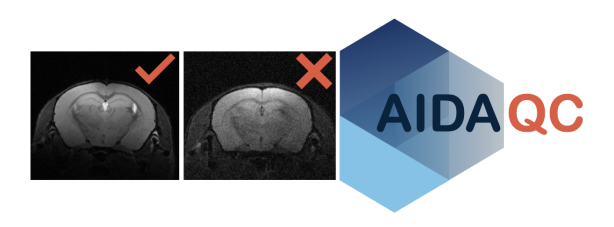
Atlas-based Imaging Data Analysis pipeline for Quality Control of Animal MRI Data

$\begin{array}{c} {\bf AIDAqc} \\ {\bf v2.1} \end{array}$

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1 Introduction

It can be challenging to acquire MR images of consistent quality or to decide in the screening of large databases, which dataset is of sufficient quality for further processing. Manual screening without quantitative criteria is strictly user-dependent and not feasible for huge databases. In contrast to clinical MRI, in preclinical, animal imaging, there is no consensus on standardization of quality control measures or categorization of good vs. bad quality images.

The Atlas-based Processing Pipeline for Quality Control of Animal MRI Data (AIDAqc) was developed for measuring and standardizing the quality of mouse brain MRI in a dynamic and novel way. AIDAqc works with T2-weighted MRI (T2w), diffusion-weighted MRI or diffusion tensor imaging (DTI), and functional MRI (fMRI).

Here, we developed a tool in Python to create a basic overview of MR image datasets including information about the SNR, temporal SNR (tSNR), spatial resolution, and movement severity (Figure 1). Currently, this tool covers T2w, DWI, and fMRI sequences.

I) Parsing:

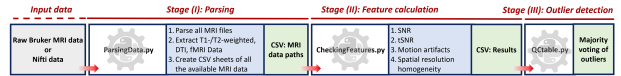
The user sets the input path and the program will parse iteratively through all subfolders with a list of all raw MR data or nifti data as its result. After parsing, only those MR files chosen by the user between the options of T2w, DTI, or fMRI data are selected, and duplicates are eliminated. Finally, CSV files are created with the storage path of every selected file, which will be the input of the next step.

II) Feature calculation:

In this step, SNR is calculated for T2w and DTI images, tSNR, and movement severity for fMRI images. Mutual information was used as a metric to calculate movement severity.

III) Outlier detection:

In this step, all of the calculated features will be statistically analyzed with the help of five methods to identify **outliers**: One class SVM, Isolation Forest, Local Outlier Factor, and Elliptic Envelope. In addition to these, a normal statistical definition of outliers based on the interquartile range is also used.



Python 3.8.12 (Libraries used: NumPy, Pandas, Matplotlib, sklearn, Nibable, Alive_progress, Os, Glob, ...)

Figure 1: Pipeline workflow: All of the necessary functions are implemented in this module, SNR is calculated by using the Chang method and also the more popular way with the help of defining regions of interest inside and outside of the brain (I) In this stage, all of the available MR files are parsed and located. (II) In the main block, here all of the parameters are calculated: SNR, tSNR, and Motion artifacts. Spatial Resolution, slice thickness, and the number of repetitions are also extracted. The final output are CSV files located in a folder called "calculated_features". (III) Five different outlier detectors, each with their own strengths and weaknesses will determine together as a "major vote" what image is considered a bad quality image.

2 Installation

- 1. Download repository by using this link. The project folder contains the python scripts necessary for the quality measurement.
- 2. Download & Install Python 3.6 or higher using Anaconda.
- 3. Importing AIDAqc environment: After the installation of anaconda navigator, we have to import the necessary environment. This can be done by importing the "aidaqc.yaml" file at the *Environments tab*, import and choosing local drive if the file is downloaded from GitHub to a local drive.

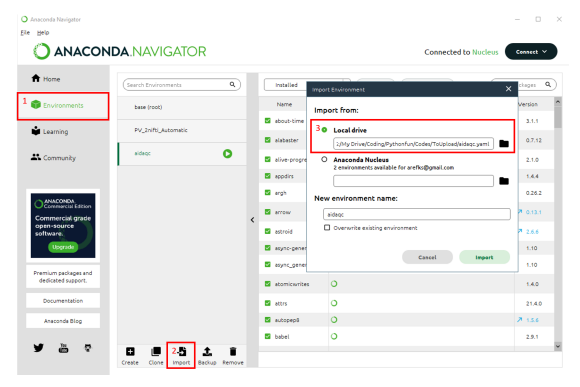


Figure 2: Importing the environment downloaded from GitHub: $Environments/Import/Local\ drive$.

You can also you directly use the anaconda terminal and typing in the folloing commands:

```
cd aidaqc
conda env create --name aidaqc --file=aidaqc.yml
```

3 Scripts

List of important Scripts:

• ParsingData.py: This is the main and only script for the user. Parser for identifying the location of T2w, DTI and fMRI files based on their

sequence name. By using python ParsingData.py -h a short explination and a list of available options will appear. An initial path can be set by the user, the program will use this path as a starting point and search for MR files in every possible subsequent folder after this initial path. The second input is a saving path which is the location where the results should be saved. Figure 7 shows an exemplary use of the author.

- FeatureCheck.py: After ParsingData.py has been used. CSV files with the corresponding addresses are created at the defined location from the user. These CSV files are used as an input for this function. But be aware that this will happen automatically from the Parsing-Data.py script this explination is just for the sake of clarification.
- **QC.py:** In this script, all the necessery functions are gathered together which are used for most parts of the other scripts. *QC.py* can be seen as the tool-box of the whole pipeline (see 3).

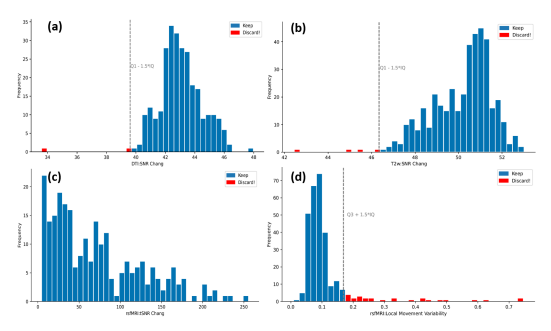


Figure 3: Exemplary statistical plots. The grey vertical line indicates the threshold of good vs bad data based on the statistical definition of "outliers". The bars with the red color indicate those files which should be discarded. (a) Histogram of SNR values of the DTI dataset (b) Histogram of SNR values of the T2w dataset (c) Histogram of tSNR values of the rsfMRI dataset (d) Histogram of the movement variability of the rsfMRI dataset, calculated based on mutual information

Attention: All program examples are only listed with the mandatory input parameters. For more details/help, call python . . . /python <command> -h. After a successful download and install of the necessary libraries, you can start using the pipeline.

4 Workflow

Here we want to show how the pipeline can be used. The steps are explained subsequently.

- 1) Download the sample data set from GIN via this link.
- 2) If not already done, download the repository from this link and follow the installation steps described in the Installation chapter.

```
■ OpenSSH SSH client

(base) kalantaria@ma-macpro-server ~ % cd /Volumes/AG_Aswendt_Projects/Student_projects

/14_Aref_Kalantari_2021/Projects/CRC/QualityControl/Codes/

(base) kalantaria@ma-macpro-server Codes %
```

Figure 4: Changing the terminal's directory to the folder containing the python scripts downloaded from GitHub.

4) activate the aidaqc environment by typing in the following command (figure 5).

conda activate aidaqc

```
■ OpenSSH SSH client

(base) kalantaria@ma-macpro-server ~ % cd /Volumes/AG_Aswendt_Projects/Student_projects ^

/14_Aref_Kalantari_2021/Projects/CRC/QualityControl/Codes/
(base) kalantaria@ma-macpro-server Codes % conda activate aidaqc

(aidaqc) kalantaria@ma-macpro-server Codes %
```

Figure 5: Activating the aidaqc environment. Note that the installation of anaconda and loading the .yaml file is a prerequisite for this step to work (see 3).

5) After activating the environment it is best to check if the environment has been installed correctly and to see if there are any errors accruing in the script. This can be done by using the help option of the function by using the following command. Additionally, a short explanation can also be seen.

python ParsingData.py -h

Figure 6: Using ParsingData.py with the help option.

6) After activating the environment the process can be started by typing in the following command using the script ParsinqData.py.

```
python ParsingData.py -i <!initial_path!> -o <!output_path!> -f nifti -s .1
```

7) After the program has parsed the files and calculated the QC features, one csv file for each available sequence will be created at the defined location set by the user. As can be seen in figure 7 the pipeline informs the user at what stage the pipeline is and how long the processing will approximately take.



Figure 7: Structural overview and summary of the pipeline. 1) Searching for all MR files available 2) Extracting the sequences related to T2w, DTI, and fMRI measurements from the parsed files 3) Duplicate MR files will be only considered once and any file address of copies are eliminated. 4) Final CSV file of stage (I) containing all addresses are created at the defined location. 5) In this part all of the file addresses of part 4 are processed sequence-based. 6) Some files which can contain faulty data or faulty structures with incomplete data won't cause any problems and will also be saved as an Error_data tab in the corresponding CSV file. 7) Same as in 6 faulty fMRI data will be filtered out. 8) Final CSV file in stage (II) containing all of the calculated QC features is saved in this stage. 9) Finally, statistical plots are created and the final results of bad quality data are saved in votings.csv

5 FAQ

Q1) How is the SNR of the T2w and DTI sequences calculated?

The SNR is calculated based on two methods, first one is based on chang method. Simply said this method calculated the SNR without needing to define regions in the image. The second one uses the standard way of calculating SNR namly defining regions of interest inside and outside of the brain. As for this pipeline, the input T2w dataset usually consists of more the one slice of the brain. For example, we have an image set with a dimension of $128 \times 128 \times 20$, the pipeline extracts the 5 best subsequent slices with the highest average value indicating a good image of the brain. Usually, the best slices are the middle ones. If the first or the last slices are the highest slices a warning like in figure 8 will be shown in the terminal and this can indicate a faulty dataset. The reason for this happening can be two things, either the dataset has just one slice which makes no problem and the error can be ignored or the dataset has more slices and it was a faulty measurement where the position of the slice was selected wrongly.



Figure 8: Warning for a dataset in which the first or the last slices have the most signal.

Q2) How is the tSNR calculated?

To calculate the temporal signal to noise ratio multiple approaches are available in the literature, most of them are using normal SNR measurements but add some calculation steps to make it legitimate to be called tSNR. To understand how it is calculated in this pipeline, let's assume a simple example again. Consider an fMRI dataset with

dimensions of $128 \times 128 \times 20 \times 500$ with 20 being the slices and 500 being the temporal time points. Similar to the T2w approach, the 4 best subsequent slices are chosen based on average image intensity. After this step, SNR is calculated based on this tSNR method. Simply said the tSNR is calculated for each voxel over the time, and then it is averaged for a region in the brain. Note that this region definition is all happening automatically based on an automatic threshold and an automatic identification of the the center of mass of the image.

$$tSNR = \frac{\mu}{\sigma} = \frac{\mu}{\sqrt{1/N\sum_{i=1}^{N} (x_i - \mu)^2}}$$
 (1)

This is done for all voxels of the defined volume, grown from the center of mass. Final tSNR is the averaged value of those.

Q3) How is the Movement severity calculated?

Mutual information (MI) was used to calculate the movement severity in the resting state MR measurements. Check out Mutual information as an image matching metric to better understand the concept of Mutual information in image analysis. To explain how MI was used in this pipeline, it's easier to use our example dataset with a dimension of $128 \times 128 \times 20 \times 500$. The question is how much movement has happened between the image at T=t and T=t+1. The four best slices are chosen based on the approach already explained in Q1 and Q2. The image of the first time point is then used as the reference image and all of the following 499 images are compared to the first one by calculating the MI between them. If the MI is high, it means the movement was small. If it's low then the movement was relatively big. The standard deviation of all the MI values is then used as a metric for the overall movement severity. So the final output observable in the CSV file are standard deviation values.

Q4) How does the parsing work in detail?

The parsing technique of this pipeline can be difficult to understand. With the help of figure 9 it gets clearer how the parsing works.

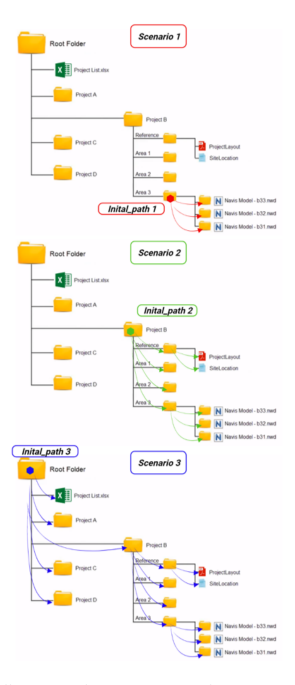


Figure 9: In this illustration, three scenarios can be seen. In each scenario the initial_path as used in figure 6 is set to different folders. The colored arrows is the exact way how the pipeline searches the folders for MR files.

Q5) Can the pipeline process other sequences as well?

For now, the pipeline can only parse T2w, DTI, and fMRI sequences. This is done by using the sequence types extracted from the header information of each measurement. In this pipeline, they are defined as: ['Dti*','EPI','RARE'] which are only relating to the sequence type used for the measurement and the names are predefined default names of Bruker's ParaVision Software. So it might be possible that other kinds of measurements use these sequences as well but are not compatible to be used with this pipeline¹. Another problem that might accrue is that if one of the permitted sequences (T2w, DTI, fMRI) is measured with multiple echo times, repetition times, separate receiver channels, repetitions for averaging purposes, or any kind of additional dimension except Slices, Time and Diffusion directions, the pipeline will not work. It is planned in the near future to add more features.

Q6) How are the SNR of the T2w and DTI sequences calculated? The only difference that we have in the DTI scans compared to the T2w scans, is the dimension of the diffusion directions. Here a small portion of the images in the diffusion direction is used to calculate the SNR similar to the T2w approach.

Q7) Why is the pipeline divided into separate stages?

The reason behind this is to increase the stability of the program and to prevent the need to run the pipeline multiple times from the beginning. As explained above, separate CSV tables are created in each stage. Stage (I) is relatively fast and it won't cause any long waiting periods to rerun that part if necessary. Stage(II) on the other hand can take more time, sometimes up to hours to finish, therefore it is possible that if an error accrues in the plotting part of the code, to simply correct the error and to read the CSV file created from the second stage and to do the plotting without the need to wait again for the whole pipeline to run again from the beginning.

Q9) What is meant by the warning: Some faulty files were found: All faulty files are available in the Errorlist?

With this warning, the pipeline just informs the user that some key files of the main image file could not be read. These can be one of

¹For example Arterial Spin Labeling (ASL) sequences, DCE and DSC sequences, etc.

the method, viso_parameters, ... files from the Bruker sequence folder. However, all these files are also listed and saved in a separate csv file in output folder.

Q9) After running the pipeline, where can I find the data which should be excluded?

In the final CSV sheet named voting.csv are all of the data listed which had at least one vote from the machine-learning outlier detectors ("Judges"). If all five outlier detectors have voted the Image as an outlier, it can be said with a high possibility that it is really bad. And usually based on experience the images with all five votes as an bad/outlier image are pure noise images, or very extreme Ghosting artefacts.

Q10) When running the help option of the function ParsingData.py, there are other options as well, can you elaborate more about them?

The -i and -o are self-explanatory. -f can be used if it should check raw bruker data or nifti data. Be aware that the tool is more stable for only nii data. The reading process of raw data can vary a lot between datasets and is not guaranteed to function for any kind of raw dataset. -s is a useful flag and it will be only used for processing nii data. Imagine you have your main T2.nii.gz file which you want to process with this tool. Also because of some other processing you already have done, there are lots of irrelevant nii files available which you don't want to include in the quality assessment for example T2.mask.nii, T2.procesed.nii, T2Mask.seperated.nii, etc. It might even be the reason of "the error" while using the tool. By using the -s flag you can set it as -s T2, and the tool will only look/parse for files ending with *T2.nii.

Author:

I have designed this pipeline to help me validate all the MR data that I use for further processing in my project. I was searching for a kind of standardization to dichotomize MR data into good and bad data. Over time I found out that this can't be done as easily as thought. The key point of this standardization tool is that one realizes good and bad can not be set with fixed global values of any kind of features like the SNR and as everything in life is, it should be looked at *relatively*. If you have any questions regarding this pipeline, feel free to contact me at:

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The end