Atlas-based Imaging Data Analysis pipeline for functional and structural MRI Data

$\begin{array}{c} \textbf{AIDAmri} \\ \textbf{v1.2} \end{array}$

Code: Leon Scharwächter, Niklas Pallast, Michael Diedenhofen, Victor Vera Frazão, Markus Aswendt Manual: Leon Scharwächter, Niklas Pallast, Tim Fleiner, Victor Vera Frazão, Markus Aswendt

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Department of Neurology
University Hospital Cologne

Contents

1	Intr	roduction	2
2	Inst	callation	4
3	Fun	actions	5
4	Bat	ch processing	6
5	Processing single files step-by-step		9
	5.1	Convert raw data	9
	5.2	Processing of T2w & T2mapping data	10
	5.3	Processing of T2 data	12
	5.4	Processing of DTI data	12
	5.5	Processing of fMRI data	15
	5.6	Peri-infarct ROI analysis	17
6	Docker usage		18
	6.1	Prerequisites	18
	6.2	Creating image	20
	6.3	Running container and mount data	21

1 Introduction

The Atlas-based Processing Pipeline for functional and structural MRI data (AIDAmri) was developed for automated processing of mouse brain MRI. AIDAmri works with T2-weighted MRI (T2w), diffusion weighted MRI or diffusion tensor imaging (DTI) and resting-state functional MRI (fMRI). The Allen Mouse Brain Reference Atlas (ARA, CCF v3) is registered on each of these MRI data sets and is used to analyse regions of interest. Furthermore, the regions of the ARA are used as seed-points for the connectivity and activity matrices. The lib User-defined ROIs and masks can be generated separately and used for analysis, e.g. stroke lesion masks and peri-infarct regions. AIDAmri comes with different atlas and template versions, which are necessary for the registration to work (Figure 1): a) annotation 50 changed

anno: original ARA CCF v3 labels (regions with grey values;100k changed to new values starting with 2000), b) anno volume 2000 rsfMRI: atlas from a) with reduced number of atlas regions by selective region fusion, 96 regions in total, split between hemispheres (right side +2000), c) same as in b) but no split, d) same as in a) but split, e) original ARA template with 50 um isotropic resolution, f) custom-made MRI template. For the complete list of atlas labels see: annoVolume+2000_rsfMRI.nii.txt

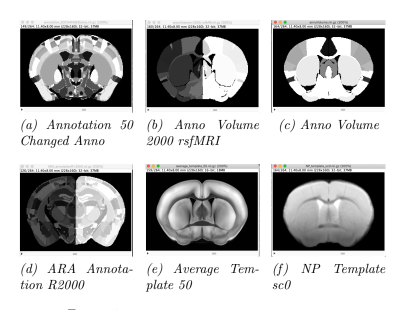


Figure 1: Atlases included in the /lib folder.

2 Installation

This installation guide refers to the native installation directly on your host machine. To install the Docker image, please read section 6.

1. Download the folders /bin and /lib by using this link /bin and /lib should be located in the same directory

The folder /bin contains the python scripts necessary for all preprocessing steps. The folder /lib contains several information about six atlases. Please do not conduct any changes to these files as it could influence the access from the python scripts.

- 2. Download & Install DSI-Studio and copy the install path into .../bin/3.2_DTIConnectivity/dsi_studioPath.txt
- 3. Download & Install FSL 5.0.1
- 4. Download & Install Cmake, open the software and click in the upper menu on $Tools \rightarrow How To Install For Command Line Use like shown in Figure 2 and follow the steps$

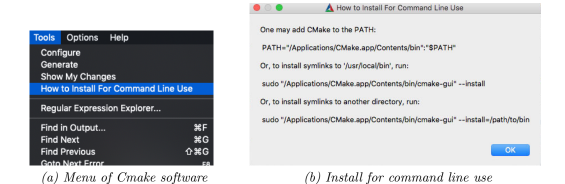


Figure 2: Installation instructions to install Cmake.

5. Download & Install Python 3.6 or higher using Anaconda and enter the following command to install necessary packages

pip install nipype==1.1.2 lmfit==0.9.11 progressbar2==3.38.0

nibabel shutil (without line break!)

Anaconda will tell you if additional packages are necessary. We recommend to install AIDAmri in a separate Anaconda environment (see the related documentation).

- 6. Install NiftyReg by conducting the following steps:
 - a. Generate your source folder .../NiftyReg/niftyreg_source
 - b. Download NiftyReg from the git by replacing <path> by your personal path and enter the following command

git clone

git://git.code.sf.net/p/niftyreg/git <path>/niftyreg_source
(without line break!)

- c. Change the folder by typing in the terminal:
 cd <path>/niftyreg_source
- d. Type in the command line git reset --hard 83d8d1182ed4c227ce4764f1fdab3b1797eecd8d
- e. Follow the steps described here

3 Functions

List of functions:

- 1_PV2NIfTiConverter: Bruker to NIfTy converter
- 2.1_T2PreProcessing: T2w MRI pre-processing including brain extraction, bias field correction and atlas registration
- 2.2_DTIPreprocessing: DTI pre-processing including brain extraction, bias field correction and atlas registration
- 2.3_fMRIPreProcessing: DTI pre-processing including brain extraction, bias field correction and atlas registration

- 3.1_T2Processing: Stroke mask calculations across all subjects per group (Incidence mapping), SNR calculations
- 3.2_DTIConnectivity: Whole brain fiber tracking using DSI Studio and calculation of diffusion measures (FA, AD, RD, MD) for every brain region
- 3.2.1_DTIdata_extract: Creates a .txt-file containing DTI values from all brain regions
- 3.3_fMRIActivity: functional connectivity analysis for all atlas regions
- 4.1_ROI_analysis: Analysis of T2w, DTI, and rs-fMRI with user-defined atlas regions, e.g.: peri-infarct regions around the stroke lesion

Attention: Please note that the Docker image does not support the GUI. All program examples are only listed with the mandatory input parameters. For more details/help, call python .../python <command> -h. The command line examples are given with the identifier testData<No.> .nii.gz and can be identically applied to other data. The test data is freely available: https://doi.org/10.12751/g-node.70e11f.

After a successful download, you can choose to either process single files manually or automate the processing for the whole dataset. In both cases, processing includes file conversion from the raw Bruker format into the NIfTI format, several preprocessing steps and registration with the Allen Brain Reference Atlas. The functions in /bin are named according to the MRI sequence to be processed (T2, DTI, fMR) and labeled with 1-4.1 in the order of processing: for example 3.1_T2Processing requires 1_PV2nIfTiConverter and 2.1_T2PreProcessing to be executed sequentially.

4 Batch processing

To process the whole project folder at once, only two scripts are necessary: At first conv2Nifti_auto.py creates a new project folder, converts all files to the NIfTI format and stores them in the new project folder, batchProc.py then applies pre-processing steps and the registration with the atlas. Remember that our test data set already is converted into the NIfTI format.

In this case only the second script needs to be applied. In general, the raw (Bruker format) data needs to be in the following structure for the first script to work: projectfolder/days/subjects/

To convert the whole project folder into the NIfTI format, open the terminal and change the directory to the /bin folder of the AIDAmri installation using cd <path to AIDAmri>/bin

Enter a similar line as in the example below to start the the first script (Bruker to NIfTI conversion). For the first script to work, the table group-Mapping.csv within /bin needs to be adjusted beforehand: specify the group name of every subject within the project folder (e.g. using Excel). This is necessary for the script to properly generate the new processed project folder structure. Remember that the algorithm does not read the first row of the table containing the titles (Subject, Group).

Example:

```
python conv2Nifti_auto.py -f /path/to/raw_dataset
-d Baseline P1 P7 P14 -g Group1 Group2
(without line break!)
```

As you can see, this script computes the conversion either for all data in the raw project folder or for certain days and/or groups specified through the optional arguments -d and -g. During the conversion a new folder called proc_data is being created in the same directory where the raw data folder is located.

After a successful Bruker to NIfTI conversion, the second script can be applied to the new project folder proc_data.

Example:

```
python batchProc.py -f /path/to/proc_data -g Group1 Group2
-d Baseline P1 P7 P14 -t T2w DTI fMRI
(without line break!)
```

This script runs every necessary script for all (pre-)processing and registration steps. The data needs to be ordered like after the Bruker2NIfTI conversion: projectfolder/days/groups/subjects/data/. Again, you can specify which days -d and groups -g to compute and which dataformat -t (T2w, DTI, fMRI) to process. Please keep in mind that the hereby executed scripts are related to each other and therefore T2w always needs to be specified before DTI.

Dependent on the size of your project, this process may take a while. After finishing, your project folder is ready for network graph analysis, e.g. using AIDA-connect.

5 Processing single files step-by-step

Open the AIDA_gui.py in the /bin folder. You can use the GUI to process the test data set, which represents one subject on one day of measurement. Type the following command line

python AIDA_gui.py

to open the window shown in Figure 3.

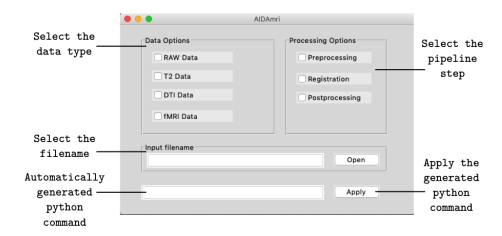


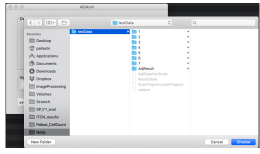
Figure 3: Description of the user interface to process the data step by step.

5.1 Convert raw data

Convert Bruker raw data to NIfTI files by specifying the folder containing all raw folders of each scan (see Figure 4). A file with exactly the same name is created in the given input folder. It contains all sorted NIfTI files. The raw data should have the same orientation as the example dataset.

python pv_conv2Nifti.py -i .../testData





(a) Options and modes

(b) Selected file

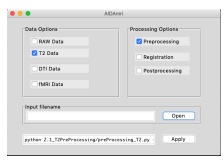
Figure 4: Convert raw data to NIfTI files by specifying the folder containing all raw folders of each scan.

Remember to move the new generated file to a new project folder, if you want to separate the raw Bruker files with the processed NIfTI files. We recommend to build a folder structure as follows: projectfolder/days/groups/subjects/data/especially if you want to use AIDAconnect for graph analysis.

5.2 Processing of T2w & T2mapping data

Apply the reorientation, bias field correction and brain extraction to the T2w data set. The automatically attached endings of the processed filenames indicate which steps have been performed. Brain extraction should be of good quality and must be manually checked or corrected by adapting the default parameter (see Figure 5).

python preProcessing_T2.py -i .../testData/T2w/testData.5.1.nii.gz



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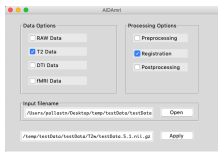
(a) Options and modes

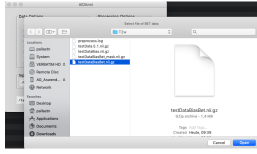
(b) Selected file

Figure 5: Apply the preprocessing to the T2 data by selecting the correct filename and options.

The next step includes the registration of the Allen Brain Reference Atlas with the brain extracted T2 dataset. It is necessary to check the registration result, e.g. by superimposing the brain extracted file with the atlas annotations (ends with ..._Anno.nii.gz) (Figure 6). There is the option to segment an additional region of interest, e.g. the stroke lesion. You can segment the region using the brain extracted dataset as reference (ends with ...BET.nii.gz). We recommend to conduct this step with itk-SNAP. The saved file should end with the extension ...Stroke_mask.nii.gz

python registration_T2.py -i .../testData/T2w/testDataBiasBet.nii





(a) Options and modes

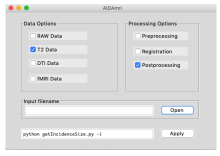
(b) Selected file

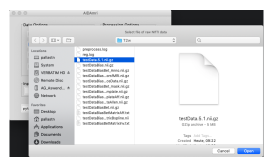
Figure 6: Apply the registration between the T2 data and the ARA by selecting the correct filename and options.

To improve the registration: try to optimize the brain extraction and generated mask (e.g. manually using ImageJ). Then run the registration again.

If you previously defined a region of interest (e.g. the stroke lesion), it is possible to calculate the region size, segmented (parental) atlas regions. Here, the segmented region .../Stroke_mask.nii.gz is overplayed with the Allen Brain Reference Atlas and saved in the file ...Anno_mask.nii.gz. Use the path to the .../T2w as input (Figure 7).

```
python getIncidenceSize_par.py -i .../testData/T2w
python getIncidenceSize.py -i .../testData/T2w
```





(a) Options and modes

(b) Selected file

Figure 7: Get the previously defined region of interest, the region size, segmented parental regions and segmented original regions by selecting the correct filename and options.

The results, such as affected regions and ROI volume are stored in the folder .../T2w in the following files:

```
affectedRegions.txt
affectedRegions.nii.gz
affectedRegions_Parental.txt
affectedRegions_Parental.nii.gz
```

5.3 Processing of T2 data

From the masks drawn on the T2-weighted images, it is possible to determine both the incidence map and the size of affective regions. For example, if a day1 folder contains multiple Mouse_1-Mouse_15 folders and the processed T2 data is in those folders, the command would be as follows

```
python getIncidenceMap.py -i .../day1 -s "Mouse*"
```

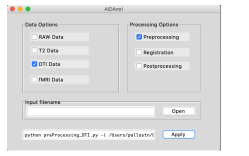
It is also possible to determine the Region size as Voxels and Volume(mm³)

```
python getRegionSize_par.py -i .../T2w
```

5.4 Processing of DTI data

The DTI processing procedure includes a dimension reduction, bias correction, a threshold application, and the subsequent brain extraction. The endings on the filenames indicate which steps have been performed (see Figure 8).

python preProcessing_DTI.py -i .../DTI/testData.7.1.nii.gz



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South lie of ray N

(a) Options and modes

(b) Selected file

Figure 8: Apply the preprocessing to the DTI data by selecting the correct filename and options.

The next step includes the registration of the Allen Brain Reference Atlas with the brain extracted DTI dataset. For processing a reference (stroke) mask, two options are available: a) Registration of a reference mask that is related to another dataset/day, e.g. to always use the same mask, append command -r <filename of ref> b) else, the algorithm will automatically use the corresponding reference mask from the respective subject folder (see Figure 9). If no mask is defined, the registration will proceed without.

python registration_DTI.py -i .../DTI/testDataSmoothMicoBet.nii.gz



(a) Options and modes

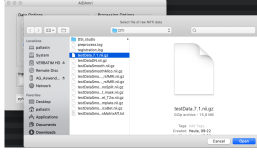
(b) Selected file

Figure 9: Apply the registration between the DTI data and the ARA by selecting the correct filename and options.

The connectivity is finally calculated using DSI-Studio. All connectivity matrices are based on the reference atlas (see Figure 10).

python dsi_main.py -i .../DTI/testData.7.1.nii.gz





(a) Options and modes

(b) Selected file

Figure 10: Process the DTI data with respect to the ARA regions by selecting the correct filename and options.

The connectivity matrices of the parental ARA, the original ARA and the related ROI are stored in the folder .../DTI/connectivity as .txt and .mat. DSI-Studio differentiates between matrices that count how many fibers pass and end in each region. The adjacency matrices can be visualised using the related plot function:

```
python plotDTI_mat.py -i .../testData/DTI/connectivity/testData*
   .connectivity.mat
(without line break!)
```

The folder .../DTI/DSI_studio also contains the diffusion value maps (e.g. FA map) registered with the atlas. This data can be extracted and saved as .txt with the region name and the corresponding FA/RD/MD/AD using the function:

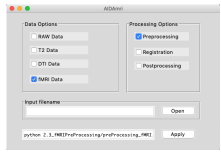
```
python DTIdata_extract.py image_file roi_file
```

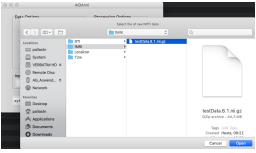
in the 3.2.1_DTIdata_extract folder. To iteratively process all subjects use the iterativeRun.py function.

5.5 Processing of fMRI data

The fMRI processing is roughly comparable to the preprocessing of the DTI datasets. Brain extraction should be of good quality and must be manually checked or corrected by adapting the given parameters (see Figure 11).

python preProcessing_fMRI.py -i .../fMRI/testData.6.1.nii.gz





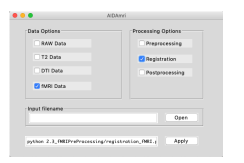
(a) Options and modes

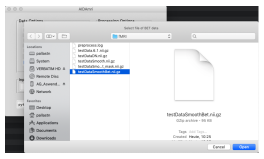
(b) Selected file

Figure 11: Apply the preprocessing to the rsfMRI data by selecting the correct filename and options.

The step includes the registration of the Allen Brain Reference Atlas with the brain extracted fMRI dataset. The result is a variety of files. An impression of the registration can be obtained by superimposing the brain extracted file with the annotations of the Allen Brain (ends with ...Anno.nii.gz) (see Figure 12)

python registration_fMRI.py -i .../testData/fMRI/testSmoothBet.nii





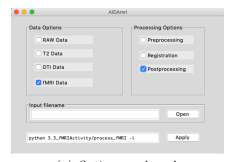
(a) Options and modes

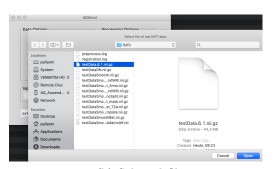
(b) Selected file

Figure 12: Apply the registration between the rsfMRI data and the ARA by selecting the correct filename and options.

If physiological data are not available, the step will be conducted without the included regression. All activity matrices are based on the reference atlas (see Figure 13).

python process_fMRI -i .../fMRI/testData.6.1.nii.gz





(a) Options and modes

(b) Selected file

Figure 13: Process the rsfMRI data with respect to the ARA regions by selecting the correct filename and options.

The activity matrices of the parental Atlas (original Atlas) are stored in the folder .../fMRI/regr as .txt and .mat with the prefix MasksTCs. and MasksTCsSplit.. The related adjacency matrices can be visualised using the related plot function:

python plotfMRI_mat.py -i .../testData/fMRI/regr/MasksTCsSplit*.mat

5.6 Peri-infarct ROI analysis

You can create custom peri-infarct masks to further analyze stroke related regions. Go to the folder bin/4.1_ROI_analysis and open the file proc_tools.py with an arbitrary editor that can open python files. Adjust all directories, paths and further specifications as described in the script. To decide which regions to include in the peri-infarct region, modify:

cortex_labels_1.txt and cortex_labels_2.txt

(for the full list of atlas labels, see ../lib/annoVolume+2000_rsfMRI.nii.txt)

Proceed with the scripts in the order 1 to 4. The first script creates peri-infarct masks for all time points:

python O1_dilate_mask_process.py

The second script aligns the peri-infarct masks in the rsfMRI and DTI space:

python 02_apply_xfm_process.py

The result of the third script depends on the imaging type: In case of rsfMRI, a MATLAB file which contains two text files is being created. 1) for each region one column with the averaged rsfMRI time series and 2) the atlas labels names. In case of DTI, a modified atlas labels file which includes individually shaped peri-infarct brain regions is being created. These new generated regions replace the original regions in the file:

python 03_create_seed_rois_process.py

The fourth script is not mandatory, but a helper tool to compare the number of voxels included in the peri-infarct region for each subject.

python 04_examine_rois.py

Attention: The scripts for peri-infarct ROI analysis are provided for analysis of time point 7 only (e.g. 7 days post stroke). For other time points, manual modifications are necessary.

6 Docker usage

Notice: This guide introduces the usage of Docker-based containers of the AIDAmri tools for Unix-based systems (i.e. Linux and MacOS). Hence, the commands shown are written for such systems and you may copy the commands into your shell including backslashes as they indicate line breaks. A suitable windows user guide will be included soon; for now we advise you to use Unix-based machines or install bash shells such as GIT bash or Anaconda bash to replicate the Unix commands. You may also use the Docker Desktop application to get access to our Docker image.

We provide a local Docker image in our repository. It fully supports every AIDAmri command line interface (CLI) tool. Please note that the AIDAmri GUI is not supported. To use a Docker container based on that image please acquaint yourself with the AIDAmri shell commands.

6.1 Prerequisites

Firstly, install Docker by clicking **here** to get your Docker distribution. We recommend to follow the **getting started tutorial** to get comfortable with the basic Docker usage and terms. For MacOS you might start the Docker Desktop application before starting Docker shell commands.

Clone the AIDAmri git repository to get all files needed to initiate the AIDAmri pipeline:

```
git clone -b dockerdev \
https://github.com/aswendtlab/AIDAmri.git
```

This automatically moves your repository to the dockerdev branch. Please note that the other branches do not contain any files to set-up the Docker.

Quick start

If you wish to quickly build and start the pipeline, a bash runfile (docker_runfile.sh) is provided.

Open a terminal and switch to the AIDAmri directory:

```
cd PATH/TO/AIDAmri
```

There, execute the runfile by typing in the following command:

bash docker_runfile.sh -b -a PATH/TO/DATAFOLDER
or

bash docker_runfile.sh -ba PATH/TO/DATAFOLDER

The -b flag initializes the building process for the Docker image called "aidamri:latest" (see below for further details). The -a flag attaches your shell to the container called "aidamri", so you will be able to use the command line interface (CLI) of the AIDAmri pipeline. If you only like to run the container without attaching to it, do not enter the flags. To attach to the running container, (i.e. the active pipeline) afterwards, use the command:

docker attach aidamri

To enter the running container, use the following command:

docker attach aidamri

You can leave a running container without stopping by pressing CTRL+P and CTRL+Q consecutively. Alternatively, you can type in exit, however this will stop and delete the container, thus you need to re-run the container after stopping it. Keep in mind, that you need to re-run the container after reboot. Use

bash docker_runfile.sh PATH/TO/DATAFOLDER

to re-run the container (building the image is not necessary if no updates were made within the dockerdev git branch). Use the -a flag as mentioned above to directly attach to your pipeline.

When built, run and attached successfully, the shell prompt should look like the following:

root@<SOME NUMBERS AND CHARACTERS>:/aida#

This means that you now are within the AIDAmri environment and can use the tool commands as described in the previous sections.

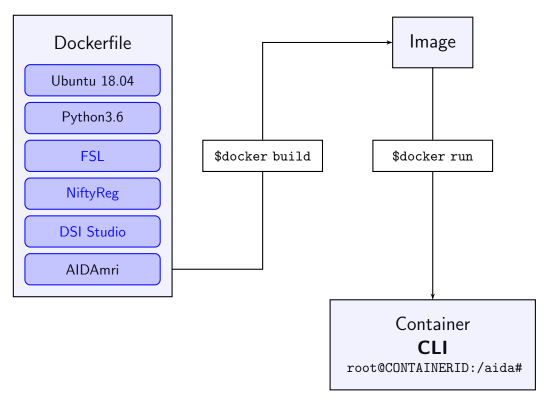


Figure 14: Docker architecture draft. The blue boxes within the Dockerfile box depict the main content layers. The boxes preceded with an \$ are command line codes. Click on texts within the boxes for further information.

6.2 Creating image

To initiate the image building process, open your shell to grant yourself access to the command line interface (CLI). Change your directory to the cloned GitHub repository.

cd PATH/TO/AIDAmri

You can check the folder contents by using the ls command. A file named Dockerfile, as well as the fslinstaller_mod.py, a bin/ and a lib/ folder should be located in this directory. Then, launch the docker daemon to build the image:

```
docker build -t aidamri:latest -f Dockerfile .
```

The created image is a template for running so-called containers (see the below section) and instantiating the pipeline. Be aware that the period at the end is

part of the command and refers to the corresponding directory. The -t flag calls for the name and the tag that you wish to give your image, in this case it is called 'aidamri' and tagged 'latest'. You may change name and tag to your liking, but keep in mind to change them accordingly in later steps that invoke the image. The -f tag calls for the Dockerfile in the current directory. Let the process run. Docker desktop should show you the image within the image tap once it is built.

You only need to run the building process once initially or after updating the github after new changes were made. If an update occurred, the building process will only update those layers that were changed, so the process will not take as long as the initial build.

6.3 Running container and mount data

Before running a container, make sure you know where exactly your data you wish to process is located on your host system. The container will be an instance written from the built image and serves as an environment for you to use the AIDAmri pipeline. To run the container, enter the following command in your command line:

```
docker run -dit \
--rm --name aidamri \
--mount type=bind,source=PATH/TO/DATA,target=/aida/DATA \
aidamri:latest
```

The first flag, <code>-dit</code> starts the container as a detached mode, hence the <code>d</code>, as well as an interactive mode, that allows you to actively use this environment, hence the <code>it</code>. Alternatively, you can directly enter the container environment by only using the <code>-it</code> flag. The <code>--rm</code> flag will delete the container after stopping it. This flag may be optional but it is recommended to use this flag to avoid container housekeeping after you complete the task. With the <code>--name</code> tag you give your container a name, in this case it is called "aidamri". Note, that the image name and the container are named individually. It is recommended to give your container a different name from the image to avoid confusion. Note further that if you wish to use more than one instance of running containers (e.g. to process multiple datasets simultaneously) you need to name each container differently.

Binding mounts create a reference to a given directory, allowing the container to access and transmute data on the host system. In this case, it should allow the pipeline to process your MRI data without copying or re-allocating your data. The --mount flag with the type=bind argument allows you to grant your container access to your target directory. Type in the absolute path at the source placeholder. Do not use relative paths. Within the container environment, a working directory

called aida/ was created. It is recommended to allocate your reference path within this directory by typing in the name of your data folder according to the target placeholder, i.e. /aida/DATA. You can give the target folder another name than the source directory as it still will refer to the source but it is advised to keep the name to avoid confusion. Be aware not to place any spaces between the commas. Pass the name and tag of your image, i.e. here aidamri:latest at the end of the command line. After initializing an ID will appear and the Docker Desktop app should show the running container. You can also check whether the container is running by typing docker container 1s.

To enter the running container, use the following command:

docker attach aidamri

You can leave a running container without stopping by pressing CTRL+P and CTRL+Q consecutively. Alternatively, you can type in exit, however, this will stop and delete the container if the --rm flag was used in the running command, thus you need to re-run the container after stopping it.

When successfully attached, the shell prompt will look like the following:

root@<SOME NUMBERS AND CHARACTERS>:/aida#

The number shows the first part of the container ID. From here, you can use the AIDAmri commands as explained in the former usage sections. As a start, change your directory to the binary folder (cd bin/) and run python batchProg.py -h to see the helping instructions of the batch processing tool and check functionality.