Übungen zur Vorlesung "Methoden der Bioinformatik"
Teil Bildanalyse (Computer Vision), BMCV Group, PD Dr. K. Rohr

Linzzfor notehooks
Wintersemester 2024/2025

Python introduction and Jupyter notebooks

Course web-page: http://www.bioquant.uni-heidelberg.de/research/groups/ biomedical-computer-vision/teaching/compmeth/Labsessions

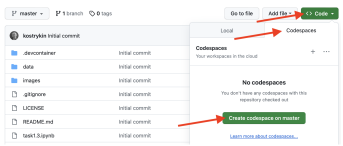
If working in the BioQuant computer room, please **use Chrome instead of Firefox** as the web browser, because the installed version of Firefox is known to have issues with GitHub. Using Firefox is fine when working on your personal computer.

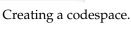
## 1 Setting up your GitHub repository

- 1. Open the course web-page in any web-browser of your choice.
- 2. Click on the link in "Create a new GitHub repository by using this link".
- 3. This will load another web-page entitled "Create a new repository". Leave everything on default and confirm the creation of the repository by clicking the "Create repository" button.
- 4. You should see a "Generating your repository" message for a few seconds and then be presented with an **overview of your repository**. You just created your first GitHub repository congrats! :)

## 2 Firing up a GitHub Codespace

- 1. Open the **overview of your repository** on GitHub. This is the web-page that you landed on after completing Task 1.
- 2. Click on the green "Code'" button, then select the "Codespaces" tab.
- 3. Click the green "Create codespace on master" button. This will load **VS Code** (Visual Studio Code) inside of your web-browser. Wait until everything is loaded, it may take about one minute.







Deleting a codespace.

Note: If, at any time, VS Code behaves weirdly (e.g., complaining about missing extensions, not loading notebooks, not finding kernels, or similar), try to reload the VS Code window. To do that, press Ctrl + 1 + P (or + P if you are on macOS) and type "Reload window ". Your work progress will be preserved. If this does not solve the issue, make sure your work is committed and pushed (see Task 3), then go to https://github.com/codespaces and delete the codespace. Then, re-create the codespace by following the steps 1–3 described above.

## 3 Your first Jupyter notebook

The left panel of **VS Code** shows an overview of **your local repository**. Right now it is identical to your GitHub repository. The assignments of this course are organized into several Jupyter notebooks. These are the task\*.ipynb files that you can see in your repository. By progressing from task to task, you will work with different notebooks.

- 1. Double click the file task1.3.ipynb inside of **VS Code** to open the notebook for this task, and follow the instructions inside the notebook. **Note:** In Jupyter notebooks, code cells can be run in an *arbitrary order*. This is very helpful for experimenting and trying out new things. Nevertheless, an assignment in this course is *only* considered "finished" when the results can be reproduced by rerunning all code cells *from top to bottom* by clicking the "Run all" button.
- 2. When finished, close your notebook. Changes are saved automatically within **your local repository**, but remember, that those changes will be lost when you close the codespace, unless you push them to your GitHub repository.
- 3. To do that, click on the "Terminal" tab at the bottom of **VS Code**, type the following Git command, and press to execute it:

```
git commit --all -m "Finish task 1.3"
```

The text "Finish task 1.3" is the **commit message**, which is arbitrary. Describe what changes you have done since your previous commit.

If you wanted, you could revert to any previous commit at a later time, and choosing an expressive message is convenient for finding the commit which you will be looking for. There also are some conventions for how a **commit message** should be formatted: It should tell in an "imperative mood" and as concisely as possible, what *the committed changes* are supposed to do<sup>1</sup>.

If done correctly, the output of the above command should be something like "1 file changed, 12 insertions(+), 1 deletion(-)", but the exact numbers may vary.

Finally, type "git push" and press to push the committed changes to your GitHub repository. If done correctly, there should be multiple lines of output, concluding with a line similar to "1d2e403..a387645 master -> master".

# 4 Writing loops in Python

Now you already know how to edit, commit, and push a Jupyter notebook.

- 1. Open the notebook task1.4.ipynb and follow the instructions in the notebook.
- 2. Commit and push your changes, then close **VS Code** (the browser tab/window).

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<sup>&</sup>lt;sup>1</sup>From the official Git documentation: https://git.kernel.org/pub/scm/git/git.git/tree/ /Documentation/SubmittingPatches?h=v2.36.1#n181

Übungen zur Vorlesung "Methoden der Bioinformatik" Teil Bildanalyse (Computer Vision), BMCV Group, PD Dr. K. Rohr Wintersemester 2024/2025

Images, Histograms, Intensity clipping

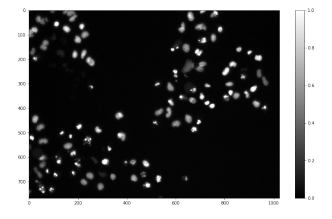
# 1 Image IO (input/output)

- 1. Open VS Code in a GitHub Codespace (see Task 2 of Lab Session 1) using the repository which you created before (see Task 1 of Lab Session 1).
- 2. Open the notebook task2.ipynb.
- 3. Enter the following code into the *first* code cell of the notebook and run it:

```
import numpy
import matplotlib.pyplot as plt
```

#### Notes:

- The first instruction should be clear from the lecture. The second instruction loads the module "matplotlib.pyplot" and makes it available by using the abbreviation "plt". This module is useful for visualizing data.
- Make sure this code cell *always* remains the very first code cell of your notebook, so it is run first when the Notebook is re-run from top to bottom.
- 4. Then, extend the notebook as follows:
  - (a) Use img = plt.imread('data/cells.png') to load an image. Note: The type of the returned object (img) is numpy.ndarray (also see the introduction on page 2). Objects of this type represent images.
  - (b) Use plt.figure() (or, e.g., plt.figure(figsize=(15,8)) if you want to specify the size of the figure) to add a *figure* to a code cell of the notebook. Then, within the same code cell, use plt.imshow(img, 'gray')<sup>1</sup> to display the image within the figure. In addition, use plt.colorbar() after the imshow-instruction to include a legend of the gray-scale encoding. Run the code cell with and without the colorbar-instruction and observe the differences. Finally, you should obtain an output like this:



<sup>&</sup>lt;sup>1</sup>The parameter "'gray'x' in "plt.imshow(img, 'gray')" specifies the color map for a *monochromatic* image (i.e. a two-dimensional array of image *intensities*, non-RGB) for visualization (e.g., gray-scale)

## Introduction: Working with numpy.ndarray objects

If img is an object of the type numpy.ndarray, then the object img has the following attributes and methods:

Attributes (data)	Methods (behaviours)	
<pre>img.ndim: Corresponds to the dimen- sion of img.</pre>	<pre>img.copy(): Tells img to return a copy of   itself.</pre>	
<pre>img.shape: If img is an image, then     the element img.shape[0] corre-     sponds to the image height (num-     ber of rows) and img.shape[1]     to the image width (number of     columns).</pre>	<ul><li>img.clip(t1, t2): Tells img to return a copy of itself using intensity clipping (see Task 3).</li><li>img.flatten(): Tells img to return a flat representation of itself (see Task 2).</li><li>(Methods are like functions which belong)</li></ul>	
	to objects.)	

Note that the above is *not* a complete list (there are many more attributes/methods<sup>2</sup> which are *not* relevant for this assignment).

## Further hints regarding numpy.ndarray objects:

- 1. The pixel in the upper left corner of the image has the coordinates (0,0), and the pixel in the lower right corner has the coordinate (width -1, height -1).
- 2. If img is an image (i.e. object of the type numpy.ndarray), then the intensity value of a pixel at position p corresponds to img[p], where p has two elements (row, column). Alternatively, you can use img[row, column] instead of img[p].

# 2 Histograms

Extend your notebook by a histogram of the previously loaded image – **Hints:** 

- 1. The function plt.hist(data) produces a histogram of a sequence of values (data).
- 2. In Python, a *sequence* is, for example, a list, a *flat* array, . . .
- 3. The image img is a *two-dimensional* array and img.flatten() returns a flat representation (sequence of all pixel values).

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<sup>&</sup>lt;sup>2</sup>see https://docs.scipy.org/doc/numpy/reference/generated/numpy.ndarray.html

# 3 Intensity clipping

Given two thresholds  $T_1$  and  $T_2$ , each pixel x, y of the image with an intensity value g(x,y) less than the threshold  $T_1$  is assigned the value  $T_1$  and each pixel with a value greater than  $T_2$  is assigned the value  $T_2$ . Pixels with a value between the two thresholds  $T_1$  and  $T_2$  remain unchanged:

$$g_{\text{clip}}(x,y) = \begin{cases} T_1 & \text{if } g(x,y) < T_1, \\ T_2 & \text{if } g(x,y) > T_2, \\ g(x,y) & \text{otherwise} \end{cases}$$

If g is the image img, y is the image row, and x is the image column, then the mathematical expression g(x, y) corresponds to img[row, column] in Python.

In this task, you will **perform intensity clipping** (i.e. compute  $g_{\text{clip}}$ , in *three* different ways) using the previously loaded image (Task 1) and visualize the result using the imshow function.

- 1. Perform intensity clipping using the method clip of numpy.ndarray.
- 2. Reproduce the behaviour of the method clip, i.e. perform intensity clipping *without* using the method clip!
  - (a) Use *two* nested for-loops (one outer loop for the image rows or columns, one inner loop for the pixels of the current row/column) and if-conditions.
  - (b) Use a *single* for-loop and if-conditions **Hint:** numpy.ndindex(img.shape) yields an iterable. The items of this iterable correspond to *all* pixel coordinates of the image img. Each item is a pair of coordinates (i.e. row and column). Use "[0]" and "[1]" to access the corresponding row and column of an item.

Include a legend of the gray-scale encoding (using "plt.colorbar()") in each figure!

# 4 Writing re-usable code (BONUS)

In the previous task, you have used loops to reproduce the behaviour of the method clip of numpy.ndarray. Now, make this code *re-usable* by putting it into a *function* which you can use anytime later. For your convenience, a skeleton of the code you need to write to implement the function is already added to the notebook (see "def clip\_image...").

#### Übungen zur Vorlesung "Methoden der Bioinformatik" Teil Bildanalyse (Computer Vision), BMCV Group, PD Dr. K. Rohr Wintersemester 2024/2025

Mean, Median, and Gaussian filtering

**Preparation.** Open the notebook task3.ipynb in VS Code (see Task 2 of Lab Session 1). Enter the "import numpy" and "import matplotlib.pyplot as plt" instructions into the *first* code cell of the notebook and run it (cf. Lab Session 2).

## 1 Linear filtering by convolution (mean filter)

In this task you will implement and test a *mean filter* (box filter).

- 1. Use imread and imshow to load and show the image data/lena.png.
- 2. Finish the implementation of the re-usable function meanfilter. The function parameters are img (the image to be filtered) and size (filter size determining the filtering neighborhood). You may assume that size is an odd number. The function must return the filtered result image. Do *not* modify the input image img!
- 3. Test your solution by using the function meanfilter for the previously loaded image (e.g., set the filter size to 3). If something does not work as expected, look for errors you have made and fix them. Repeat fixing and testing until everything works.
- 4. Compare your result for filter size 5 with the correct result image data/lena\_mean-filter5.png. Use imread to load the image. For a *quantitative* comparison of two images, use the instruction "assert numpy.allclose(img1, img2, atol=1/255)" where "img1" and "img2" are the two images **Notes:** 
  - (a) The instruction "assert condition" interrupts the code execution if condition is False, rising your attention to an error. Otherwise, nothing happens.
  - (b) In contrast to a comparison using the mathematical equality operator ("img1 == img2"), a comparison using allclose tolerates *numerical inaccuracies*. The parameter "atol=1/255" specifies the maximum tolerated difference per pixel. Since the PNG file quantifies image intensities as multiples of 1/255, errors lower than 1/255 cannot be distinguished from numerical inaccuracies.

#### **Hints:**

- 1. To create a new numpy.ndarray object representing an image with height shape [0] and width shape [1], initially filled with zeros, you can use numpy.zeros(shape).
- 2. You can use two nested for-loops: An *outer* for-loop to iterate over all pixels of the image and an *inner* for-loop to iterate over all pixels of the filtering neighborhood.
- 3. To iterate over all pixels of the image or the filtering neighborhood using a for-loop, ndindex can be used (cf. Lab Session 2, Task 3.2(b)).
- 4. Bear in mind the border problem, i.e. you should not access pixels where the neighborhood is partially outside the image.
- 5. Also consider the schematic illustration in Figure ??.

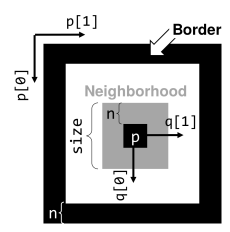


Figure 1: Schematic illustration of a size  $\times$  size filtering *neighborhood* and the image *border* with  $n = \lfloor (\text{size} - 1)/2 \rfloor$ , assuming that size is an *odd* number  $\geq 3$ . The image pixel p corresponds to the *center* of the filtering neighborhood (i.e. the *origin* of the coordinate system of the filtering neighborhood).

## 2 Non-linear filtering (median filter)

In this task you will implement and test a *median filter*.

- 1. Finish the implementation of the re-usable function medianfilter. The function parameters are img (the image to be filtered) and size (filter size). You may assume that size is an odd number. The function must return the filtered result image. Do *not* modify the input image img!
- 2. Analogously to Task 1, test your solution by using the function medianfilter and *quantitatively* compare your result for filter size 5 with the correct result image data/lena\_medianfilter5.png.

#### **Hints:**

- 1. You can use a list to store the pixel values of the filtering neighborhood. An empty list is created by "list()" or the shorthand expression "[]".
- 2. The function sorted(sequence) returns a sorted sequence (e.g., a list, a *flat* numpy.ndarray). Alternatively, both list and numpy.ndarray objects provide the method sort(). Using this method tells the object to sort itself. **Examples:**

Example 1	Example 2	Output
<pre>data = [4, 3, 8, 2] print(sorted(data))</pre>	<pre>data = [4, 3, 8, 2] data.sort() print(data)</pre>	[2, 3, 4, 8]

If data is a numpy.ndarray object with data.ndim == 2, then invoking the method data.sort() sorts the values of each row of data independently from the other rows.

### 3 Using pre-implemented filters

- 1. Load the module scipy.ndimage using the instruction "import scipy.ndimage".
- 2. Use the following pre-implemented filters in this module and include the filtering results (via imshow) into your notebook:
  - (a) Use scipy.ndimage.uniform\_filter(img, size) for a mean filter.
  - (b) Use scipy.ndimage.median\_filter(img, size) for a median filter.
  - (c) Use scipy.ndimage.gaussian\_filter(img, sigma) for a Gaussian filter (where "sigma" is the standard deviation of the Gaussian function).
- 3. Compare the results obtained using the functions in 3.2(a) and 3.2(b) with those you obtained in Tasks 1 and 2. What are the main differences? Do you have an explanation? Use a *Markdown* cell to write your answer, i.e. change the cell type!

## 4 Slicing and benchmarking (BONUS)

In this task, you will implement a filtering function which is *faster* than those you implemented in Tasks 1 and 2. You will also learn how to *benchmark* the run time of code.

- 1. Decide which filtering method you want to accelerate (mean or median filter). If you are unsure, choose the one whose solution you are more confident with.
- 2. Finish the implementation of the re-usable function fastfilter. The function parameters are img (the image to be filtered) and size (filter size). The function must return the filtered result image. Do *not* modify the input image img!

**Important:** Use the hints below to confine your code to only a single for-loop:

- (a) If img is an object of the type numpy.ndarray (e.g., an image), then img[i0:i1, j0:j1] corresponds to the rectangular subsection of the image (also called *slice*) ranging from row i0 to i1-1 and column j0 to j1-1 (all inclusive). The subsection itself also is of type numpy.ndarray.
- (b) For mean filters: The mean value of an numpy.ndarray object can be computed using its mean() method (e.g., img.mean() if img is an object of the type numpy.ndarray).
- (c) For median filters: The method flatten() of numpy.ndarray objects (cf. Lab Session 2) yields a flat representation (which can be sorted).
- 3. Test your solution by using the function fastfilter for the previously loaded image and *quantitatively* compare your result to that you obtain using meanfilter or medianfilter, respectively.
- 4. Use the instruction "%timeit fastfilter(img, 9)" to benchmark the run time of your fastfilter implementation. Use a similar instruction to benchmark the run time of meanfilter or medianfilter, respectively.
- 5. Document your observations and try to think of an explanation (use Markdown!)

## **Edge detection: Derivative operators**

**Preparation.** Open the notebook task4.ipynb in VS Code (see Task 2 of Lab Session 1). Enter the "import numpy" and "import matplotlib.pyplot as plt" instructions into the *first* code cell of the notebook and run it (cf. Lab Session 2).

#### 1 Prewitt filter

- 1. Use imread and imshow to load and show the image data/lena.png.
- 2. Compute the partial derivatives  $g_x$  and  $g_y$  of an image g(x,y) by convolving the image with Prewitt derivative operators (see Figure 1). To this end, finish the implementation of the re-usable functions prewitt\_h and prewitt\_v. The functions are supposed to compute and return the *convolution* of the input image img with the horizontal and vertical Prewitt derivative operators, respectively. **Hints:** 
  - (a) Do *not* modify the input image! Avoid the border problem by computing the partial derivatives  $g_x$  and  $g_y$  only for those pixels which have their neighborhood completely inside the image.
  - (b) If you do not know where to start, start from your implementation of the mean filter (Lab Session 3, Task 1.2). The  $3 \times 3$  mean filter performs a convolution using a uniform filter mask (weighting each image pixel equally, dividing by  $3 \times 3 = 9$ ). You only need to change the weights based on the values of "q"!
- 3. Test your implementations for prewitt\_h and prewitt\_v by including images of the computed partial derivatives into your notebook. Use colorbar() after each imshow-instruction to also include a legend of the gray-scale encoding.
- 4. Quantitatively compare your results obtained using prewitt\_h and prewitt\_v with the correct result images data/lena\_prewitt\_h.tiff and data/lena\_prewitt\_v.tiff, respectively (for quantitative image comparison, recall Lab Session 3, Task 1).

**Important hint:** Use skimage.io.imread instead of imread to load a TIFF file. However, remember to first *load* the skimage.io module (use "import skimage.io").

**Rule of thumb:**<sup>1</sup> There is no reason for not *always* using skimage.io.imread (and not using plt.imread) except that you have to remember to load the module.

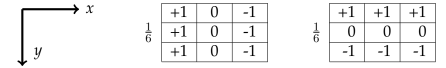


Figure 1: Prewitt derivative operators

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<sup>&</sup>lt;sup>1</sup>e.g., for the exam  $\ddot{-}$ 

## 2 Edge detection

1. Compute the *magnitude* of the image gradient

$$\left\|\nabla g\left(x,y\right)\right\| = \sqrt{g_x^2\left(x,y\right) + g_y^2\left(x,y\right)}$$

and include the resulting image into the notebook. Hints:

- (a) When working with objects of the type numpy.ndarray (e.g., images), mathematical operations are *propagated* to the intensity values of the image. For example, if img is an image of the type numpy.ndarray, then the expression img\*2 yields an image with *doubled* intensities (in comparison to img).
- (b) The square root of an intensity value (or all values in an image that is a numpy.ndarray object) can be computed using the numpy.sqrt function (e.g., numpy.sqrt(value) or numpy.sqrt(img)).
- Quantitatively compare your result with the correct result image data/lena\_prewitt\_gradmag.tiff.

## 3 Sobel filter (BONUS)

Repeat Task 1 using the Sobel filter instead of the Prewitt filter:

- 1. Compute the partial derivatives  $g_x$  and  $g_y$  by convolving the image g(x, y) with Sobel derivative operators (see Figure 2). To this end, implement the re-usable functions sobel\_h and sobel\_v analogously to prewitt\_h and prewitt\_v.
- 2. Test your implementations for sobel\_h and sobel\_v by including images of the computed partial derivatives into your notebook (also include color legends).
- Quantitatively compare your results using sobel\_h and sobel\_v with the correct result images data/lena\_sobel\_h.tiff and data/lena\_sobel\_v.tiff, respectively.

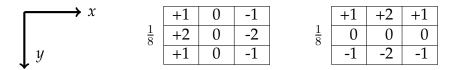


Figure 2: Sobel derivative operators