photo receptors in our eyes!

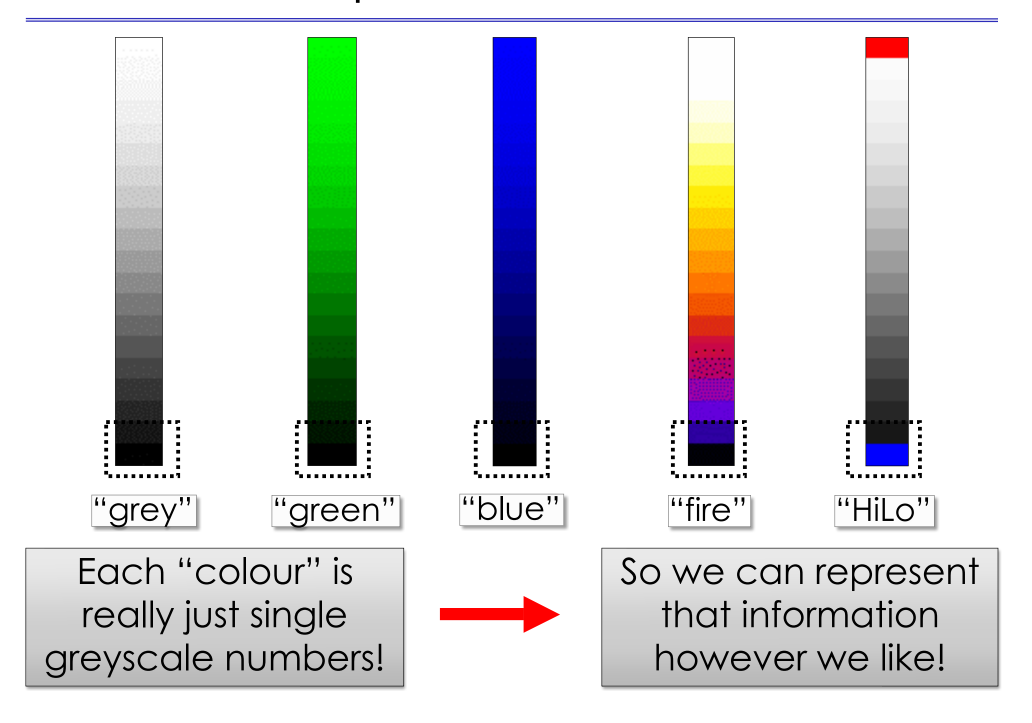


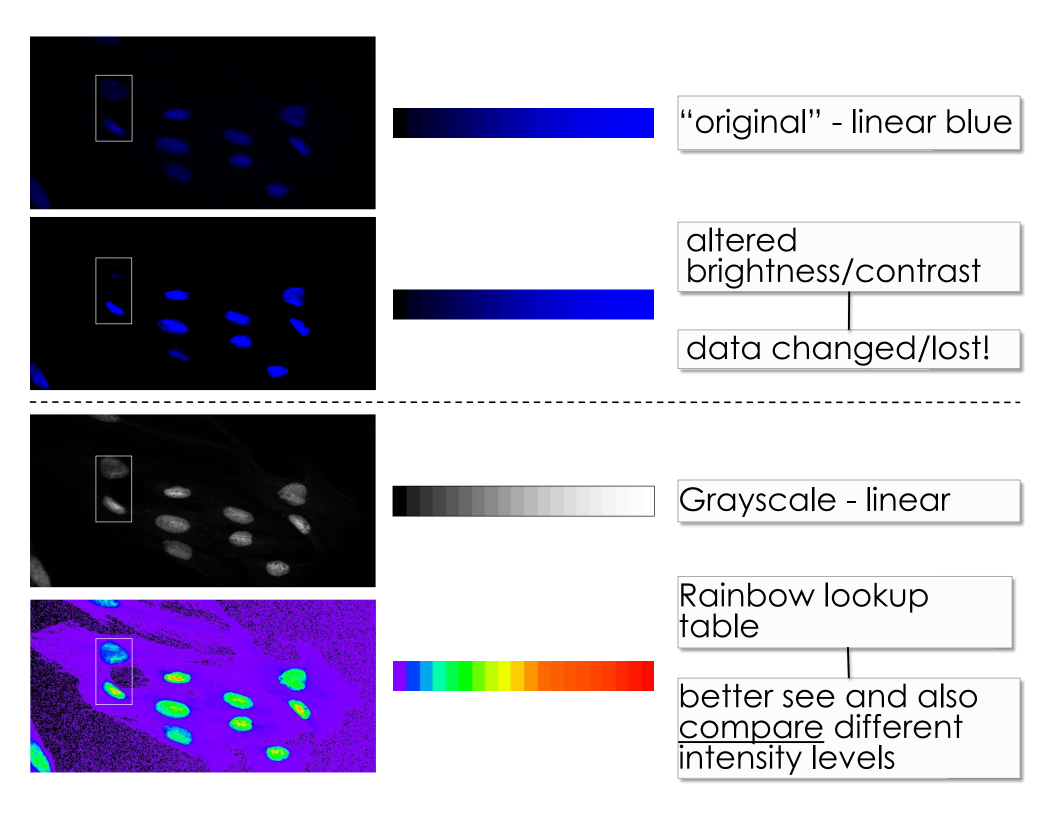
Each "colour"
is really just
single
greyscale
numbers!

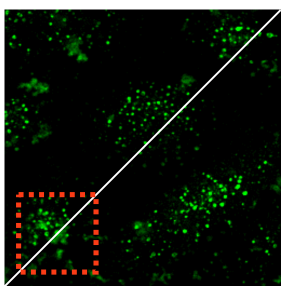


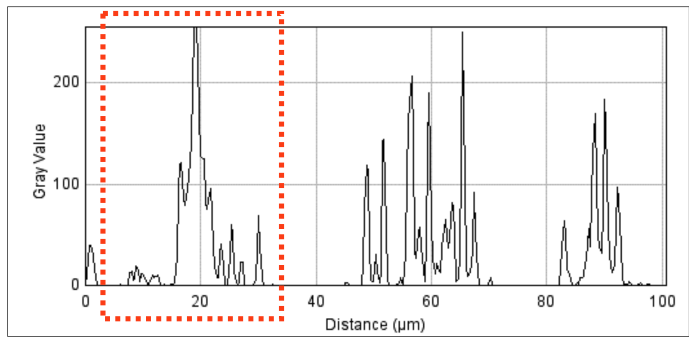


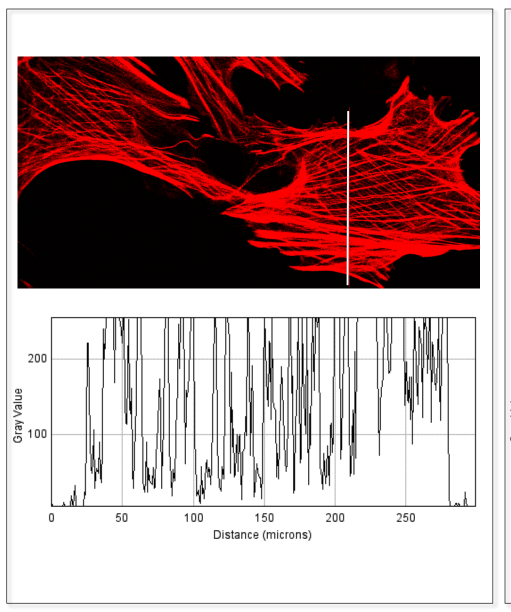
Lookup Tables / Palettes

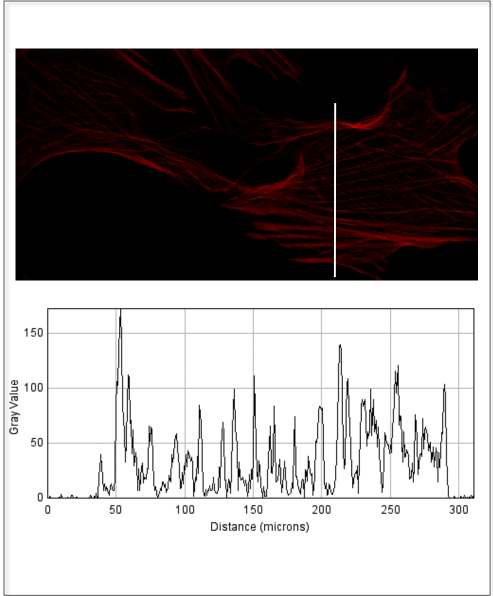




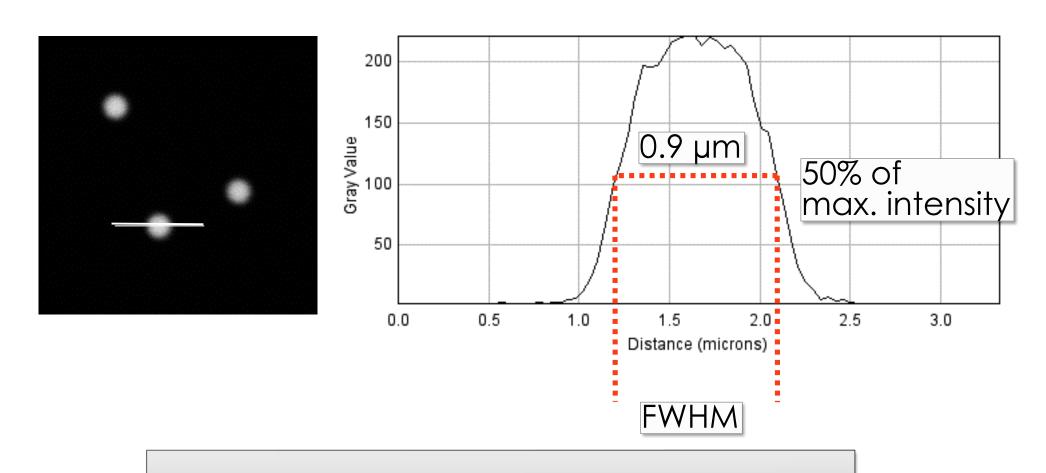




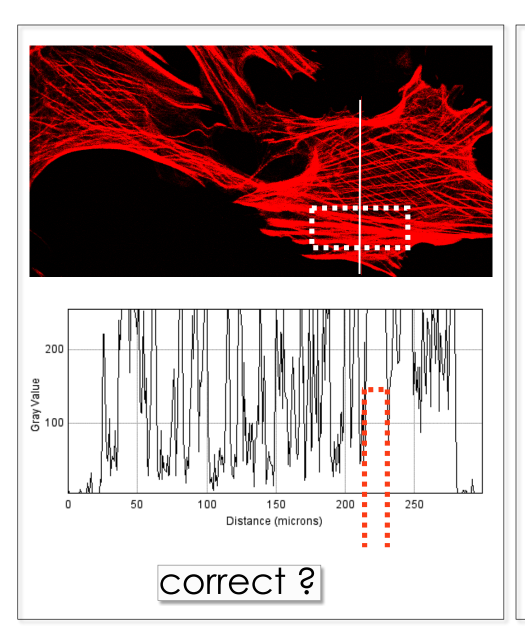


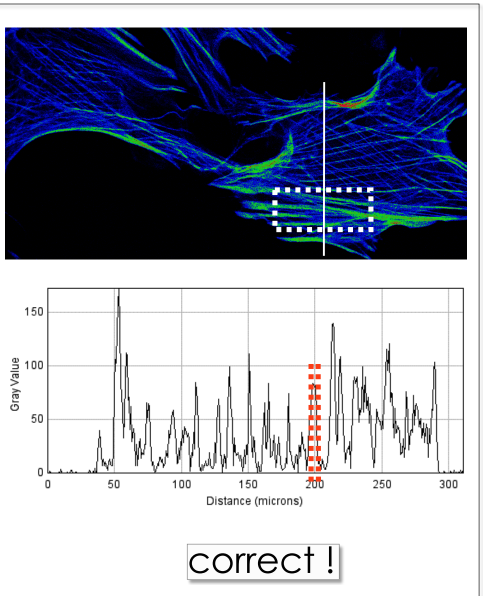


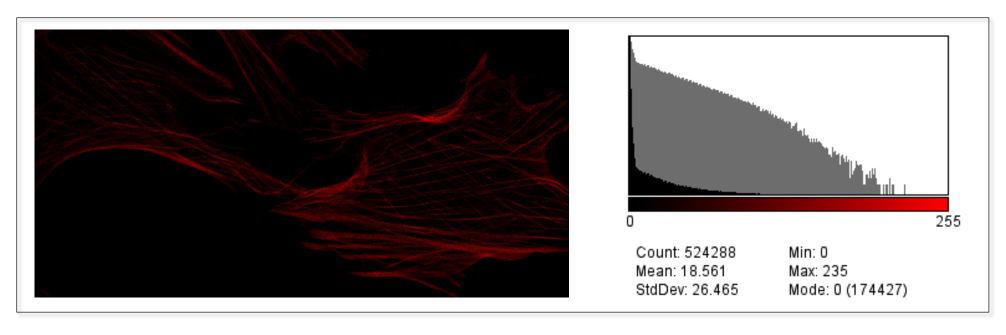
for measurements

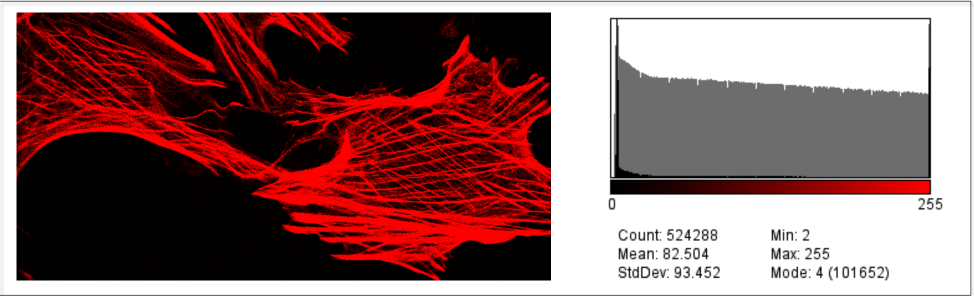


FWHM= "Full Width at Half Maximum"



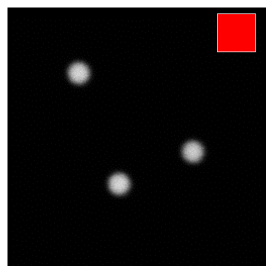


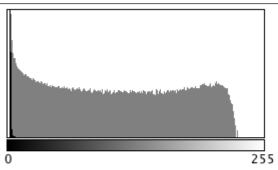




2D Histogram =

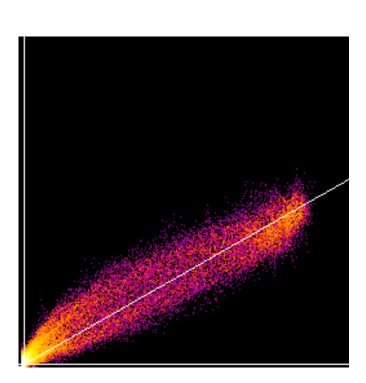
Scatterplot or cytofluorogram

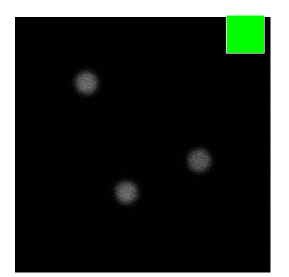


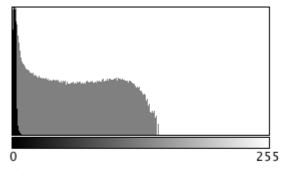


Count: 589824 Min: 2 Mean: 5.782715 Max: 229

StdDev: 22.748384 Mode: 2 (480434)



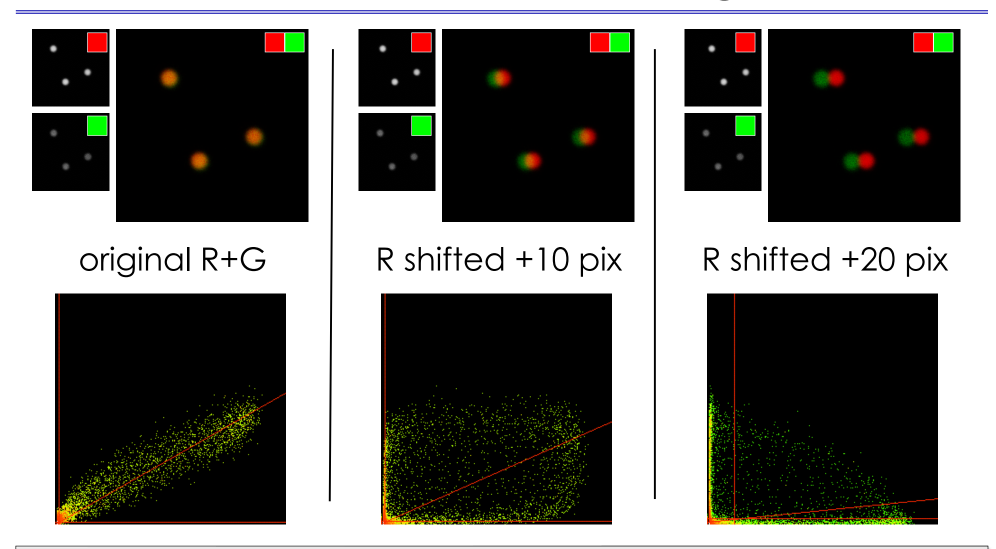




Count: 589824 Min: 0 Mean: 3.928024 Max: 153

StdDev: 13.113667 Mode: 1 (140607)

Scatterplot / 2D Histogram



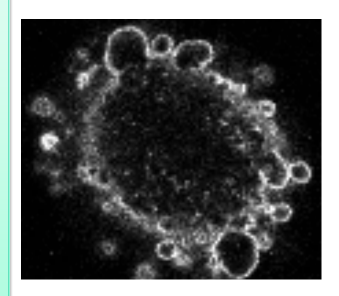
Find a way to <u>visualise</u> what you <u>actually want to see</u>:

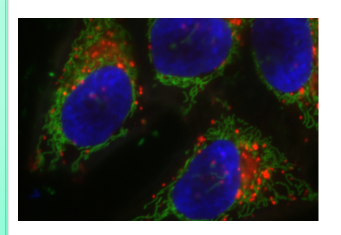
Here, we don't care <u>WHERE</u> the beads are;

We care if they are in the <u>same place or not!</u>

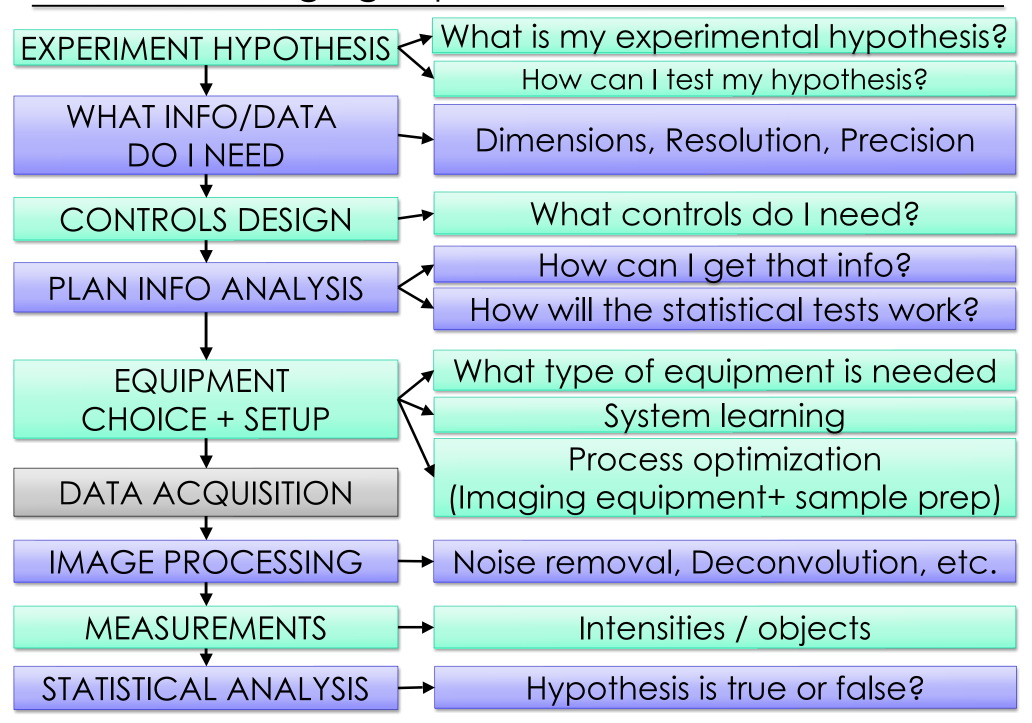
Imaging Experiment Planning:

- ✓ What BIOLOGY am I trying to measure?
 - Hypothesis?!!?
- ✓ Do I need 3D, 4D, xD information
 - Resolution?
 - Sampling: Space, Time, Intensity
- ✓ Choose appropriate microscope
 - Don't always use Confocal LSM
- ✓ Optimise microscope system
 - get best data from your sample
- ✓ Do the right controls!!!
- ✓ Measure Something
 - Statistics to test hypothesis
 - How many data points/images/cells?





Imaging Experiment Work Flow



Practical Session 1c



Getting to know "Fiji" better – Fiji is just ImageJ http://pacific.mpi-cbg.de

File - Open Samples - Neuron

RGB colour space:

- ✓ <u>Colour channels</u> Image-Colour-Channels Tool, Split channels etc.
- ✓ LookUp Tables/Paletts: Image Lookup tables, or LUT toolbar icon
- ✓ <u>Line Profile</u>: Analyze Plot Profile
- ✓ <u>Histogramm</u>: Analyze-Histogram or Plugins-Analyze-2D Histogram
- ✓ Intensity Scale: Analyze Tools Calibration bar

Basics of Quantitative Image Analysis

What you need to know about Image Processing... but never thought to ask ... continued

Session 2

- ✓ Filtering Images in the spatial, frequency and time domain
- ✓ Segmentation finding and measuring objects in images

Session 3

- ✓ Detect Info Loss, Colocalization Analysis and more
- ✓ Whatever you find interesting



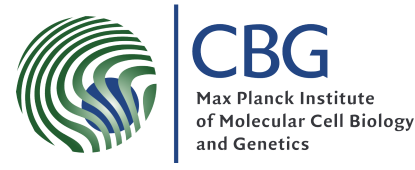
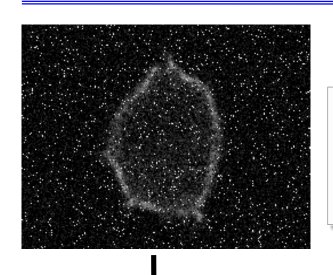




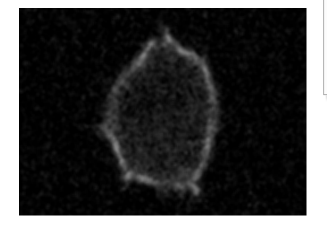


Image processing in the spatial domain



A) Introduction

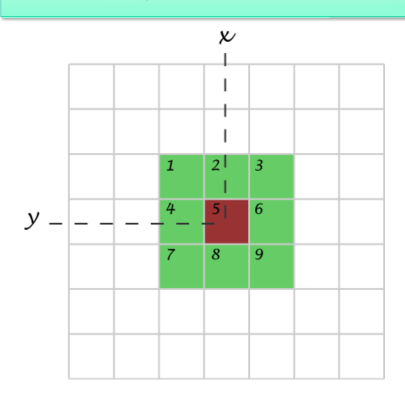
- Neighbourhood
- Operation on neighbourhood



- B) Spatial filters
 - Mean and Median filter
 - Edge detection

A. Introduction

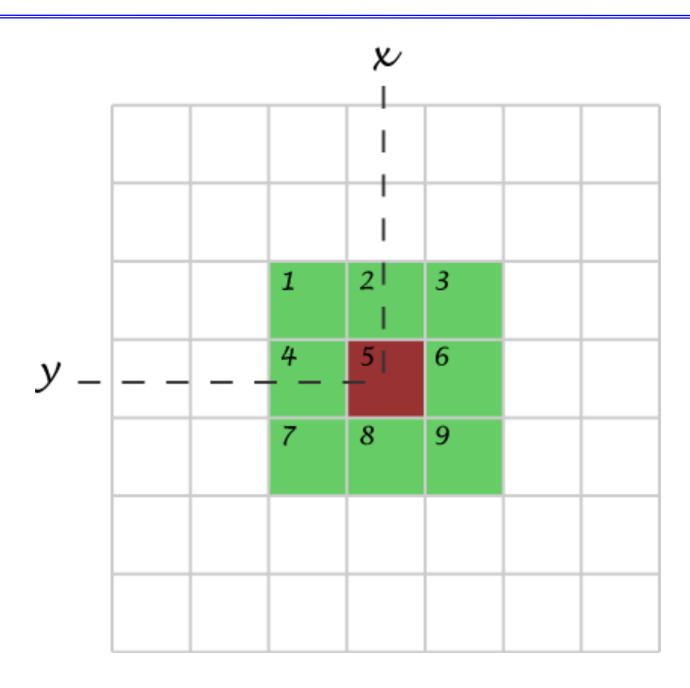
"Transformation or set of transformations where a new image is obtained by neighbourhood operations."

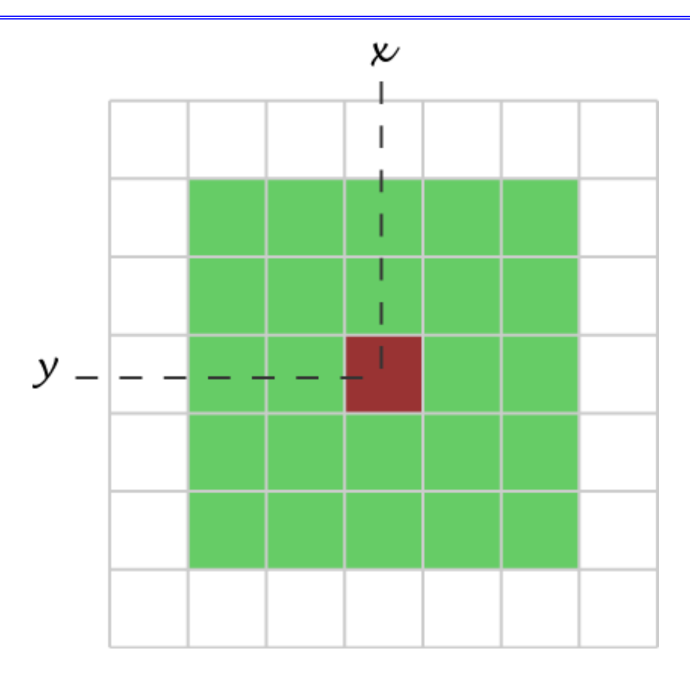


The Intensity of a pixel in the new image depends on the intensity values of "neighbour pixels"

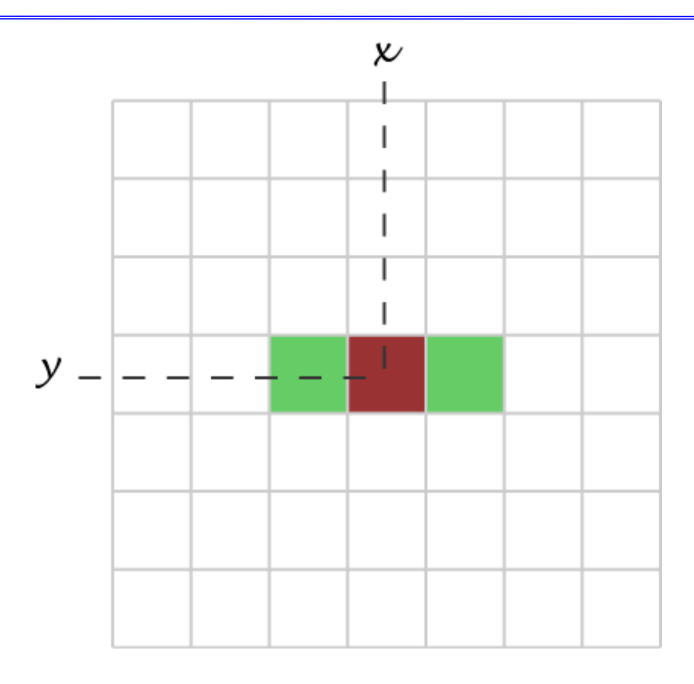
Neighbourhood (or kernel): pixels that matter

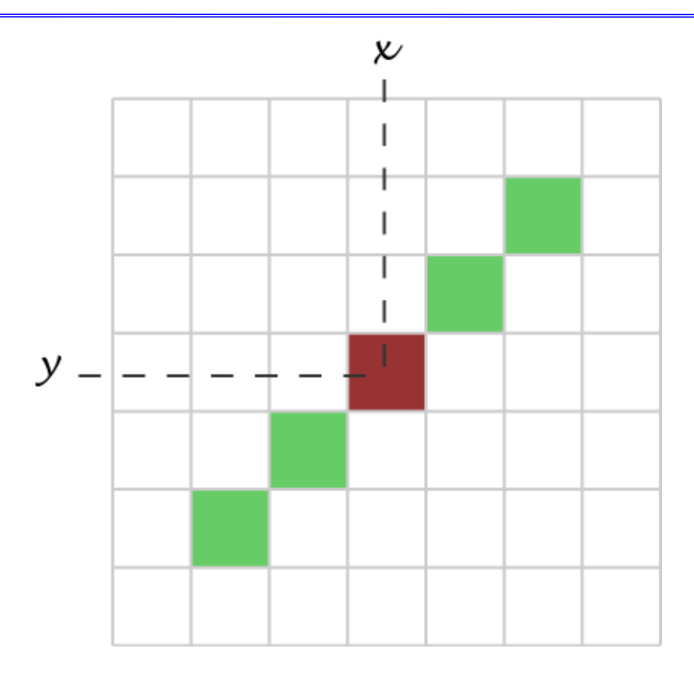
3 x 3



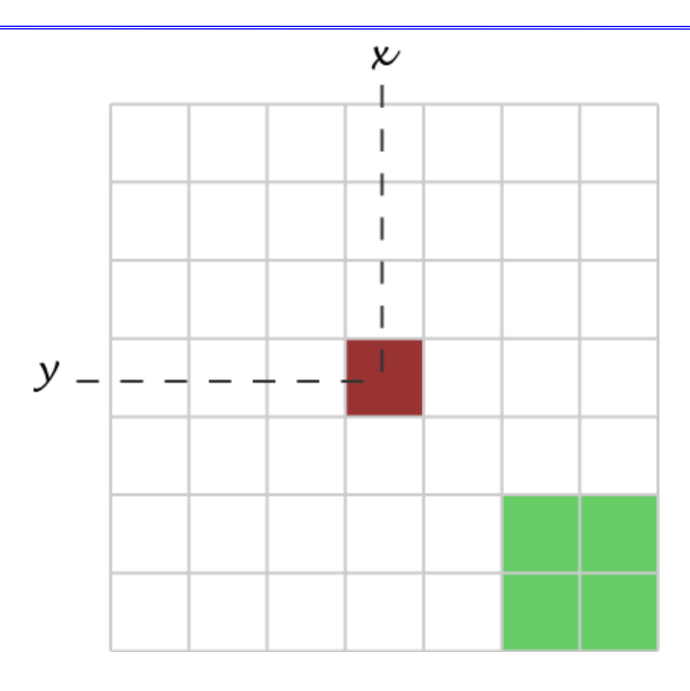


1 x 3

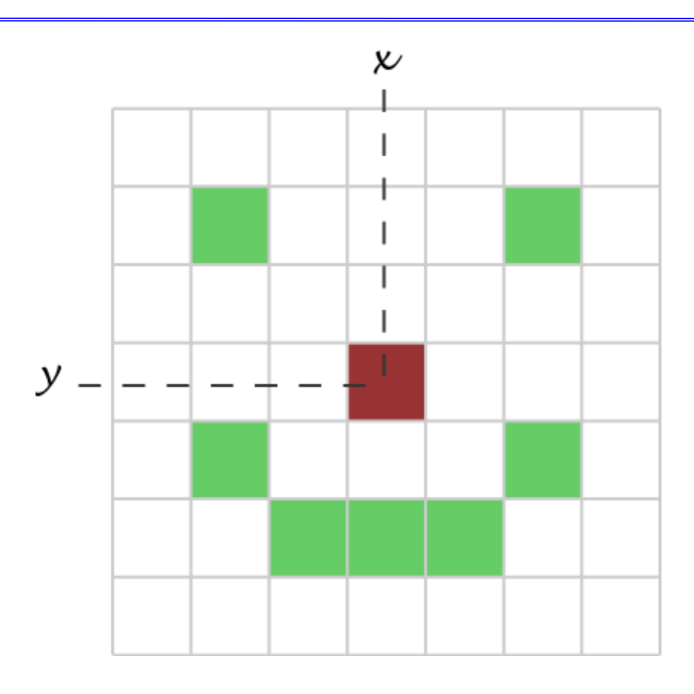




2 x 2; shift

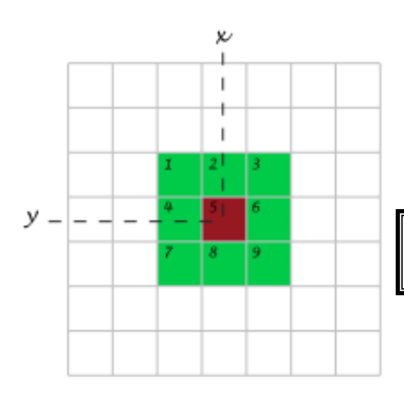


Misc.



B: Filtering - the mean filter

Simplest filter: The value of a pixel is replaced by the intensity mean of the neighbourhood pixels.

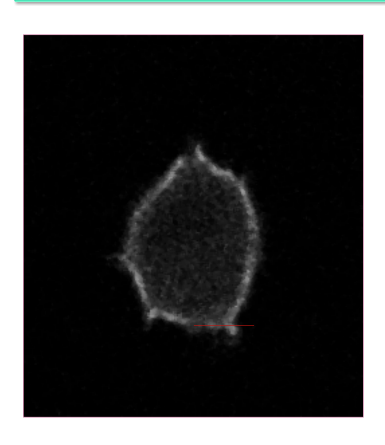


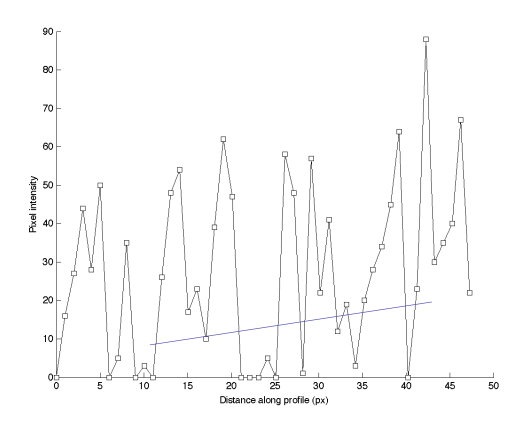
3x3 example:

$$\alpha_1^* = \frac{1}{9}(a_1 + a_2 + a_3 + a_4 + a_5 + a_6 + a_7 + a_8 + a_9)$$

The mean filter

Noise removal - typically Gaussian or Poisson noise.





Appears for weak labeling, short exposure time, confocal = few photons detected

The mean filter

The mean filter is a linear filter!

$\alpha_{1,1}$	$\alpha_{\text{1,2}}$	α _{1,3}
$\alpha_{2,1}$	α _{2,2}	$\alpha_{2,3}$
$\alpha_{3,1}$	$\alpha_{3,2}$	$\alpha_{3,3}$

"The new pixel value depends on a linear combination of neighbourhood pixel values"

(The order of several linear filters in sequence does not matter)

The mean filter

Main property: low-pass filter (smooths small objects)

- kernel size influence
- number of successive applications
- + simplest filter fast
- + it's a linear filter
- + averages noise, does not eliminate it
- + works againstGaussian and Poissonnoise

- blurs images small details are lost (low pass filter)
- smoothes edges dramatically
- fails for salt & pepper noise

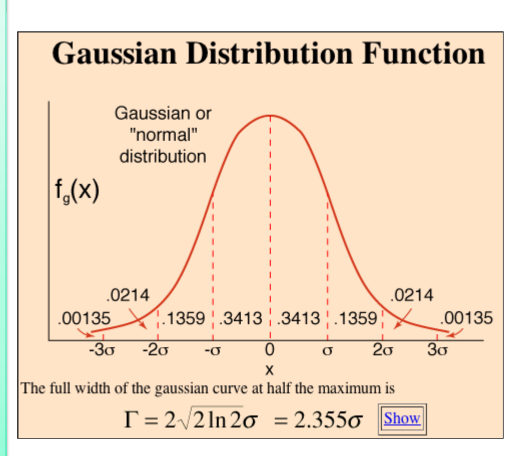
Linear filtering - Properties

- ✓ Applying a linear filter to an image is the same as: applying it to all <u>parts</u>, then summing the results.
- ✓ When applying a succession of linear filters:
 the <u>order</u> filters are applied in does not matter.
- ✓ Mathematical framework underlying it: <u>Convolution</u>.
- ✓ We can also reverse the process: <u>Deconvolution</u>

The Gaussian filter

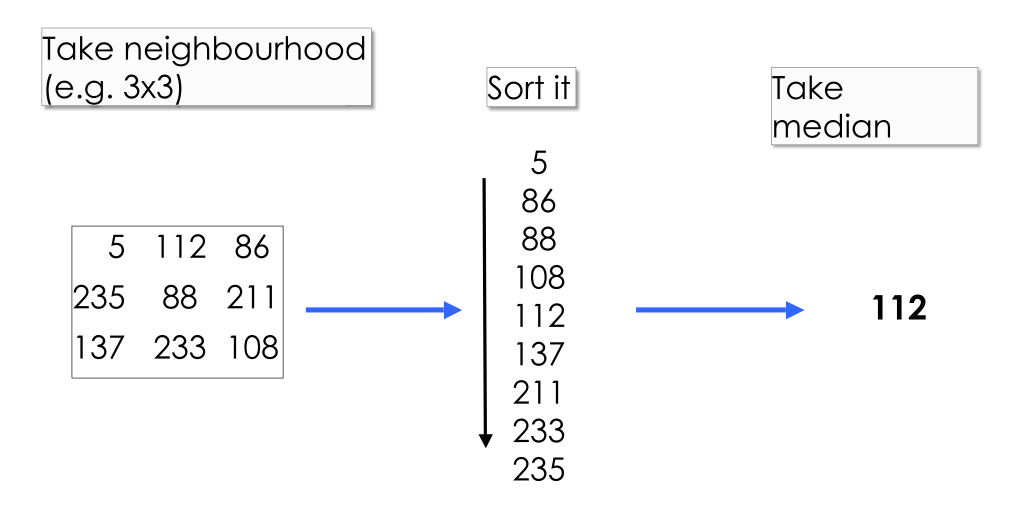
Gaussian Curve - Bell Shaped function

- ✓ smooths Poisson noise
- ✓ linear Filter
- ✓ makes more mathematical sense than mean filter?
- ✓ ...properly spatially sampled image, looks like PSF
- ✓ can vary the sigma value:
 number of pixels
- ✓ varying degree of blur.



The median filter

The value of a pixel is replaced by the median of the pixel intensity in neighbour pixels



The median filter

noise elimination

Original:

5 9 6 6 9 5 9 9 5 9 7 8 7 9 8 9 8 6 7 9 9

9 9 7 200 9 6 9

6 5 8 6 9 6 7 9 7 9 9 8 6 7

7 9 5 6 7 6 6

outlier

Median filtered:

0	5	6	6	6	7	0
5	8	7	7	7	9	7
8	9	8	8	7	9	7
6	8	8	8		9	6
6	8	8	9	8	7	6
6	7	7	8	6	7	6
0	7	6	6	6	6	0

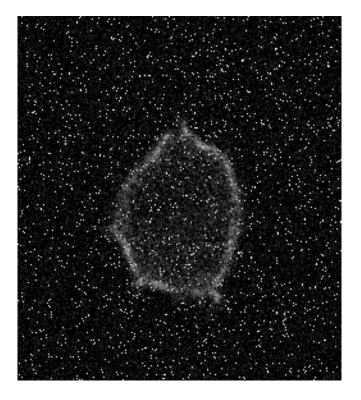
The outlier value has been completely removed from the dataset

The median filter - what is it good for?

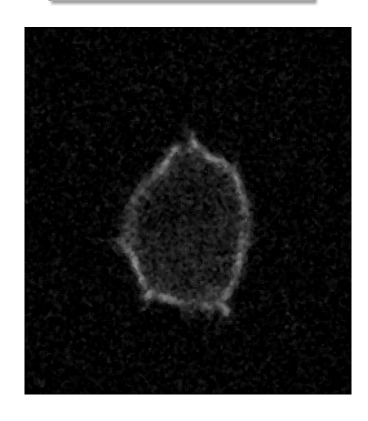
"Salt & pepper" noise * removal

* Typically appears for very weak labeling, high detector gain etc.

Original:



Median filtered:



The median filter

- + Typically good for "Salt & pepper" noise removal
- + Eliminates noise
- + Edge-preserving

- Slower than mean (not such a problem anymore... computers are fast)
- NOT linear



Practical Session 2a

Simple Image Filtering

- (1) File Open Samples bat cochlea volume
- (2) File Import URL...

http://pacific.mpi-cbg.de/samples/colocsample1bRGB_BG.tif

(1) Convolve a simple binary image

- ✓ Process Filters Convolve (play with different kernels)
- ✓ Process Filters Gaussian Blur (change sigma, in px)

(2) Noisy sample image

- ✓ Mean and Median Filter (change pixel number, kernel size)
- ✓ Gaussian Blur ... and Gaussian Blur again... and...

The Fourier transform

The Fourier transform is a way to obtain a new representation of the data (a bit like the 2D histogram from earlier)

It is best suited for data with repetitive patterns, as it highlights those

And ... don't worry about the maths for now...

The Fourier transform

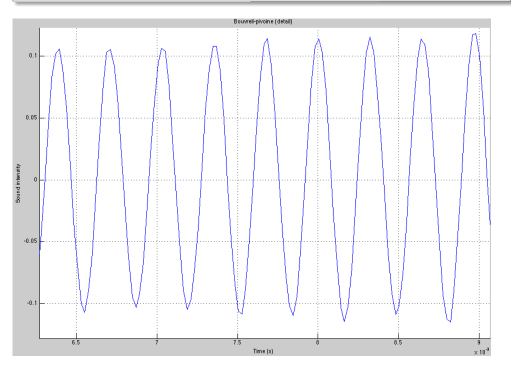
Bird song

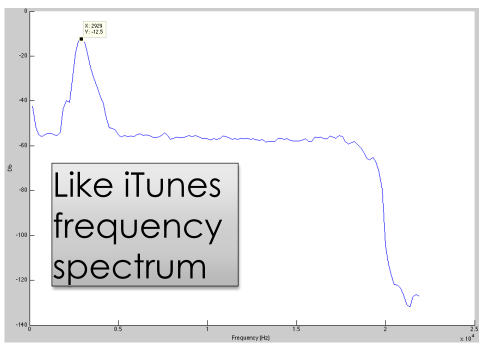
Detail of the signal:

Delay between peaks:~ 0.35 ms

FFT of this looks like:

Peak in FFT: ~ 3 kHz



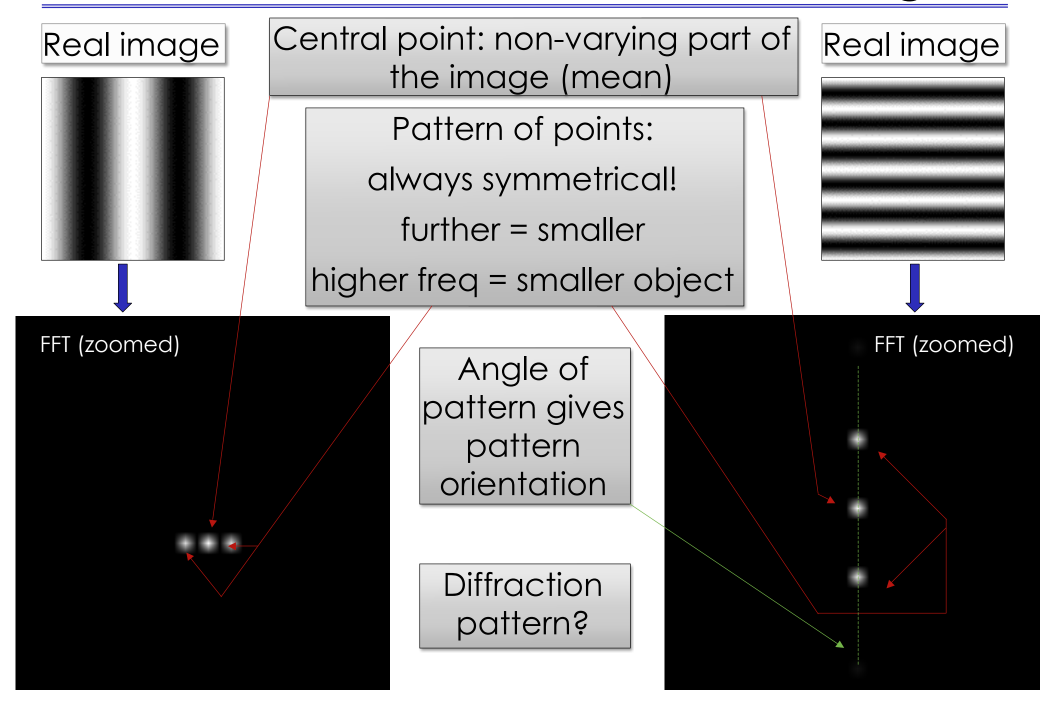


Equivalence: spatial domain vs. Fourier or Freq. domain

1 / 3000 = 0.33 ms

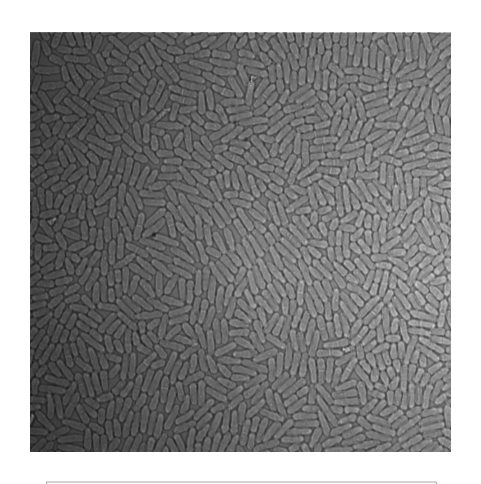
Peak in FFT gives frequency or peroidicity of pattern

The Fourier transform – in 2D images

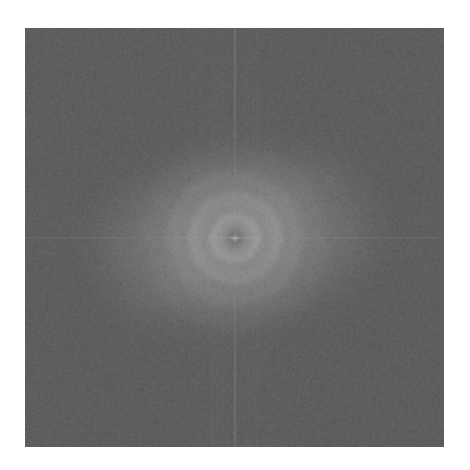


The Fourier transform – in 2D images

Real images... are rarely that clear



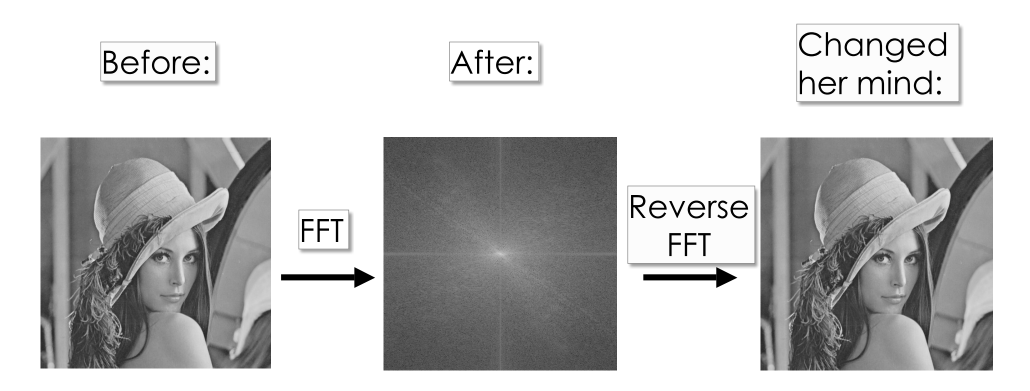




FFT

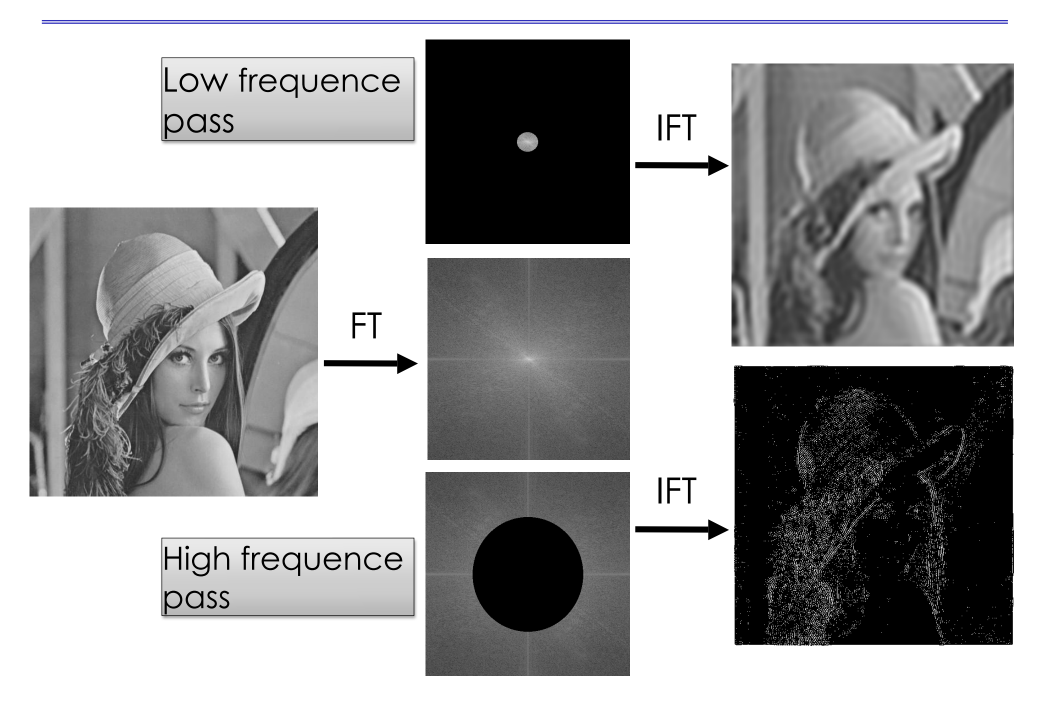
The inverse Fourier transform

Fourier image and real image contain same information \rightarrow so it's possible to reverse the process:

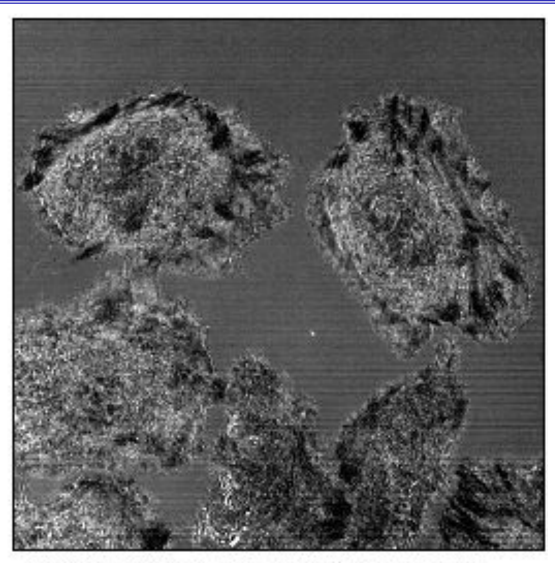


Same thing happens physically in a microscope. FT image is in the Back Focal Plane of Objective!

Can use as a filter for detail:



... a filter for periodic noise:

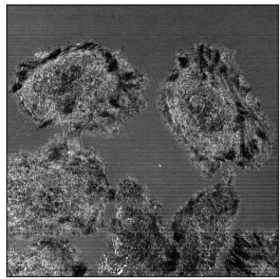


The original image. Reflectance mode of the confocal using the 458 nm line of an Ar laser. Note the horizontal lines.

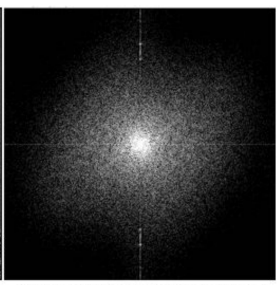
... a filter for periodic noise:

Laser intensity noise from a bad AOTF...

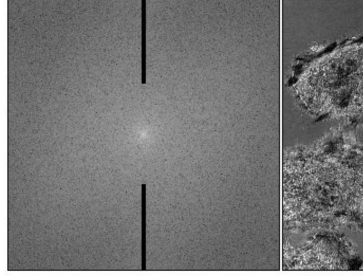
can be removed by frequency filtering in the correct spatial direction.



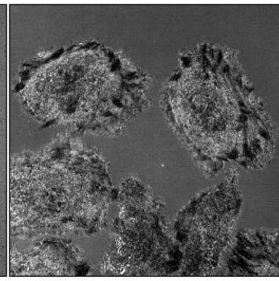
The original image. Reflectance mode of the confocal using the 458 nm line of an Ar laser. Note the horizontal lines.



The power spectrum calculated by ImageJ, contrast enhanced to show the bright spots that represent the X axis fluctuation.



The power spectrum with masks drawn on it.



The inverse transform applying the masks.

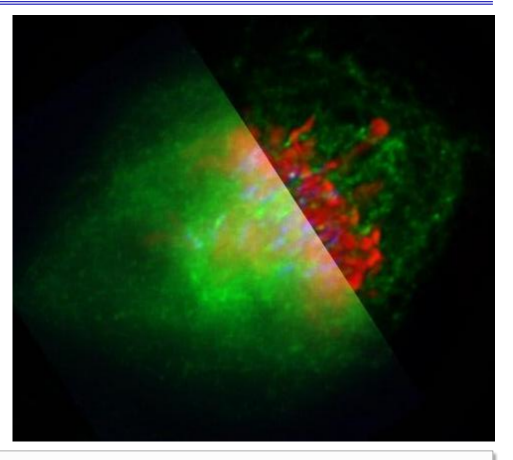
... during "Deconvolution":

Take Image and PSF image

- + Do Fourier transforms
- + Image FT / PSF FT
- + Reverse FT of result

_

Deconvolved image with much improved contrast and less out of focus signal.



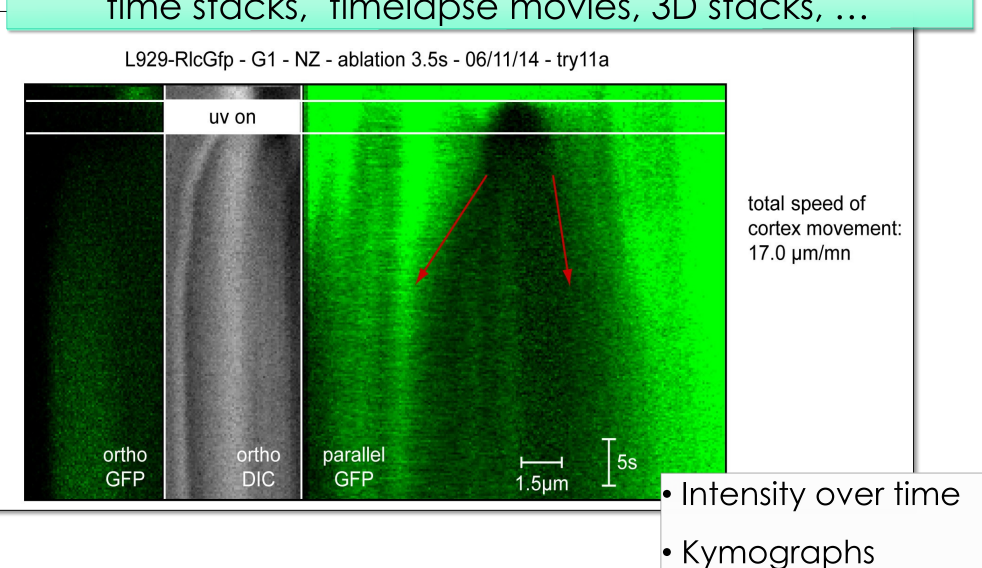
A metaphase human cell stained for DNA (red), centromeres (blue) and the anaphase promoting complex/cyclosome (green). Recorded by Claire Acquaviva, Pines lab

Left part: original data

Right part: deconvolved with Huygens Professional.

Time? Just another dimension

Dealing with multiple images files: time stacks, timelapse movies, 3D stacks, ...



Motion blur

Motion blur = average over time
Does this happen in your sample? Frame Rate?





Practical Session 2b

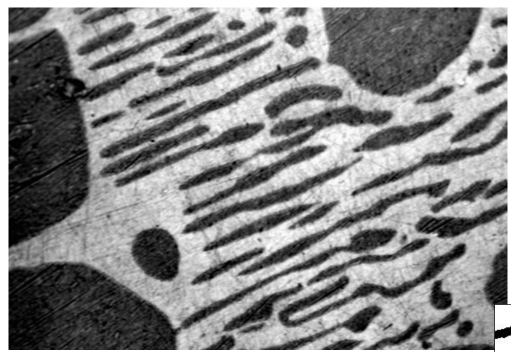


Getting to know "Fiji" better –
Fiji is just ImageJ
http://pacific.mpi-cbg.de

File - Open Samples - Bridge

Fourier Image Filtering

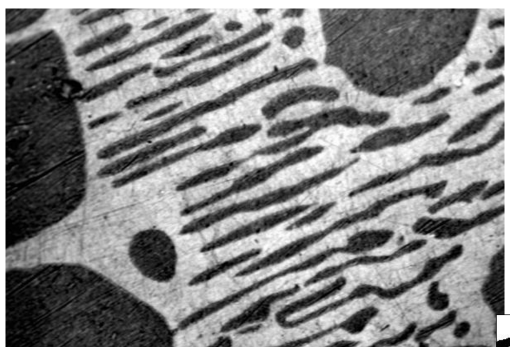
- ✓ FFT, filter out parts, Inverse FFT: Mess up the image. Can you extract high and low frequency information?
- ✓ <u>Use circle selection and Edit Fill</u>: Set foreground colour to black.
- ✓ FFT bandpass filter



"Greyscale" image

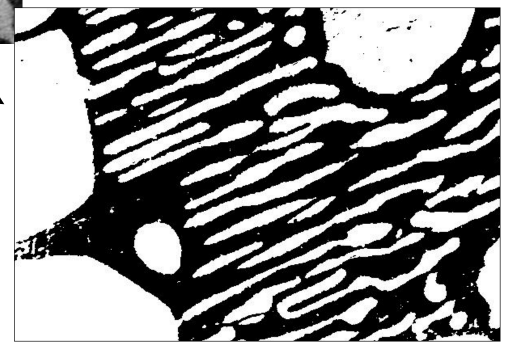
Foreground background





"Scalar Intensity" image

"Binary" image

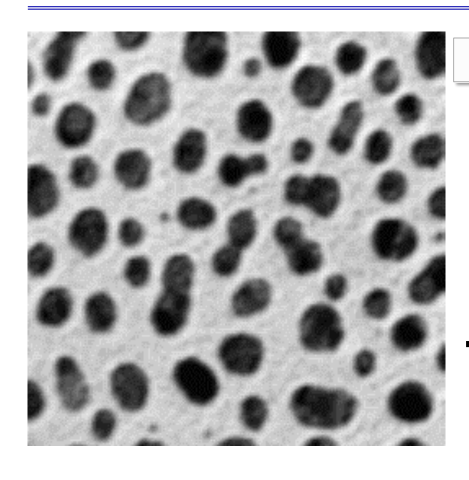


1	65	13	55	2
2	3	34	2	1
4	0	31	1	2
1	33	3	54	3
56	3	2	1	34

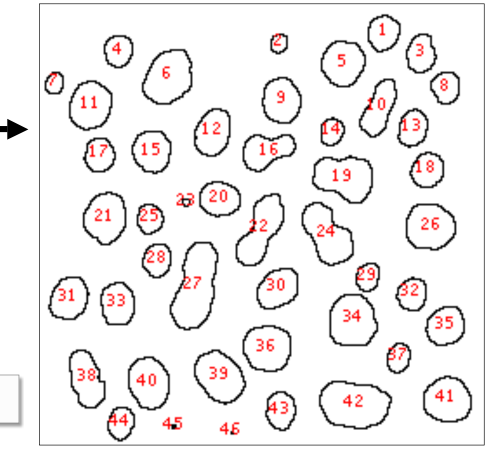
"Scalar Intensity" image

"Binary" image

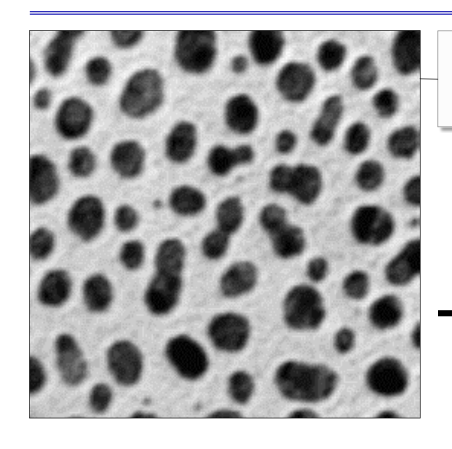
0	1	1	1	0
0	0	1	0	0
0	0	1	0	0
0	1	0	1	0
1	0	0	0	1



"Scalar Intensity" image

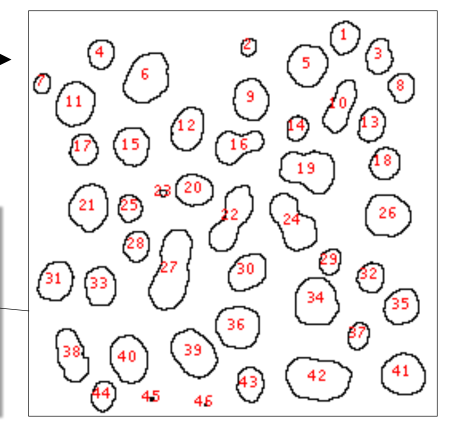


Labeled objects

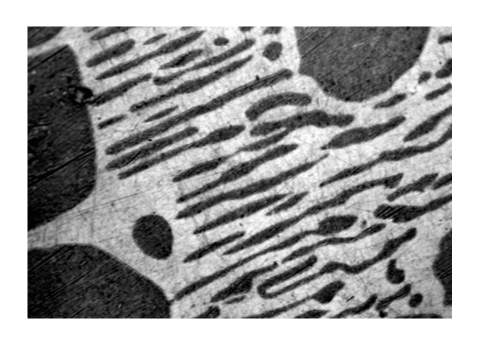


High Information Content 65536 pixels, 0-255 value

Lower Information Content, but easier to interpret biological meaning... 45 "objects" with properties: size, shape, intensity etc.



"Thresholding" (Intensity Histogram Split)



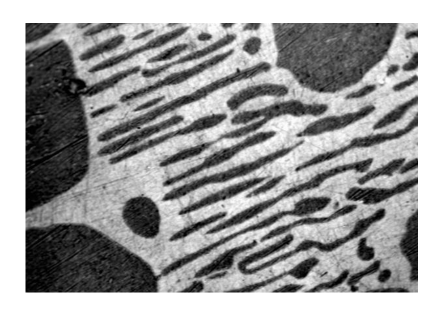
Clear difference between foreground and background?

Image not very noisy?

Choose an intermediate grey value = "threshold"

Determines foreground and background.

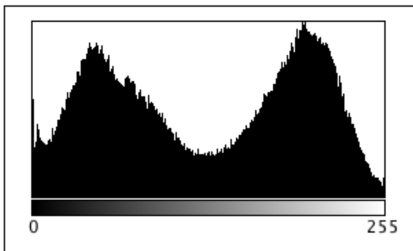
"Thresholding" (Intensity Histogram Split)



How to choose the grey level for thresholding?

Look at pixel intensity histogram of whole image...

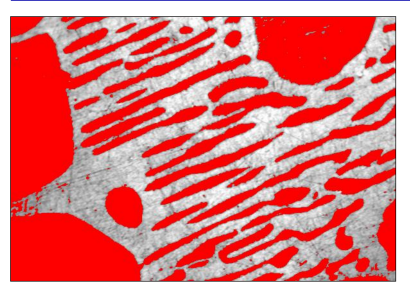
Is there an obvious place?



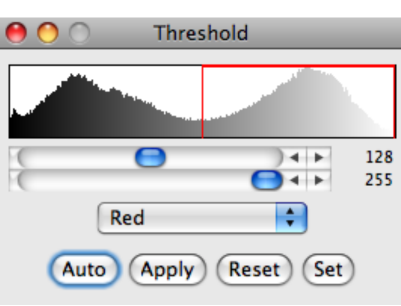
Count: 247200 Min: 0 Mean: 126.159284 Max: 255

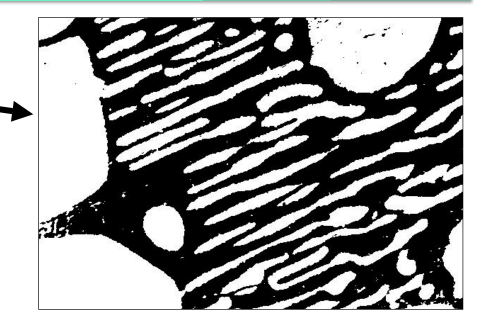
StdDev: 73.220749 Mode: 196 (1820)

"Thresholding" (Intensity Histogram Split)



Histogram is bimodal, so put threshold in the trough between the peaks!

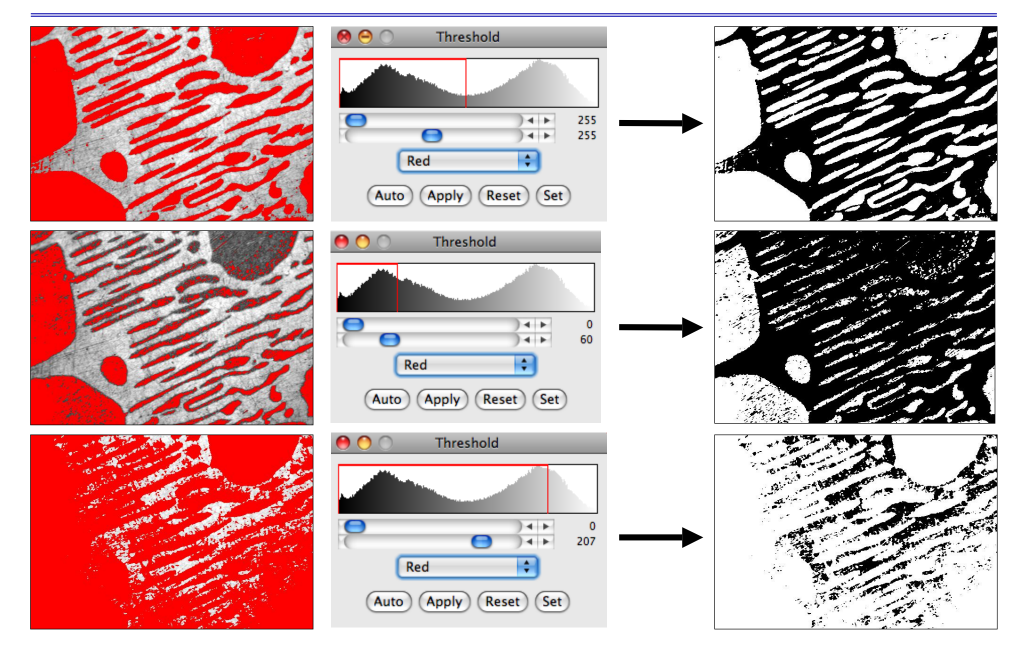




Note, in this case:

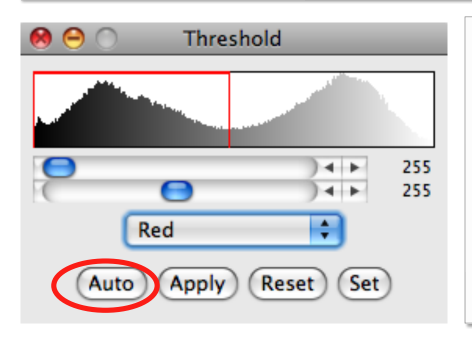
Foreground = "dim" objects Background = "bright" objects

"Dumb Global Threshold" (Subjective - User Biased)



Computed Global Threshold Objective - Reproducible

ImageJ - Image - Adjust - Threshold - Auto (=Make Binary):

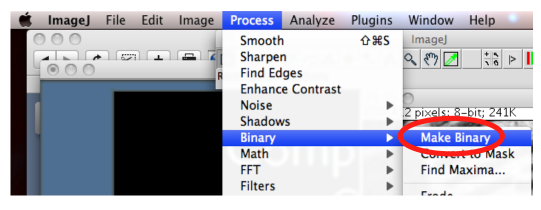


Initial guess of Threshold, T

Compute mean pixel intensity of background and foreground

Tnew = 0.5 x (mean of foregrnd + mean of bkgrnd)

Iterate until Tnew no longer changes.



Note:

Manual threshold set?

Make Binary uses
that dumb threshold!



Practical Session 2c

Simple Image Segmentation

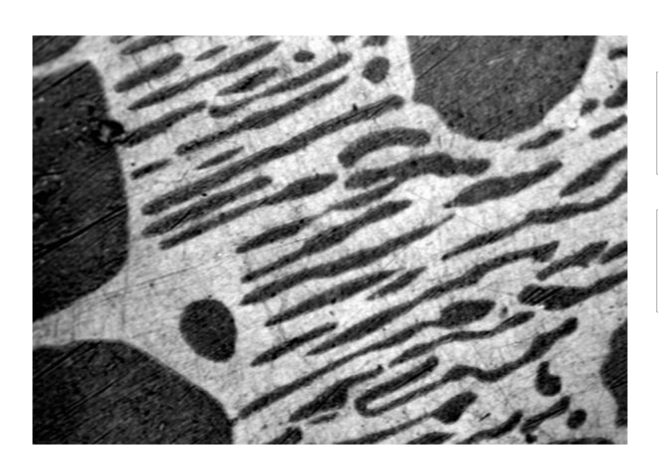
- (1) File Open Samples Blobs (inverse)
- (2) File Open Samples Clown

(1) The sholds

- √ Image Lookup Tables Invert LUT
- ✓ Process Binary Make Binary (default method)
- ✓ Image Adjust threshold: Adjust the thresholds, then set
 them to make binary
- ✓ Image Adjust Auto Threshold and Auto Local Threshold
- ✓ Many more methods, and "local" method

(2) Statistical Region Merging

Edge Detection: The Sobel filter



Images may contain objects

These objects have edges

How can we find the edges?

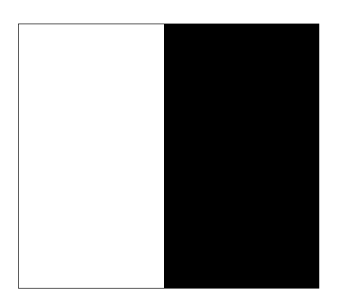
Edge Detection

What is an "edge"?

"Hard Edge" - Adjacent black / white pixels

<u>"Soft / Fuzzy Edge"</u> - common in images. Especially for small diffraction limited objects like vesicles/membranes.

Noise makes edges look softer

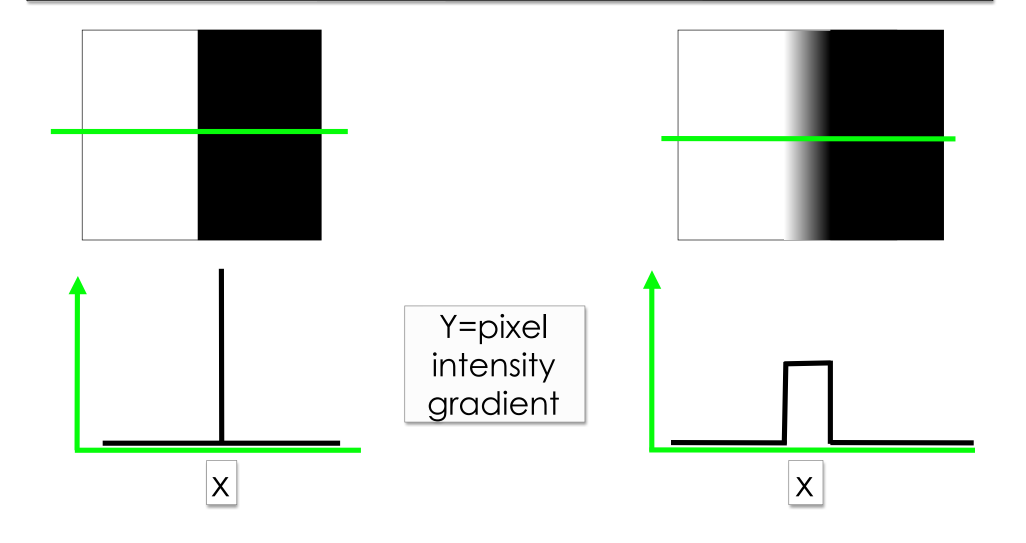




Edge Detection "Image Gradient"

What is a "Gradient Image"?

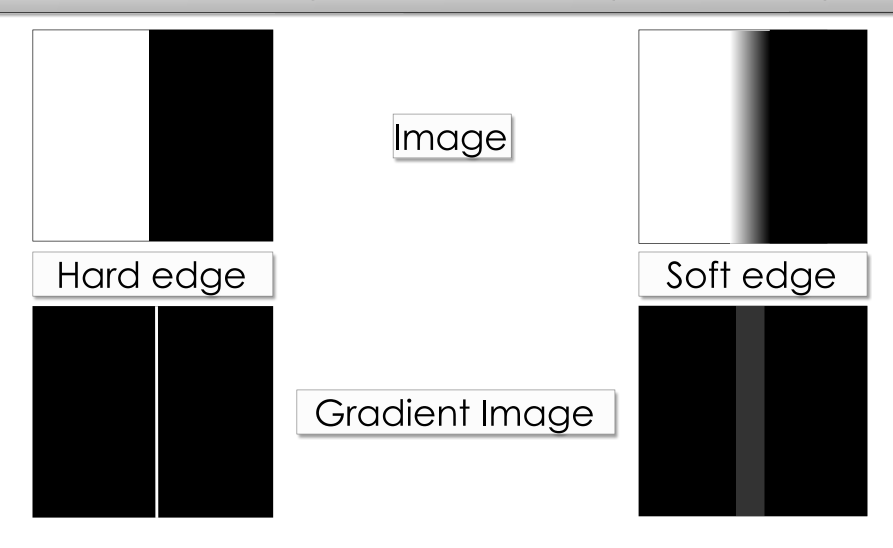
Rate of change of pixel intensity (1st derivative)



Edge Detection "Image Gradient"

What is a "Gradient Image"?

Rate of change of pixel intensity (1st derivative)



"Image Gradient" - How?

Sobel filter - 3x3 convolution filter pair in x AND y

- ✓ find edges with x and y components
- ✓ compute total gradient magnitude
- ✓ approximates 1st derivative of image

-1	0	+1
-2	0	+2
-1	0	+1

+1	+2	+1
0	0	0
-1	-2	-1

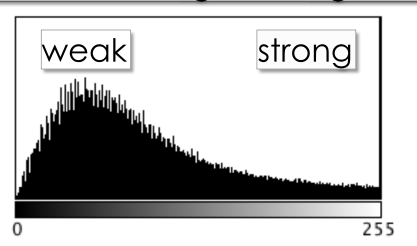
$$output = \sqrt{g_x^2 + g_y^2}$$

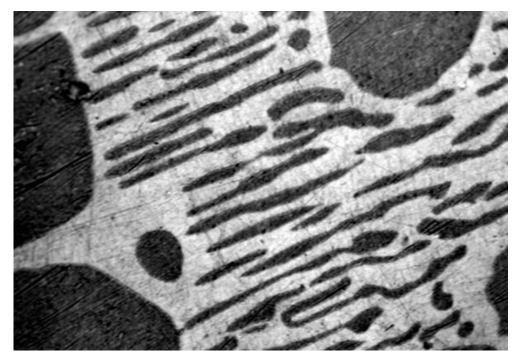
Gradient Image - Real Sample:

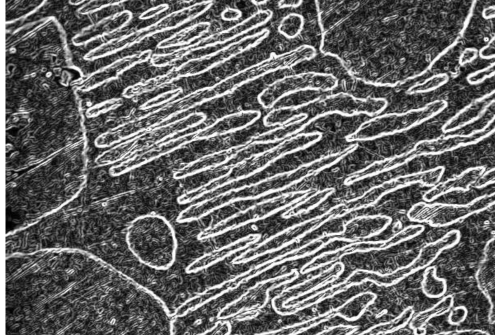
Real / Biological images:

- ✓ Sobel filter
- ✓ many edges
- ✓ many weak edges from noise

gradient image histogram



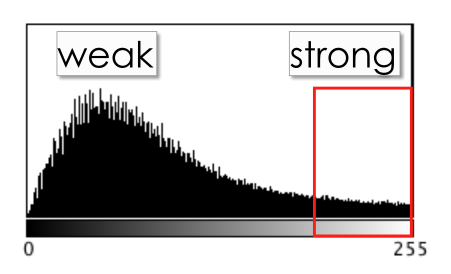


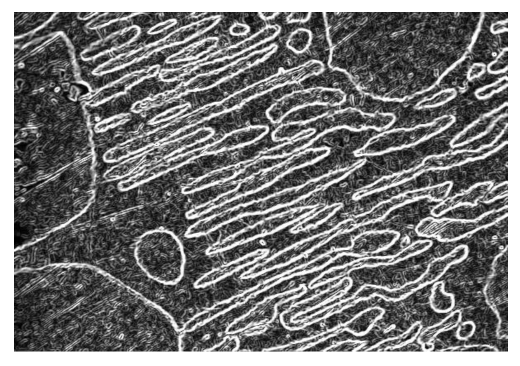


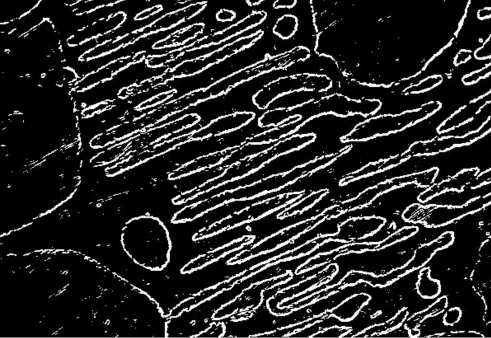
Gradient Image - Strong Edges?

Remove weak edges?

- ✓ Threshold the gradient image
- ✓ Smoothing filter beforehand







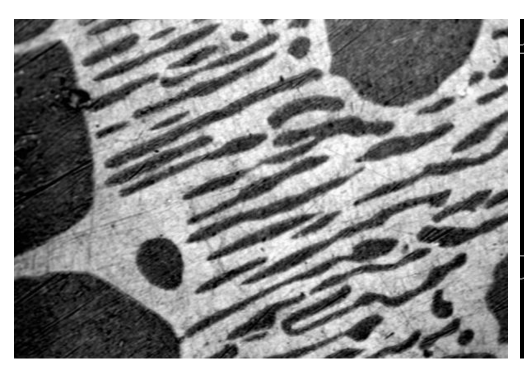
"Canny" Edge Detection

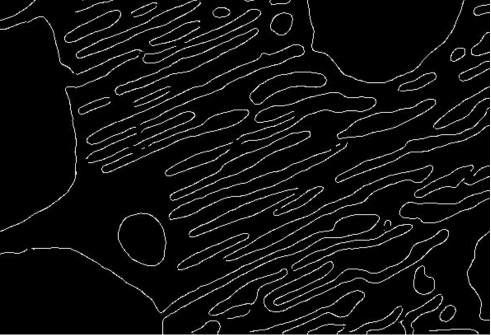
Remove weak/noisy edges - keep strong

Gaussian smooth image + hysteresis threshold gradient image

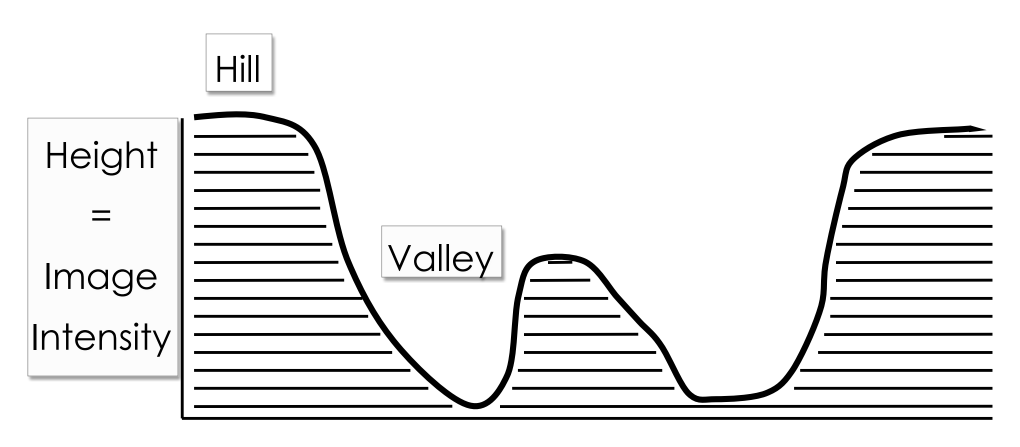
Make edges sharp - 1 pixel wide

Non maximal suppression of gradient image





... mountains, lakes and oceans



View From the Side

... mountains, lakes and oceans

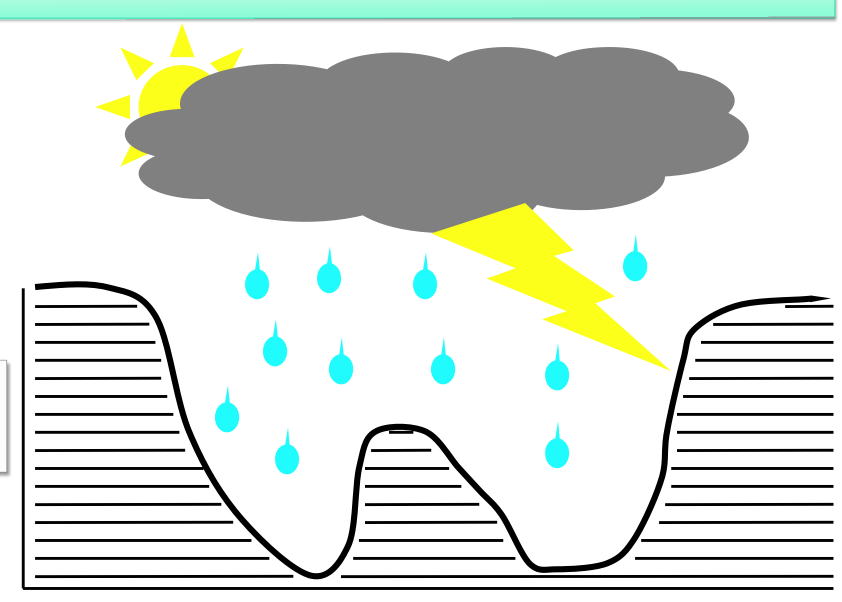


Image Intensity

... mountains, lakes and oceans

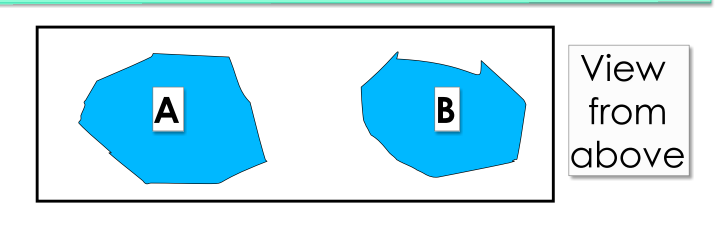
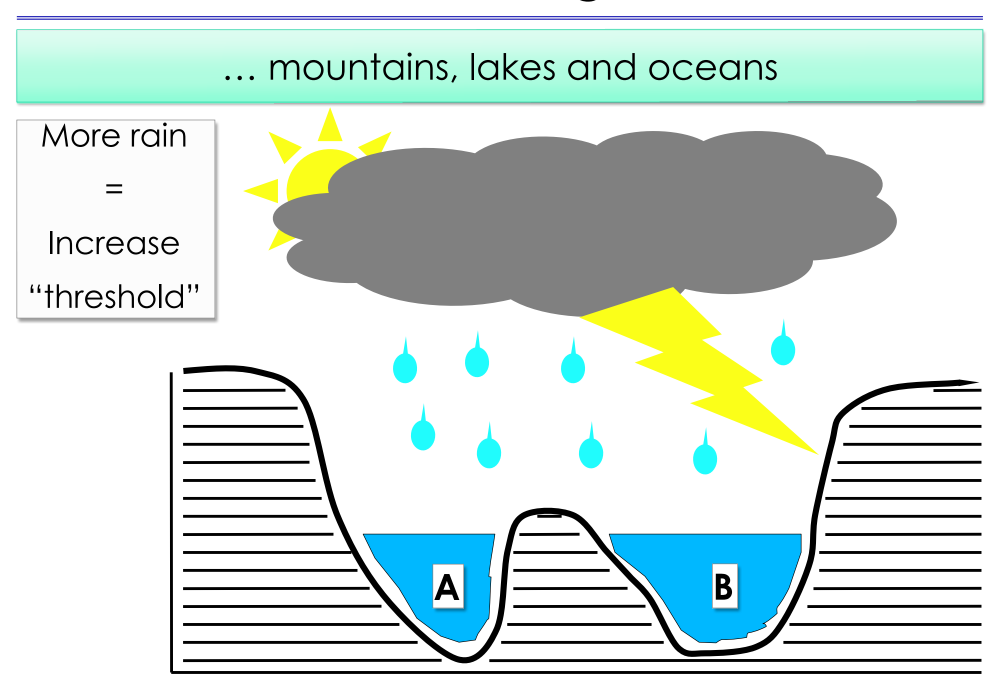


Image Intensity

A

B

2 flooded areas



... mountains, lakes and oceans

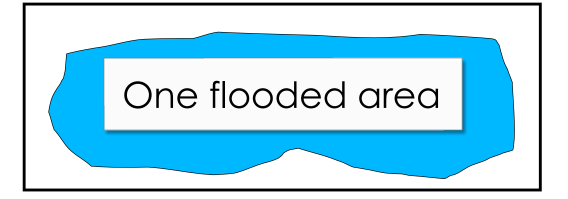
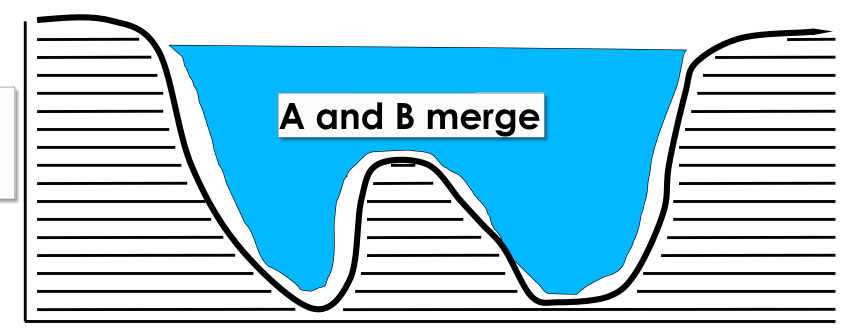
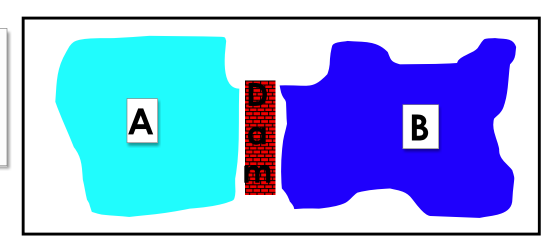


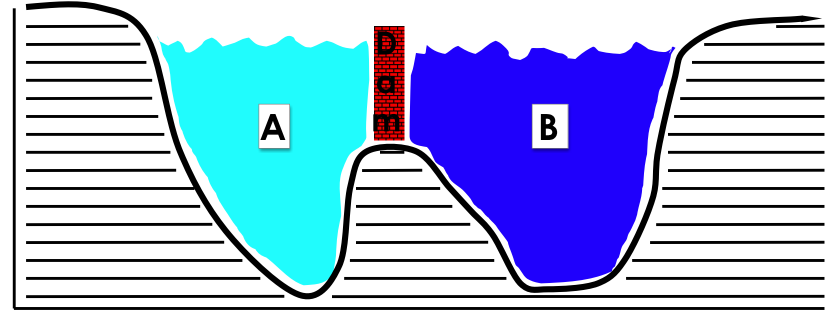
Image Intensity



... mountains, lakes and oceans

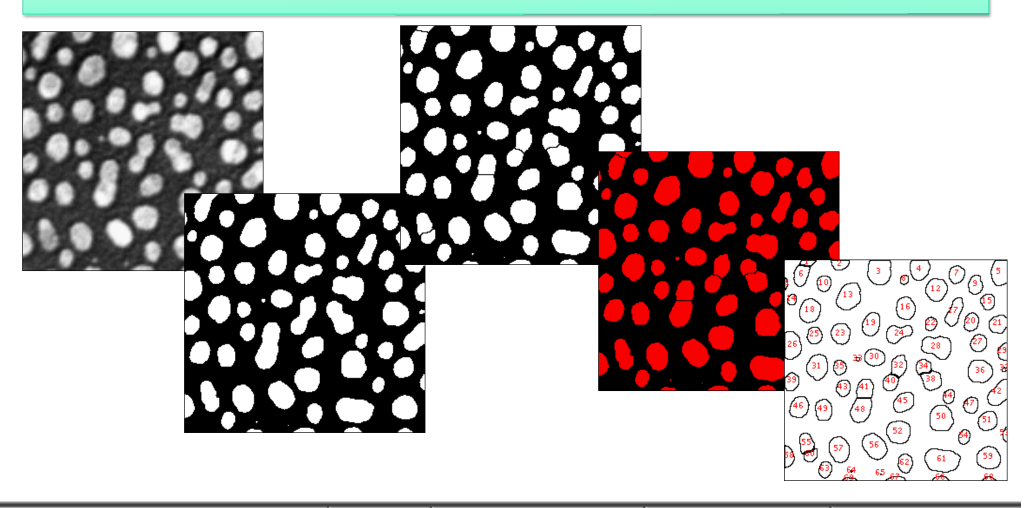
Make a "Dam" at the "Watershed line"





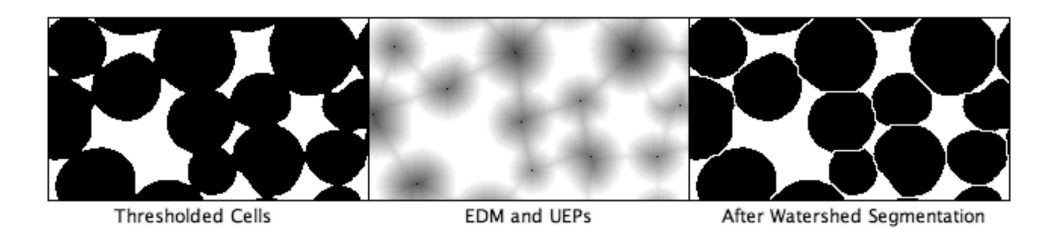
Watershed - to find object number

Sometimes there are just too many to count by hand



Slice	Count	Total Area	Average Size	Area Fraction
blobs-bin-WShed-inv.tif	69	22159.000000	321.144928	34.1

Watershed to separate touching objects



- ✓ Euclidian Distance Map
- ✓ Ultimate Eroded Points
- ✓ Fill with water from UEP until hits edge of object, or dams between objects

Practical Session 2d



Getting to know "Fiji" better –

Fiji is just ImageJ (Batteries included)

http://pacific.mpi-cbg.de

File - Open Samples - Blobs

Watershed Segmentation and Analysis

- ✓ <u>Invert, Make Binary, Watershed, Threshold, Analyze Particles:</u> Separate and measure touching objects
- ✓ Search the Wiki for NucleiWatershedSegmentation tutorials

Links and Further Reading

Standard Text Book:

Digital Image Processing 2nd Ed., Gonzalez and Woods, Prentice Hall

Fiji and ImageJ:

- ✓ Fiji Wiki and docs: http://pacific.mpi-cbg.de (also:Installation)
- ✓ <u>ImageJ home:</u> http://rsb.info.nih.gov/ij/
- ✓ ImageJ Doc.Wiki: http://imagejdocu.tudor.lu/doku.php
- ✓ MacBioPhotonics plugins collection for microscopy:

http://www.macbiophotonics.ca/downloads.htm

Image Processing Facility

- ✓ <u>Intranet</u> Services and Facilities Image Processing Facility
- ✓ Wiki info for beginners tips software documentation:

https://wiki.mpi-cbg.de/wiki/imagepro/index.php/Main_Page

Imaging Facility Network (IFN): https://ifn.mpi-cbg.de

Email: ipf(at)mpi-cbg.de