MIRACL Documentation

Release 2.2.6

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CHAPTER

ONE

ABOUT

Find important information about MIRACL here. Use the Next button at the bottom of the page or jump to a topic directly by chosing it from the sidebar menu or TOC.

1.1 MIRACL in a nutshell

MIRACL (Multi-modal Image Registration And Connectivity anaLysis) is a general-purpose, open-source pipeline for automated:

- 1. Registration of cleared and imaging data (ex. LSFM and MRI) to atlases (ex. Allen Reference Atlas)
- 2. 3D Segmentation and feature extraction of cleared data
- 3. Tract-specific or network-level connectivity analysis
- 4. Statistical analysis of cleared and imaging data
- 5. Comparison of dMRI/tractography, virus tracing, and connectivity atlases
- 6. Atlas generation and Label manipulation

1.1.1 Program structure

MIRACL is structured into Modules and Workflows.

Modules

The pipeline is comprised of different Modules depending on their respective functionality. Functions for each module are grouped together:

Module	Functionality
connect	Connectivity
conv	Conversion (Input/Output)
reg	Registration
seg	Segmentation
lbls	Labels
utilfn	Utilities
sta	Structure Tensor Analysis
stats	Statistics

An example of using a module would be to run the clar_allen function which performs a CLARITY whole-brain registration to Allen Atlas on a nifti image (down-sampled by a factor of five):

```
$ miracl reg clar_allen -i niftis/SHIELD_05x_down_autoflor_chan.nii.gz -o ARI -m_

→combined -b 1
```

The above command uses the -i flag to select the nifti file, -o to specify the orientation of the image, -m to register to both hemispheres and -b to include the olfactory bulb.

Workflows

The workflow (*flow*) module combines multiple functions from the above modules for ease of use to perform a desired task.

For example, a standard reg/seg analysis could look like this:

First perform registration of whole-brain CLARITY data to ARA:

\$ miracl flow reg_clar -h

Then perform segmentation and feature extraction of full resolution CLARITY data:

\$ miracl flow seg -h

Or structure tensor analysis:

```
$ miracl flow sta -h
```

1.2 Licence

MIRACL is licensed under the terms of the GNU General Public License v3.0.

MIRACL is distributed in the hope that it will be useful, but **WITHOUT ANY WARRANTY**; without even the implied warranty of **MERCHANTABILITY** or **FITNESS FOR A PARTICULAR PURPOSE**. You should have received a copy of GNU General Public License v3.0 along with HippMapp3r.

The code is released for academic research use only. For commercial use, please contact .

1.3 Citing MIRACL

Citation guidelines for using MIRACL in your work.

1.3.1 MIRACL publication

If you use MIRACL in your work please cite our paper:

APA

Goubran, M., Leuze, C., Hsueh, B., Aswendt, M., Ye, L., Tian, Q., Cheng, M.Y., Crow, A., Steinberg, G.K., McNab, J.A., Deisseroth, K., and Zeineh, M. (2019). Multimodal image registration and connectivity analysis for integration of connectomic data from microscopy to MRI. Nature communications, 10(1), 5504.

BibTeX

```
@article{goubran2019multimodal,
    title={Multimodal image registration and connectivity analysis for integration of_
    connectomic data from microscopy to MRI},
    author={Goubran, Maged and Leuze, Christoph and Hsueh, Brian and Aswendt, Markus and_
        -Ye, Li and Tian, Qiyuan and Cheng, Michelle Y and Crow, Ailey and Steinberg, Gary K_
        -and McNab, Jennifer A and Deisseroth, Karl and Zeineh, Michael},
        journal={Nature communications},
        volume={10},
        number={1},
        pages={5504},
        year={2019},
        publisher={Nature Publishing Group UK London}
}
```

1.3.2 Tools used by MIRACL

Some of our functions build on or use these tools (please cite their work if you are using them):

Tool	Function
Allen Regional Atlas	Atlas registration
Allen Connectivity Atlas	Connectivity analyses
ANTs	Registration
Fiji	Segmentation
c3d	Image processing
FSL	Diffusion MRI processing
MRtrix3	Tractography
TrackVis/DTK	Tractography

1.4 MIRACL in research

A selection of publications that used or referenced **MIRACL**:

Hsueh, B., Chen, R., Jo, Y., Tang, D., Raffiee, M., Kim, Y. S., ... Goubran, M., & Deisseroth, K. (2023). Cardiogenic control of affective behavioural state. Nature, 1-8.

Qu, L., Li, Y., Xie, P., Liu, L., Wang, Y., Wu, J., ... & Peng, H. (2022). Cross-modal coherent registration of whole mouse brains. Nature Methods, 19(1), 111-118.

Georgiadis, M., Schroeter, A., Gao, Z., Guizar-Sicairos, M., Liebi, M., Leuze, C., ... & Rudin, M. (2021). Nanostructure-specific X-ray tomography reveals myelin levels, integrity and axon orientations in mouse and human nervous tissue. Nature communications, 12(1), 2941.

Wang, X., Zeng, W., Yang, X., Zhang, Y., Fang, C., Zeng, S., ... & Fei, P. (2021). Bi-channel image registration and deep-learning segmentation (BIRDS) for efficient, versatile 3D mapping of mouse brain. Elife, 10, e63455.

Boillat, M., Hammoudi, P. M., Dogga, S. K., Pages, S., Goubran, M., Rodriguez, I., & Soldati-Favre, D. (2020). Neuroinflammation-associated aspecific manipulation of mouse predator fear by Toxoplasma gondii. Cell reports, 30(2), 320-334.

Pallast, N., Wieters, F., Nill, M., Fink, G. R., & Aswendt, M. (2020). Graph theoretical quantification of white matter reorganization after cortical stroke in mice. NeuroImage, 217, 116873.

Ito, M., Aswendt, M., Lee, A. G., Ishizaka, S., Cao, Z., Wang, E. H., ... & Steinberg, G. K. (2018). RNA-sequencing analysis revealed a distinct motor cortex transcriptome in spontaneously recovered mice after stroke. Stroke, 49(9), 2191-2199.

1.5 Acknowledgements

Huge thank you to:

- Vanessa Sochat (@vsoch) for creating the Docker & Singularity containers for the pipeline
- Newton Cho, Jordan Squair and Stéphane Pagès for helping optimize the segmentation workflows & troubleshooting
- The members of AICONS Lab for their feedback and contributions

GitHub contributers (ordered by number of commits):



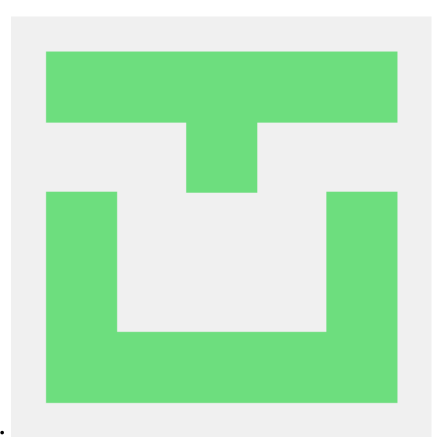
anthonyprinaldi



vsoch



ishita1988



kelvin-jok



chrisroat

PR's:

• dariocalvarez

1.6 AICONS Lab

The Artificial Intelligence and COmputational NeuroSciences (AICONS) Lab is located at the Sunnybrook Research Institute of the University of Toronto and is part of the Black Centre for Brain Resilience and Recovery, Harquail Centre for Neuromodulation, and Temerty Centre for AI Research and Education in Medicine.

Our work combines technical and translational research, focusing on the development of novel AI, computational and imaging tools to probe, predict and understand neuronal and vascular circuit alterations, and model brain pathology in neurological disorders, including *Alzheimer's disease, stroke and traumatic brain injury*.

For more information visit our official webpage.

CHAPTER

INSTALLING AND RUNNING MIRACL

We provide instructions on how to install and run MIRACL using either of the following methods:

Important: Docker is our recommended method for running **MIRACL** on local machines and servers. We recommend **Singularity** to run **MIRACL** in a cluster environment (e.g. Compute Canada).

Attention: Support for installing **MIRACL** locally (i.e. on your host system directly without using **Docker** or **Singularity**) will be phased out in future versions of the software

Docker

Singularity

Local

Windows

We provide a build script to automatically create a **Docker** image for you that can be run using **Docker Compose**. This method does not require a manual installation of **MIRACL** and works on Linux, macOS and Windows (using WSL 2).

Tip: This is our recommended method for running MIRACL on local machines and servers

Docker is well suited if you want to run **MIRACL** on a local machine or local server. If you need to run **MIRACL** on a cluster, see our instructions for installing **Singularity**. If you don't have Docker installed on your computer, do that first. Make sure your installation includes **Docker Compose** as it is required to run the build script we provide. Note that **Docker Compose** is included as part of the **Docker Desktop** installation by default.

First, it is important to understand how the container is built. There is a base image in the docker folder that installs **Python** and dependencies. Then the Dockerfile in the base of the repository builds the mgoubran/miracl image from that base. When the build happens, it cats the version.txt file in the repository to save a versioned base, but then the build uses the tag revised-base-latest that is always the latest base. The base container is built from this folder and pushed manually, while the main container is built and pushed automatically via the **CircleCI** Recipe. Thus, if you want to update the base, you will need to see the README.md in that folder and push new images.

This will build a **Docker** image of **MIRACL** based on its latest version using our default naming scheme. For custom names and specific versions see below for our Additional build options section.

Clone the **MIRACL** repo to your machine:

\$ git clone https://www.github.com/mgoubran/MIRACL
\$ cd MIRACL

Build the latest MIRACL image using the build script we provide:

\$./build.sh

Error: Make sure that the script can be executed. If it can't and you are the owner of the file, use chmod u+x build.sh to make it executable. Prefix with sudo if you are not the owner of the file or change permissions for g and/or o.

Once the image has successfully been built, run the container using **Docker Compose**:

\$ docker compose up -d

Note: Note that the **Docker Compose** syntax is different if you installed it using the standalone method. **Compose standalone** uses the -compose syntax instead of the current standard syntax compose. The above command would thus be docker-compose up -d when using **Compose standalone**.

The container is now running and ready to be used.

Interactively shell inside:

```
$ docker exec -it miracl bash
```

Files that are saved while using **MIRACL** should be saved to volumes mounted into the container in order to make them persistent. To mount volumes, just add them to the docker-compose.yml in the base directory under volumes.

Danger: Do not delete the volume that is already mounted which mounts your **.Xauthority**! This is important for X11 to work correctly.

Example:

```
volumes:
    - '/home/mgoubran/.Xauthority:/home/mgoubran/.Xauthority'
    - '/home/mgoubran/mydata:/home/mgoubran/mydata'
```

Exit your container and navigate to your MIRACL folder. Use Docker Compose to stop the container:

\$ docker compose down

Note: Note that the **Docker Compose** syntax is different if you installed it using the standalone method. **Compose standalone** uses the -compose syntax instead of the current standard syntax compose. The above command would thus be docker-compose up -d when using **Compose standalone**.

Naming is done automatically when using our build script which includes a default naming scheme. By default, the image is named mgoubran/miracl:latest and the container is tagged with miracl.

You can easily change the defaults if your usecase requires it by running our build script with the following options:

\$./build -i <image_name> -c <container_name>

Options:

-i, Specify image name (default: mgroubran/miracl)
-c, Specify container name (default: miracl)

Example:

\$./build -i josmann/miracl -c miracl_dev_version

Tip: Use ./build -h to show additional options

By default, **Docker** images will be built using the latest version of **MIRACL**. If you need to build a **Docker** image based on a specific version of **MIRACL**, do the following:

1. Clone the MIRACL repository and navigate to the MIRACL folder:

\$ git clone https://www.github.com/mgoubran/MIRACL
\$ cd MIRACL

2. Cloning the repository will download all tags/versions. List them with:

\$ git tag -1

Example output:

v1.1.1 v2.2.1 v2.2.2 v2.2.3 v2.2.4 v2.2.5

3. Decide which tag/version of MIRACL you want to use and check it out as a new branch:

\$ git checkout tags/<tag_name> -b <branch_name>

Example:

\$ git checkout tags/v2.2.4 -b miracl_v2.2.4

4. If you are reverting to a version of MIRACL >= 2.2.4, you can build the image for your chosen version by running the build script with the -t flag:

\$./build.sh -t

Note: If you want to build an image for a version of MIRACL <= 2.2.4 either follow the build instructions of the particular version or download the latest build script using e.g. wget https://raw.githubusercontent.com/AICONSlab/MIRACL/master/build.sh (overwrites current build script if present) and run it with the -t flag.

5. From here you can follow our instructions for building **MIRACL** from scratch starting with docker compose up -d. Our script will automatically detect the version of the branch you checked out and tag the image accordingly.

Unlike **Docker**, **Singularity** is well suited to run in a cluster environment (like Sherlock at Stanford or Compute Canada). We provide the latest version of **MIRACL** as a **Singularity** container that can be conveniently pulled from cloud storage.

Tip: This is our recommended method for running **MIRACL** in a SLURM cluster environment such as Compute Canada or Sherlock @ Stanford

First, log in to the cluster:

\$ ssh -Y <username>@<cluster>

<cluster> could be sherlock.stanford.edu or cedar.computecanada.ca for example

Once logged in, change the directory to your scratch space and pull (download) the **Singularity** container:

\$ cd \$SCRATCH \$ singularity pull miracl_latest.sif library://aiconslab/miracl/miracl:latest

Attention: singularity pull requires **Singularity** version 3.0.0 or higher. Please refer to our *Troubleshoot-ing section* ("Can I build a Singularity container from the latest MIRACL image on Docker Hub") if you are using an older version of **Singularity**.

To shell into the container use:

```
$ singularity shell miracl_latest.sif bash
```

Use the -B flag to bind a data directory to the container:

\$ singularity shell -B /data:/data miracl_latest.sif bash

See also:

For running functions on clusters please check our Singularity tutorials for Compute Canada and Sherlock

Warning: Support for this installation method will be discontinued in future versions of **MIRACL**. We recommend to use **Docker** or **Singularity** instead.

Steps to setup/run MIRACL on a Linux/macOS machine:

\$ git clone https://github.com/mgoubran/MIRACL.git miracl

Tip: Alternatively, you can download the zip file containg the repo and uncompress it

Next, change directories into the newly created miracl folder:

\$ cd miracl

Create your virtual MIRACL environment and activate it:

Attention: To setup a virtual environment you need **Anaconda** for **Python 2.7**. It can be downloaded from their official website

\$ conda create --name miracl python=3.7.4 pip
\$ conda activate miracl

Install dependencies:

\$ pip install -e .

Next, download the depends folder from our Dropbox link and place it either inside the linux_depends or mac_depends folder:

```
$ mv ~/Downloads/depends.zip miracl/.
$ cd miracl
$ unzip depends.zip
$ rm depends.zip
```

This folder contains compiled versions of **ANTS** and **c3d** for Linux or Mac OS. Before continuing, make sure to change the permissions.

This can be done by running:

```
$ chmod -R 755 <path/to/depends>/*
```

In order to run the pipeline, some symbolic links must be added to access certain commands. Inside the miracl folder, run:

Make sure <path/to/depends> is replaced with the directory path that leads to the depends directory.

Place the atlases folder (which got downloaded together with the depends folder) inside the miracl folder:

```
$ mv ~/Downloads/atlases.zip miracl/.
$ cd miracl
$ unzip atlases.zip
$ rm atlases.zip
```

This folder contains the Allen Atlas data needed for registration and connectivity analysis.

First, download **Fiji/ImageJ** from their offical website.

Then do:

```
$ cd depends
$ wget https://downloads.imagej.net/fiji/latest/fiji-linux64.zip
$ unzip fiji-linux64.zip
$ rm fiji-linux64.zip
```

Next, install additional plugins by going to Help -> Update and clicking on the Manage update sites button.

Choose the following update sites:

- 3D ImageJ Suite: http://sites.imagej.net/Tboudier
- Biomedgroup: https://sites.imagej.net/Biomedgroup
- IJPB-plugins: http://sites.imagej.net/IJPB-plugins

Download FSL and install it:

```
$ wget https://fsl.fmrib.ox.ac.uk/fsldownloads/fslinstaller.py
$ sudo python fslinstaller.py
```

For the visualization of nifti files and labels we recommend ITKSNAP or the nifti plugin for Fiji/ImageJ.

If you have diffusion MRI data download and install MRtrix3:

```
$ sudo apt-get install git g++ python python-numpy libeigen3-dev zlib1g-dev libqt4-

opengl-dev libg11-mesa-dev libfftw3-dev libtiff5-dev
$ git clone https://github.com/MRtrix3/mrtrix3.git
$ cd mrtrix3
$ ./configure
$ ./build
$ ./set_path
```

To end a **MIRACL** session, deactivate your virtual environment:

\$ conda deactivate

To update MIRACL, navigate into your MIRACL base folder (e.g. \$ cd miracl) and run:

\$ git pull

You should be good to go!

Warning: Support for installing **MIRACL** locally in the WSL will be discontinued in future versions of **MIRACL**. We recommend to use **Docker** or **Singularity** instead.

To install **MIRACL** on your Windows system, Windows Subsystem for Linux (WSL) must be installed. WSL2 is preferred. From there, the usual steps to install **MIRACL** on a Linux based system will be used with a few tweaks.

Hint: Follow the below steps if you want to install **MIRACL** in your WSL instance locally. If you prefer to use **Docker** to run **MIRACL** on Windows follow our installation instructions for **Docker** instead.

The Windows Subsystem for Linux (WSL) creates an environment that allows users to run versions of Linux without having to set up a virtual machine or a different computer.

To install WSL, users can follow the instructions from Microsoft. More comprehensive instructions can be found here. Upgrading from WSL 1 to WSL 2 is recommended, due to WSL 2's benefits.

Note: You may ignore this step if you have a preferred Linux distribution that is already installed in your WSL2

A Linux distribution (distro), like Ubuntu, is an operating system based on the Linux kernel.

Now that WSL (either 1 or 2) is installed, the **Ubuntu 22.04** distro can be installed. To install Ubuntu, open the Windows Store app, search for "**Ubuntu 22.04**", and select the Get button. You could also use this link.

The Ubuntu distro should have **Python 3** installed. To ensure that this is the case, update all packages installed in the WSL:

```
$ sudo apt update
$ sudo apt -y upgrade
```

We can see which version of **Python 3** is installed by typing:

```
$ python3 -V
```

The output in the terminal window will show the version number.

pip is required to install software packages in Python. It can be installed by running the following command:

\$ sudo apt install -y python3-pip

You could also use Anaconda to install the packages but we found that installing and using pip was more straightforward.

To actually install MIRACL, follow the local installation instructions for Linux and macOS.

To use **MIRACL's** graphical user interface (GUI), **Xming** must be installed. **Xming** is a display server for Windows computers, that is available for use by anyone. It can be downloaded from SourceForge.

Before running **MIRACL's** GUI, run Xming. In the terminal window where **MIRACL's** GUI will be run, input the following command:

```
$ export DISPLAY=$DISPLAY:localhost:0
```

Now that everything is installed, MIRACL can be run via the WSL. To run:

- 1. Open WSL via terminal
- 2. Navigate to the folder where you would like to run MIRACL from
- 3. Activate the environment containing miracl:

\$ source activate miracl
\$ miraclGUI

An accompanying Jupyter notebook for this tutorial can be found here.

CHAPTER

THREE

TUTORIALS

The tutorial structure generally matches the module/function structure of **MIRACL**. Refer to the *Legend section* for syntax information.

Choose the tutorial for the function you are interested in from its respective module in the sidebar/TOC or go to our *Getting started section* for a tutorial on **MIRACL's** general usage.

Note that not all functions have tutorials yet...we are working on it!!!

3.1 Legend

In the docs/tutorials, code examples are written as follows:

```
$ miracl -h
```

Code blocks look like this:

```
usage: miracl [-h] {connect,conv,flow,lbls,reg,seg,sta,stats,utils} ....
positional arguments:
  {connect,conv,flow,lbls,reg,seg,sta,stats,utils}
    connect
                        connectivity functions
    conv
                        conversion functions
                        workflows to run
    flow
   lbls
                        label manipulation functions
                        registration functions
   reg
                        segmentation functions
   seq
   sta
                        structure tensor analysis functions
    stats
                        statistical functions
   utils
                        utility functions
optional arguments:
  -h, --help
                        show this help message and exit
```

Inline code is marked as: \$ miracl -h

Admonitions are displayed as colored text boxes. This is an example of what a 'tip' admonition would look like:

Tip: The above -h flag can be used with each of MIRACL's modules/functions

We use brackets to denote text as follows:

- {}: Used for variabels.
 - Example: niftis/downsample{factor}x.nii.gz
- <>: Used for placeholder text in examples that you need to replace with your own information.
 - Example: \$ ssh <username>@cedar.computecanada.ca
- []: Placeholders for flag arguments used in command-line scripting.
 - Example: \$ miracl flow sta -f [Tiff folder] -o [output nifti]
- []: Denotes flags in the command-line help menus.
 - Example: \$ miracl [-h]

Files and directories (or generally paths) are denoted like this: example_dir/example_file.nii.gz Names of executable programs are marked as follows: **MIRACL**

Lastly, links are highlighted in blue (purple when clicked): link to MIRACL's README

3.2 Getting started

MIRACL can be run in the command-line or by using its GUI.

3.2.1 Command-line

Note: If you have installed MIRACL locally, run source activate miracl first to start its virtual environment.

To look at available modules, invoke the help menu:

```
$ miracl -h
```

The following menu should be printed to the terminal:

```
usage: miracl [-h] {connect,conv,flow,lbls,reg,seg,sta,utils} ...
positional arguments:
  {connect,conv,flow,lbls,reg,seg,sta,utils}
   connect
                        connect functions
    conv
                        conv functions
                        workflows to run
    flow
   lbls
                        Label manipulation functions
   reg
                        registration functions
                        segmentation functions
   seg
                        STA functions
    sta
                        STA functions
    sta
   utils
                        Utils functions
optional arguments:
                        show this help message and exit
  -h, --help
```

If you want information about a particular module you can call it with -h. Let's use the **conv** module as an example. Invoke its help menu using:

\$ miracl conv -h

You should get:

```
usage: miracl conv [-h] {tiff_nii,nii_tiff,set_orient,gui_opts} ...
positional arguments:
    {tiff_nii,nii_tiff,set_orient,gui_opts}
    tiff_nii convert Tiff stacks to Nii
    nii_tiff convert Nii volume to Tiff stack
    set_orient Set orientation tag with GUI
    gui_opts GUI options
optional arguments:
    -h, --help show this help message and exit
```

For accessing the help menu of a specific function in the conv module, say tiff_nii, type:

\$ miracl conv tiff_nii -h

You should get:

```
usage: Converts Tiff images to Nifti
A GUI will open to choose your:
    - < Input CLARITY TIFF dir >
_ _ _ _ _ _ _ _ _ _ _ _
For command-line / scripting
Usage: miracl conv tiff_nii -f [Tiff folder]
Example: miracl conv tiff_nii -f my_tifs -o stroke2 -cn 1 -cp C00 -ch Thy1YFP -vx 2.5 -
→vz 5
required arguments:
  -f dir, --folder dir Input CLARITY TIFF folder/dir
optional arguments:
  -d , --down
                         Down-sample ratio (default: 5)
  -cn , --channum
                         Chan # for extracting single channel from multiple channel data.
\leftrightarrow (default: 0)
                         Chan prefix (string before channel number in file name). ex: COO
  -cp , --chanprefix
  -ch , --channame
                         Output chan name (default: eyfp)
  -o , --outnii
                         Output nii name (script will append downsample ratio & channel
\rightarrow info to given name)
 -vx , --resx
                         Original resolution in x-y plane in um (default: 5)
  -vz , --resz
                         Original thickness (z-axis resolution / spacing between slices)
\rightarrow in um (default: 5)
 -c [ ...], --center [ ...] Nii center (default: 0,0,0) corresponding to Allen atlas
→nii template
```

(continues on next page)

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-dz ,downzdim	Down-sample in z dimension, binary argument, (default: 1) => yes
-pd ,prevdown	Previous down-sample ratio, if already downs-sampled
-h,help	Show this help message and exit

To run the function with an input dir called input_tiff_dir, a down-sampling factor of 5 and an output called test, you would type:

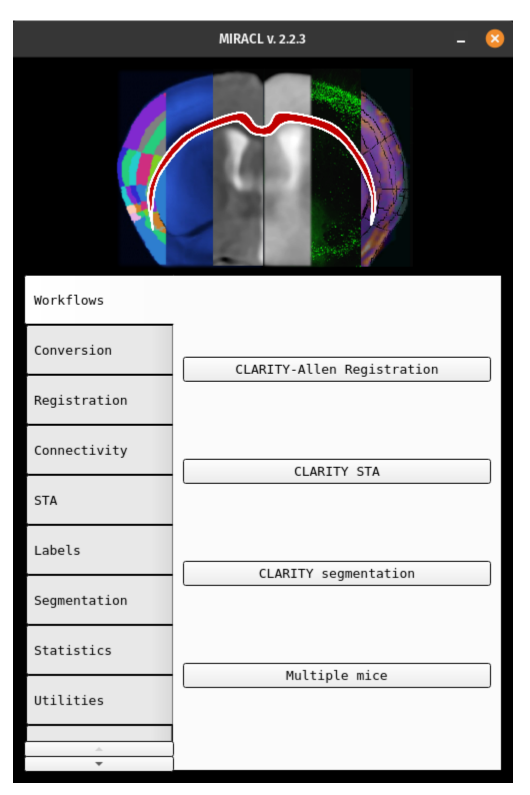
\$ miracl conv tiff_nii -f input_tiff_dir -d 5 -o test

3.2.2 GUI

To run the main GUI:

\$ miraclGUI

The GUI should open:



To get the GUI of a specific function, run it without arguments, for example:

\$ miracl conv tiff_nii

Not all functions have GUIs yet... we are working on it!!!

See also:

Check the rest of the tutorials for more detailed documentation on modules and functions

3.3 Workflows

The workflow (flow) module combines multiple functions from various modules for ease of use to perform a desired task.

For example, a standard reg/seg analysis could look like this:

First perform registration of whole-brain CLARITY data to ARA:

\$ miracl flow reg_clar -h

Then perform segmentation and feature extraction of full resolution CLARITY data:

\$ miracl flow seg -h

Or structure tensor analysis:

\$ miracl flow sta -h

Note that not all functions have tutorials yet...we are working on it!!!

3.3.1 ACE Workflow

AI-based Cartography of Ensembles (ACE) pipeline highlights:

- 1. Cutting-edge vision transformer and CNN-based DL architectures trained on very large LSFM datasets (link to sample data and *refer to example section*) to map brain-wide neuronal activity.
- 2. Optimized cluster-wise statistical analysis with a threshold-free enhancement approach to chart subpopulationspecific effects at the laminar and local level, without restricting the analysis to atlas-defined regions (link to sample data and *refer to example section*).
- 3. Modules for providing DL model uncertainty estimates and fine-tuning.
- 4. Interface with MIRACL registration to create study-specific atlases.
- 5. Ability to account for covariates at the cluster level and map the connectivity between clusters of activations.

Main Inputs

Control and Treated directories, containing whole-brain 3D LSFM datasets for multiple subjects. OR A single directory containing a single subject's whole-brain 3D LSFM dataset.

CLI

To get more information about the workflow and its required arguments use the following command on the cli:

\$ miracl flow ace -h

The following information will be printed to the terminal:

```
usage: miracl ace (-s SINGLE_TIFF_DIR |
                  (-c CONTROL_BASE_DIR CONTROL_TIFF_DIR_EXAMPLE -e EXPERIMENT_BASE_DIR_
→ EXPERIMENT_TIFF_DIR_EXAMPLE))
                  -sao SA_OUTPUT_FOLDER -sam {unet,unetr,ensemble}
                  (--overwrite | --no-overwrite)
  1) Segments images with ACE
  2) Registers tissue cleared data (down-sampled nifti images) to Allen Reference mouse
→brain atlas
  3) Voxelizes high-resolution segmentation maps to downsample into Allen atlas.
→resolution
  4) Warps voxelied segmentation maps from native space to Allen atlas
  5) Generates group-wise heatmaps of cell density using the average of voxelized and
→warped segmentation maps in each group
 6) Computes group-level statistics/correlation using cluster-wise analysis on_
→voxelized and warped segmentation maps
Single or multi method arguments:
-s SINGLE_TIFF_DIR, --single SINGLE_TIFF_DIR
                        path to single raw tif/tiff data folder
-c CONTROL_BASE_DIR CONTROL_TIFF_DIR_EXAMPLE, --control CONTROL_BASE_DIR CONTROL_TIFF_
→DIR_EXAMPLE
                        FIRST: path to base control directory. SECOND: example
                        path to control subject tiff directory
-e EXPERIMENT_BASE_DIR EXPERIMENT_TIFF_DIR_EXAMPLE, --experiment EXPERIMENT_BASE_DIR_
→ EXPERIMENT_TIFF_DIR_EXAMPLE
                        FIRST: path to base experiment directory. SECOND:
                        example path to experiment subject tiff directory
                      overwrite existing output files for comparison
--overwrite
                        workflow
                      do not overwrite existing output files for comparison
--no-overwrite
                        workflow. This flag can be used to run only the stats
                        analysis (if the subject-only steps have already been
                        run).
required arguments:
-sao SA_OUTPUT_FOLDER, --sa_output_folder SA_OUTPUT_FOLDER
                        path to output file folder
-sam {unet,unetr,ensemble}, --sa_model_type {unet,unetr,ensemble}
                        model architecture
utility arguments:
-ua U_ATLAS_DIR, --u_atlas_dir U_ATLAS_DIR
                        path of atlas directory (default:
                        '/code/atlases/ara/')
```

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Use -hv **or** --help_verbose flag **for** more verbose help **and** view other ACE modules arguments

Note: There are a number of optional arguments including TFCE cluster-wise analysis parameters that can be provided to the respective function invoked by the workflow. These arguments have been ommitted here for readability but can be viewed by running miracl flow ace -hv.

Flag	Parameter	Туре	Description
-s,single	SINGLE_TIFF_DIR	str	path to raw tif/tiff data folder
-c,control	CONTROL_BASE_DIR, CON- TROL_TIFF_DIR_EXAMPLE	(str, str)	path to base control directory, example path to control subject tiff directory
-e, experiment	EXPERIMENT_BASE_DIR, EX- PERIMENT_TIFF_DIR_EXAMPLE	(str, str)	path to base experiment directory, example path to experiment subject tiff directory
-sam, sa_model_type	{unet,unetr,ensemble}	str	model architecture
overwrite no-overwrite		bool	whether to overwrite existing output files for comparison workflow

Main outputs

clar_allen_reg # registration output / pre-liminary files conv_final # conversion (tiff to nifti) output reg_final # main registration output seg_final # segmentation output including model(s) outputs and uncertainty estimates vox_final warp_final heatmap_final cluster_final # cluster-wise analysis output including p_value and f_stats maps corr_final # correlation analysis output including correlation maps and p_value maps

Executes:

```
seg/ace_interface.py
conv/miracl_conv_convertTIFFtoNII.py
reg/miracl_reg_clar_allen.sh
seg/miracl_seg_voxelize_parallel.py
reg/miracl_reg_warp_clar_data_to_allen.sh
stats/miracl_stats_heatmap_group.py
stats/miracl_stats_ace_interface.py
```

Example of running ACE on single subject (segmenation + registration + voxelization + warping) (link to sample data):

```
$ miracl flow ace \
    -s ./non_walking/Newton_HC1/cells/ \
    -sao ./output_dir \
    -sam unet \
    -overwrite
```

Example of running ACE flow on multiple subjects:

```
$ miracl flow ace \
    -c ./non_walking/ ./non_walking/Newton_HC1/cells/ \
    -e ./walking/ ./walking/Newton_UI1/cells/ \
    -sao ./output_dir \
    -sam unet \
    -overwrite
```

Example of running only ACE segmentation module on one single subject (link to sample data):

```
$ miracl seg ace \
   -sai ./Ex_561_Em_600_stitched/ \
   -sao ./output_dir \
   -sam unetr
```

Example of running only ACE cluster wise analysis on voxelized and warped segmentation maps (link to sample data):

```
$ miracl stats ace \
    -c ./ctrl/ \
    -e ./treated/ \
    -sao ./output_dir \
```

Jupyter notebook

An accompanying Jupyter notebook for this tutorial can be found *here*.

3.3.2 CLARITY whole-brain registration to Allen Atlas

The registration workflow relies on an autofluorescence channel input (tiff files), and can perform whole-brain or hemisphere registrations to the Allen Atlas.

This workflow performs the following tasks:

- 1. Sets orientation of input data using a GUI
- 2. Converts TIFF to NII
- 3. Registers CLARITY data (down-sampled images) to Allen Reference mouse brain Atlas

- 4. Warps Allen annotations to the original high-res CLARITY space
- 5. Warps the higher-resolution CLARITY to Allen space (if chosen)
- 6. Test data
- 7. A test dataset (CLARITY autofluorescence channel) for registration is found here under data: https://www. dropbox.com/sh/i9swdedx7bsz1s8/AABpDmmN1uqPz6qpBLYLtt8va

Main outputs

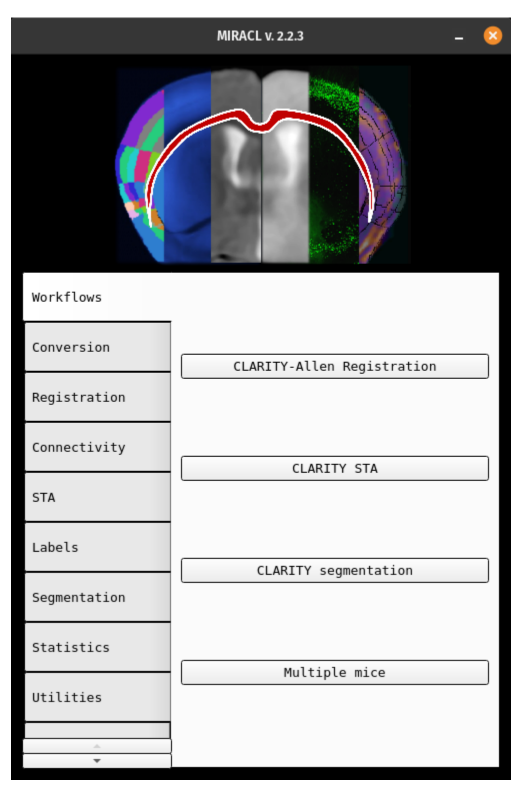
File	Description		
<pre>reg_final/clar_allen_space.nii.gz</pre>	CLARITY data in Allen reference space		
<pre>reg_final/clar_downsample_res{vox}um.nii.gz</pre>	CLARITY data downsampled and oriented to 'standard'		
<pre>reg_final/annotation_hemi_{hemi}_{vox}um_clar_down nii.gz</pre>	Allen labels registered to downsampled CLAR-ITY		
<pre>reg_final/annotation_hemi_{hemi}_{vox}um_clar_vox. tif</pre>	Allen labels registered to oriented CLARITY		
<pre>reg_final/annotation_hemi_{hemi}_{vox}um_clar. tif</pre>	Allen labels registered to original (full-resolution) CLARITY		

GUI

Run:

\$ miraclGUI

and choose CLARITY-Allen Registration from the Workflows tab:



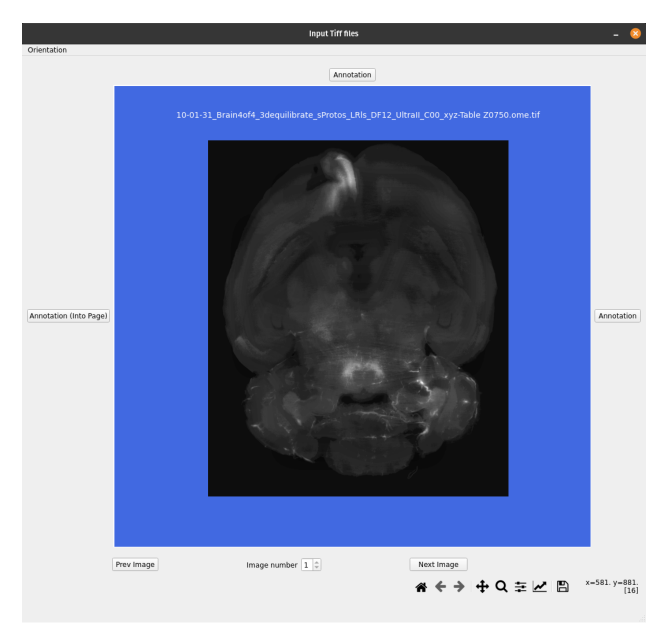
Or run:

\$ miracl flow reg_clar

Choose the input tiff folder with the auto fluorescence channel from the pop-up menu:

Look in:	/home/josmann/data/prtSPIM/2020111	L9_Brain0	- 0 0) 🖓 🕼 🗉 🗏
Computer josmann	Ex 488 Em 0 Stitched	Size	Type Folder Folder	Date Modified 8 Feb 26:48:08 8 Feb 23:11:36
Directory:				<u>Choose</u>
Files of type:	Directories			 Cancel

The following GUI will appear which opens the data and lets you set its orientation manually:



You can navigate through the data using the bar bellow, by specifying the slice number or using the arrows:

Prev Image	Image number 1 🛊 🚽	Next Image
		☆ < → ⊕ Q ≑ 🗹 🖺

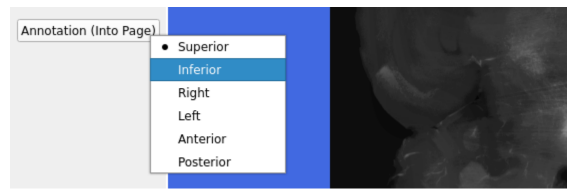
First choose the data plane (axial, coronal or sagittal):

Orientation	
Axial	
Sagittal	
Coronal	
	10-01-31_Brain4of4_

Then choose the orientation at the top and right of the image:

	Annotation	
		 Superior Inferior
		Right
		Left
and the second		Anterior
· · · ·		Posterior
and the second se		

Next, choose the orientation for scrolling through the slices (going into the page), can confirm the orientation by changing the image number at the bottom (enter higher number and press Enter), or using the Next or Prev image buttons:



Finally close the GUI:



Next, set the tiff conversion parameters:

Nii conversion options 🗕 🙁				
No Dir selected	Select output directory (def = working dir)			
out nii (def = clarity)				
downsample ratio (def = 5)				
channel #				
channel prefix				
channel name (def = eyfp)				
in-plane res (def = 5 um)				
z res (def = 5 um)				
center (def = $0 \ 0 \ 0$)				
Help	Enter			
	Run			

Conversion parameters description:

Parameter	Description Default	
out nii	Output nifti name	clarity
downsample ratio	Downsample factor for conversion	5
channel #	Number for extracting single chan- nel from multiple channel data (leave blank if single channel data/tiff files)	0
channel prefix	<pre>String before channel number in file name (leave blank if single chan- nel). For example, if tiff file name has _C001tif for channel 1 and _C002tif for channel 2, to choose channel 1 if it's the auto flu- orescence channel:</pre>	Channel prefix not invoked if not provided
channel name	Output channel name	eyfp
in-plane res	Original resolution in x-y plane in um	5
z res	Thickness (z-axis resolution/spacing between slices) in um	5
center	Center of nifti file	0 0 0

Next, choose the registration options:

Reg options – 😣			
No file selected	Select input custom atlas [blank (def)]		
No file selected	Select input custom labels [blank (def)]		
Hemi [combined (def)/split]			
Labels resolution [vox] (def = 10 'um')			
olfactory bulb incl. (def = 0)			
side [blank (def) / rh / lh]			
Help	Enter		
	Run		

Registration parameters description:

Parameter	Description	Default
Hemi	Warp Allen labels with hemisphere split (Left labels are different from Right labels) or combined (Left and Right lables are the same i.e. mir- rored). Accepted inputs are: • combined • split	combined
Labels resolution [vox]	Voxel size/resolution of labels in um accepted inputs are: 10, 25 or 50	10
olfactory bulb	If olfactory bulb is in- cluded in the dataset. Accepted inputs are: 0 (not included) 1 (included)	0
Side	Only if registering hemisphere, else leave blank. Accepted inputs are: rh (right hemisphere) lh (left hemi- sphere)	None

Command-line

Usage:

\$ miracl flow reg_clar -f [Tiff folder]

Example:

\$ miracl flow reg_clar -f my_tifs -n "-d 5 -ch autofluo" -r "-o ARS -m combined -v 25"

Arguments (required):

f. Input Clarity tif dir/folder

Optional arguments (remember the quotes):

```
Conversion to nii (invoked by -n " "):
d. [ Downsample ratio (default: 5) ]
cn. [ chan # for extracting single channel from multiple channel data (default: 0) ]
cp. [ chan prefix (string before channel number in file name). ex: C00 ]
ch. [ output chan name (default: eyfp) ]
vx. [ original resolution in x-y plane in um (default: 5) ]
vz. [ original thickness (z-axis resolution / spacing between slices) in um (default: 5).
-→]
c. [ nii center (default: 5.7 -6.6 -4) corresponding to Allen atlas nii template ]
Registration (invoked by -r " "):
o. Orient code (default: ALS)
   to orient nifti from original orientation to "standard/Allen" orientation
m. Warp allen labels with hemisphere split (Left different than Right labels) or.
→combined (L & R same labels / Mirrored)
   accepted inputs are: <split> or <combined> (default: combined)
v. Labels voxel size/Resolution of labels in um
   accepted inputs are: 10, 25 or 50 (default: 10)
1. image of input Allen Labels to warp (default: annotation_hemi_split_10um.nii.gz -_
\rightarrow which are at a resolution of 0.01mm/10um)
   input could be at a different depth than default labels
   If 1. is specified (m & v cannot be specified)
```

Visualize results

Run:

\$ miracl reg check

Usage:

Example:

\$ miracl reg check -f reg_final -v itk -s clarity

Arguments:

Required:

-f, Input final registration folder

Optional:

-v, Visualization software: itkSNAP 'itk' (default) or freeview 'free'
-s, Registration Space of results: clarity (default) or allen

3.3.3 STA workflow

Run the structural tensor analysis (STA) workflow for fiber quantification and tracking.

Attention: Run workflow after running the CLARITY-Allen registration first

Workflow for STA:

- 1. Converts Tiff stack to nii incl. down-sampling
- 2. Uses registered labels to create a seed mask at the depth (ontology level) of the desired label (seed)
- 3. Creates a brain mask
- 4. Runs STA analysis using the seed and brain masks
- 5. Computes virus intensities for all labels at that depth

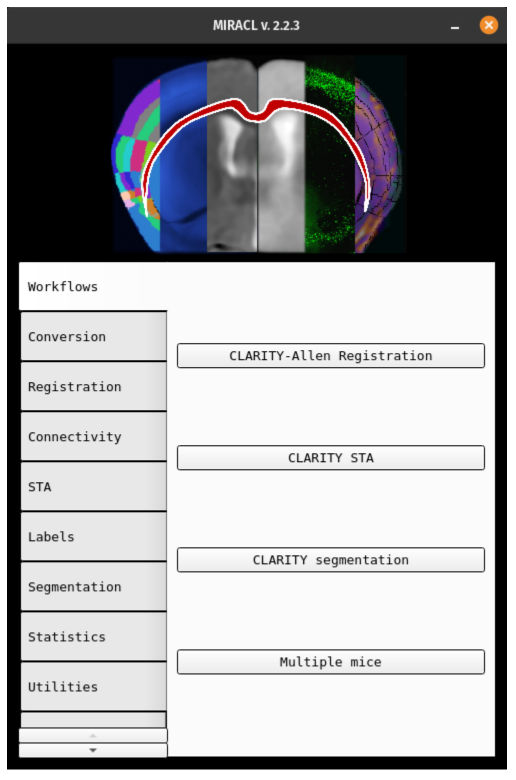
Executes:

```
conv/miracl_conv_convertTIFFtoNII.py
lbls/miracl_lbls_get_graph_info.py
lbls/miracl_lbls_generate_parents_at_depth.py
utils/miracl_extract_lbl.py
utils/miracl_create_brainmask.py
sta/miracl_sta_track_primary_eigen.py
lbls/miracl_lbls_stats.py
```

Main Outputs

File	Description
<pre>clarity_sta_{label}_seed/dog{dog}_gauss{gauss}/ filter_ang{angle}.trk</pre>	Tract file
<pre>virus_signal_stats_depth_{depth}.csv</pre>	Virus stats csv
<pre>sta_streamlines_density_stats_depth_{depth}.csv</pre>	Streamline density stats csv

GUI



From the main GUI (invoked with: \$ miraclGUI), select Workflows -> CLARITY STA:

The following window will appear:

STA workflow	• •
No file selected	Select Brain mask (optional)
No file selected	Select Seeding label mask (optional)
No Dir selected	Select Input tiff folder
No Dir selected	Select CLARITY final registration folder
Out nii name (def = clarity)	
Seed label abbreviation	
hemi (combined or split)	
Derivative of Gaussian (dog) sigma	
Gaussian smoothing sigma	
Tracking angle threshold	
Downsample ratio (def = 5)	
chan # (def = 1)	
chan prefix	
Out chan name (def = AAV)	
Resolution (x,y) (def = 5 um)	
Thickness (z) (def = 5 um)	
Downsample in z (def = 1)	
Dilation factor for x axis (def = 0)	
Dilation factor for y axis (def = 0)	
Dilation factor for z axis (def = 0)	
Step length (def 0.1)	
Use 2nd order range-kutta method for tracking (def 0)	
Output directory	
Help	Enter
	Run

Hint: To open the STA workflow menu directly use: \$ miracl flow sta

Click on Select Input tiff folder and choose the folder that contains the virus channel from the dialog window.

Then choose the registered Allen labels inside the final registration folder (reg_final) from the dialog window by clicking on Select CLARITY final registration folder.

Next choose the output file name (Output nii name), e.g. Mouse05. Our script will automatically append down-sample ratio and channel info to the given name.

Set the tracking parameters:

Parameter	Description	Default
Seed label abbreviation	From Allen atlas ontology, for the seed region. Examples: Combined hemispheres: • CP for Caudoputamen • PL for Prelimbic Area Right hemisphere: • RCP for Right Caudoputamen • RPL for Right Prelimbic Area	Required. Function will exit with error 1 if not provided.
hemi	Labels hemisphere. Accepted inputs are: • combined`(both) • split (left or right)	combined
Derivative of Gaussian (dog) sigma	Example: 1	0.5,1.5
Gaussian smoothing sigma	Example: 0.5	0.5,2
Tracking angle threshold	Example: 35	25,35
Use 2nd order runge-kutta method for tracking	Use 2nd order runge-kutta: • 0 (don't use) • 1 (use)	0

And the tiff conversion parameters:

Parameter	Description	Default
Downsample ratio	Set the downsample ratio.	5
chan #	For extracting single channel from multiple channel data.	1
chan prefix	String before channel number in file name. Example:	C00
Resolution (x,y)	Original resolution in x-y plane in um.	5
Thickness	Original thickness (z-axis resolu- tion/spacing between slices) in um.	5
Downsample in z	Downsample in z dimension. Bi- nary: • 0 (no) • 1 (yes)	1

Users can also input their own brain mask, as well as their own seeding mask. Both masks would respectively replace the automatically generated brain mask and regional mask used for the tractography. Users also have the option to dilate the seed mask across any of the three dimensions, by a value (indicated by the Dilation factor fields).

Attention: Note that the following parameters are required:

- tiff folder
- output nii name
- Seed label abbreviation
- CLARITY final registration folder
- hemi
- Derivative of Gaussian
- Gaussian smoothing sigma
- Tracking angle threshold

After choosing the parameters, first press Enter to save them and then Run to start the workflow.

Command-line

Usage:

```
$ miracl flow sta -f [ Tiff folder ] -o [ output nifti ] -l [ Allen seed label ] -m [_

→hemisphere ] -r [Reg final dir] -d [ downsample ratio ]
```

Example:

```
$ miracl flow sta -f my_tifs -o clarity_virus -l PL -m combined -r clar_reg_final -d 5 -
→c AAV g 0.5 -k 0.5 -a 25
```

Or for right PL:

```
$ miracl flow sta -f my_tifs -o clarity_virus -l RPL -m split -r clar_reg_final -d 5 -c_

→AAV -g 0.5 -k 0.5 -a 25
```

Arguments:

```
arguments (required):
  -f: Input Clarity tif folder/dir (folder name without spaces)
  -o: Output nifti
  -1: Seed label abbreviation (from Allen atlas ontology)
  -r: CLARITY final registration folder
  -m: Labels hemi
  -g: Derivative of Gaussian (dog) sigma
  -k: Gaussian smoothing sigma
  -a: Tracking angle threshold
optional arguments:
  -d: Downsample ratio (default: 5)
  -c: Output channel name
  -n: Chan number for extracting single channel from multiple channel data (default: 0)
  -p: Chan prefix (string before channel number in file name). ex: C00
  -x: Original resolution in x-y plane in um (default: 5)
  -z: Original thickness (z-axis resolution/spacing between slices) in um (default: 5)
```

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```
-b: Brain mask (to replace brain mask automatically generated by workflow)
-u: Seed mask (in place of regional seed mask generated by workflow)
-s: Step length
-downz: Downsample in z
-dilationfx: Dilation factor for x (factor to dilate seed label by)
-dilationfy: Dilation factor for y (factor to dilate seed label by)
-dilationfz: Dilation factor for z (factor to dilate seed label by)
-rk: Use 2nd order range-kutta method for tracking (default: 0)
-out_dir: Output directory
```

Jupyter notebook

An accompanying Jupyter notebook for this tutorial can be found here.

3.3.4 CLARITY whole-brain segmentation

There are multiple segmentation functions for different data (stains/channels):

- virus
- cFos
- sparse
- nuclear

The segmentation workflow relies on an output from the registration workflow, but the segmentation wrapper function can be run without running the registration workflow.

This workflow performs the following tasks:

- 1. Segments neurons in cleared mouse brain of sparse or nuclear stains in 3D
- 2. Voxelizes segmentation results into density maps with Allen Atlas resolution
- 3. Computes features of segmented image and summarizes them per label

It executes:

```
seg/miracl_seg_clarity_neurons_wrapper.sh
seg/miracl_seg_voxelize_parallel.py
seg/miracl_seg_feat_extract.py
```

Main outputs

File	Description
<pre>segmentation/seg.{tif,mhd} or seg_nuclear.{tif,mhd} segmentation/seg_bin.{tif,mhd} or seg_bin_nuclear. {tif,mhd}</pre>	Segmentation image with all labels (cells) Binarized segmentation image
<pre>voxelized_seg.{tif,nii}</pre>	Segmentation results voxelized to ARA reso- lution
<pre>voxelized_seg_bin.{tif,nii}</pre>	Binarized version
<pre>clarity_segmentation_features_ara_labels.csv</pre>	Segmentation features summarized per ARA labels

Hint: Results can be opened in Fiji for visualization

GUI

Select from the main GUI menu (invoked from the cli: \$ miraclGUI) or run:

\$ miracl flow seg

The following window will appear:

	Seg options 🛛 🗕 🙁
No file selected	Select Registered allen labels (clar_vox.tif) in the reg_final dir
No Dir selected	Select Input tiff dir
seg type [virus (def), cfos, sparse or nuclear]	
channel prefix (ex = C001)	
labels voxel size	
Help	Enter
	Run

Click on Select registered labels (...clar_vox.tif) in the reg_final dir to choose the registered labels annotation_hemi_{side}_**um_clar_vox.tif to summarize segmentation features where:

- {side} -> combined or split
- ** is the resolution -> 10, 25 or 50

The following window will appear:

Look in:	/home/josmann/data/projec4/downsample_03	/reg_final 🔻	00	0	A	::	
Computer	allen labels to clar mosaic.png	f 3.88 GiB 24GiB z 35MiB	tif File gz File gz File tif File tif File gz File gz File tif File	Date 7 Oct 6 Oct 6 Oct 7 Oct 7 Oct 7 Oct 7 Oct 6 Oct	29 21 20 21 29 28 29 28):10:: .:03: .:54: .:06: .:57: .:02:: .:02:: .:52:: .:05:: .:05:: .:55::	17 12 15 11 52 53 36 58
File <u>n</u> ame: Files of type:	annotation_hemi_combined_10um_clar_vox.tif All Files (*)					<u>O</u> pe Canc	

Next, click on select input tiff dir to select folder with Thy1-YFP or other channel:

Look in:	🚞 /home/josmann/data/prtSPIM/202011	19_Brain0	- 0 0	0 🕼 🗉 🗏
Computer josmann	Ex 488 Em 0 Stitched	Size	Type Folder Folder	Date Modified 8 Feb 26:48:08 8 Feb 23:11:36
Directory:				
Files of type:	Directories			 Cancel

Lastly set the segmentation parameters:

Parameter	Description	Default
seg type	Channel type: • virus • cFos • sparse (like Thy1 YFP) • nuclear (like PI)	virus
channel prefix	Channel prefix and number if multiple channels. Example: Filter0001	None
labels voxel size	Registered labels voxel size in um: • 10 • 25 • 50	10

Click Enter and Run to start the segmentation process.

Command-line

Usage:

```
$ miracl flow seg -f [ Tiff_folder ]
```

Example:

Arguments:

```
arguments (required):
f. Input Clarity tif folder/dir [folder name without spaces]
t. Channel type: sparse (like Thy1 YFP) or nuclear (like PI)
optional arguments (don't forget the quotes):
Segmentation (invoked by -s " "):
p. Channel prefix & number if multiple channels (like Filter0001)
Feature extraction (invoked by -e " "):
1. Allen labels (registered to clarity) used to summarize features
    reg_final/annotation_hemi_{hemi}_{vox}um_clar_vox.tif
```

3.4 Conversion

Functions to convert between image file formats e.g. converting tiff image files to nifti format.

Note that not all functions have tutorials yet...we are working on it!!!

3.4.1 Tiff to Nifti

Function to convert Tiff images to Nifti format for analysis or visualization.

GUI

Run:

\$ miracl conv tiff_nii

The following window will open:

Tiff to Nii conversion	- 😣
No Dir selected	Select Input tiff folder
Output dir (def = working dir)	
Out nii name (def = clarity)	
Downsample ratio (def = 5)	
chan # (def = 1)	
chan prefix	
Out chan name (def = eyfp)	
Resolution (x,y) (def = 5 "um")	
Thickness (z) (def = 5 "um")	
center (def = 0,0,0)	
Downsample in z (def = 1)	
Prev Downsampling (def = 1 -> not downsampled)	
Help	Enter
	Run

Click on Select Input tiff folder to choose the input tiff folder:

Look in:	/home/josmann/data/prtSPIM/2020111	19_Brain0	- 0 (0 🕼 🗉 🗏
Computer	Ex 488 Em 0 Stitched	Size	Type Folder Folder	Date Modified 8 Feb 26:48:08 8 Feb 23:11:36
Directory:				<u>C</u>hoose
Files of type:	Directories			 Cancel

Next set the desired parameters or use the default by leaving the fields blank:

Parameters	Description	Default	
Out nii name	Output file name.	clarity	
Downsample ratio	Downsample factor for nifti images.	5	
chan # and prefix	Use if tiff files have more than one channel. For ex- ample, given the names	Not invoked if not provided	-Tabl
	10-04-06_R923_06R1ca_647_100p Z1284.ome.tif for channel 1 and		
	10-04-06_R923_06R1ca_647_100p Z1284.ome.tif for channel 2 with the latter being the desired channel for conversion, chan # would be 2 and chan prefix would be C00.		-Table
Out chan name	Output channel name.	eyfp	
Resolution (x,y)	Original resolution in x-y plane in um.	5	
Thickness	Original thickness (z-axis resolu- tion/spacing between slices) in um.	5	
center	Center of Nifti file.	0 0 0	
Downsample in z	Down-sample in z dimension. Bi- nary: • 0 (no) • 1 (yes)	1	
Prev Downsampling	 Previous downsample ratio if already downsampled. Accepted inputs are: 0 (downsampled) 1 (not downsampled) 	1	

After choosing the parameters press Enter to save them then Run to start the conversion process.

Tip: After the conversion is done, nifti (nii/nii.gz) files can be visualized using the ITKsnap software

Command-line

Usage:

```
$ miracl conv tiff_nii -f [ Tiff_folder ]
```

Example:

```
miracl conv tiff_nii -f my_tifs -o stroke2 -cn 1 -cp COO -ch Thy1YFP -vx 2.5 -vz 5
```

Required arguments:

-f dir, --folder dir Input CLARITY TIFF folder/dir

Optional arguments:

```
-d, --down
                      Down-sample ratio (default: 5)
-cn, --channum
                      Chan # for extracting single channel from multiple channel data_
\leftrightarrow (default: 1)
                      Chan prefix (string before channel number in file name). ex: COO
-cp, --chanprefix
                      Output chan name (default: eyfp)
-ch, --channame
-o, --outnii
                      Output nii name (script will append downsample ratio & channel info.
\rightarrowto given name)
                      Original resolution in x-y plane in um (default: 5)
-vx, --resx
-vz, --resz
                      Original thickness (z-axis resolution / spacing between slices) in_
\rightarrowum (default: 5)
-c [ ...], --center [ ...]
                      Nii center (default: 0,0,0 ) corresponding to Allen atlas nii
→template
-dz, --downzdim
                     Down-sample in z dimension, binary argument, (default: 1) => yes
-pd, --prevdown
                     Previous down-sample ratio, if already down-sampled
-h, --help
                      Show this help message and exit
```

3.5 Registration

Registration functions for e.g. registering data (down-sampled images) to Allen Reference mouse brain atlas. Note that not all functions have tutorials yet...we are working on it!!!

3.5.1 CLARITY-Allen registration

This function will do the following:

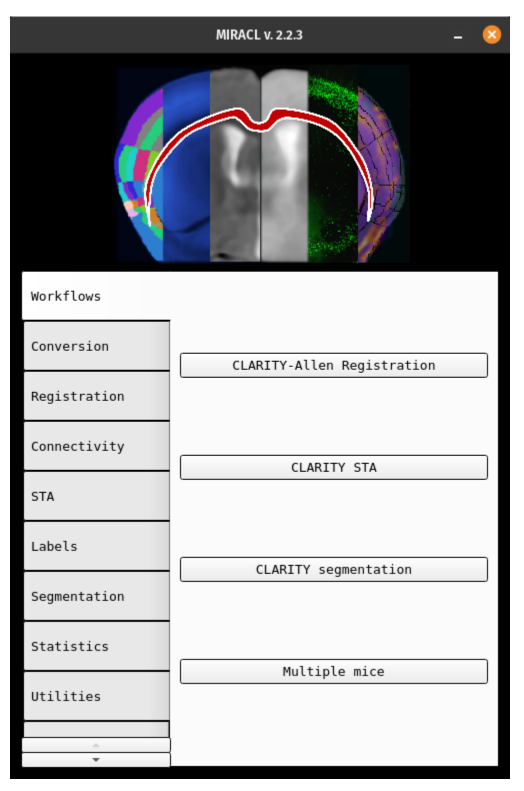
- 1. Registers CLARITY data (down-sampled images) to Allen Reference mouse brain atlas
- 2. Warps Allen annotations to the original high-res CLARITY space
- 3. Warps the higher-resolution CLARITY to Allen space

GUI

To open the main registration menu, open MIRACL's main menu first by running:

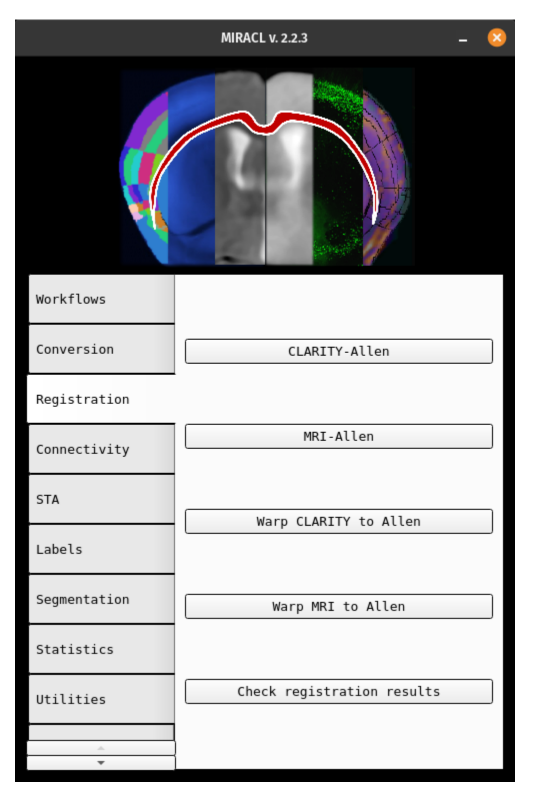
\$ miraclGUI

MIRACL's main menu will open:



Select the Registration tab on the left for the main registration menu.

The main registration window will look like this:



From here you can select CLARITY-Allen to start the registration. The Reg options menu will open:

	Reg options 🛛 🗕 😣
No file selected	Select Down-sampled auto-fluorescence (or Thy1) channel
Output directory (def = working dir)	
Orient code (def = ASL)	
Labels Hemi [combined (def)/split]	
Labels resolution [vox] (def = 10 'um')	
olfactory bulb incl. (def = 0)	
side (def = None)	
extra int correct (def = 0)	
Help	Enter
	Run

Tip: To open the above Reg options menu directly, run: \$ miracl reg clar_allen

The registration will be run on downsampled CLARITY Nii images. You can provide the folder containing these files in the first field. **This parameter is required to run the registration**. You can use **MIRACL's** conversion methods to create the downsampled files if you do not have them yet.

Flag	Parameter	Description	Default	
-i	Input down-sampled CLARITY Nii	Preferably auto- fluorescence channel data (or Thy1_EYFP if no auto chan). File name should have **x_down like 05x_down (meaning 5x downsampled). Example: • combined • split	Required. Script exits with error 1 if not pro- vided.	

All remaining parameters are optional. If left blank, their respective default values will be used:

Flag	Parameter	Description	Default
-1	Output directory	Directory the output (re- sults) will be written to.	<working_directory>/ reg_final</working_directory>
-0	Orient code	Code to orient nifti from original orientation to 'standard/Allen' orienta- tion.	ALS
-m	Labels hemi	Chose to register to one or both hemispheres. Warps Allen labels with hemi- sphere split (L differ from R labels) or combined (L and R have the same la- bels i.e. are mirroed). Ac- cepted inputs are: • combined • split	combined
-V	Labels resolution [vox]	Labels voxel size/resolution in um. Accepted inputs are: • 10 • 25 • 50	10
-b	Olfactory bulb included	Specify whether the olfac- tory bulb is included in brain. Accepted inputs are: • Ø (not included) • 1 (included)	0
-S	Side	 Provide this parameter if you are only registering one hemisphere instead of the whole brain. Accepted inputs are: rh (right hemisphere) 1h (left hemisphere) 	None
-p	Extra int correct	If utilfn intensity correc- tion has already been run, skip correction inside reg- istration. Accepted inputs are: • Ø (don't skip) • 1 (skip)	0

After providing the parameters click Enter to save them and Run to start the registration process.

Once the registration is done the final files will be located in the output folder (default: <working_directory>/

reg_final). Files created in intermediate steps will be located in a folder called <working_directory>/ clar_allen_reg.

Command-line

The command-line version has additional functionality that is not included in the GUI version:

Attention: Note that the above listed -i parameter (input down-sampled CLARITY Nii) is also required for the command-line

Usage:

Example:

```
$ miracl reg clar_allen -i downsampled_niftis/SHIELD_03x_down_autoflor_chan.nii.gz -o_

ARI -m combined -b 1
```

Jupyter notebook

An accompanying Jupyter notebook for this tutorial can be found *here*.

3.5.2 MRI whole-brain registration to Allen Atlas

This registration method performs the following tasks:

- 1. Registers in-vivo or ex-vivo MRI data to Allen Reference mouse brain Atlas
- 2. Warps Allen annotations to the MRI space

GUI

Invoke with \$ miraclGUI and select from main menu or run:

\$ miracl reg mri_allen_nifty

The following window will open:

Reg options 🛛 🗕 📀					
No file selected	Select In-vivo or Ex-vivo MRI				
Orient code (def = RSP)					
Labels Hemi [combined (def)/split]					
Labels resolution [vox] (def = 10 'um')					
olfactory bulb incl. (def = 0)					
skull strip (def = 1)					
No orient (def = 0)					
Help	Enter				
	Run				

Click on Select In-vivo or Ex-vivo MRI and choose the input MRI nii (preferable T2-w) using the dialog window. Then set the registration options:

Parameter	Description	Default
Orient code	Orient nifti from original orientation to 'standard/Allen' orientation.	RSP
Labels Hemi	Warp allen labels with hemisphere split (Left different than Right la- bels) or combined (Left and Right labels are the same/mirrored). Ac- cepted inputs are: • split • combined	combined
Labels resolution [vox]	Labels voxel size/resolution in um. Accepted inputs are: • 10 • 25 • 50	10
Olfactory bulb included	Specify whether the olfactory bulb is included in brain. Accepted inputs are: • 0 (not included) • 1 (included)	0
skull strip	 Strip skull. Accepted inputs are: 0 (don't strip) 1 (strip) 	1
No orient	No orientation needed (input image in 'standard' orientation). Accepted inputs are: • 0 (orient) • 1 (don't orient)	0

Click Enter and Run to start the registration process.

Command-line

Usage:

Example:

```
$ miracl reg mri_allen_nifty -i inv_mri.nii.gz -o RSP -m combined -v 25
```

Arguments:

arguments (required):
i. input MRI nii
Preferably T2-weighted
optional arguments:
o. orient code (default: RSP)
to orient nifti from original orientation to "standard/Allen" orientation
<pre>m. hemisphere mirror (default: combined)</pre>
warp allen labels with hemisphere split (Left different than Right labels) or
\hookrightarrow combined (L & R same labels / Mirrored)
accepted inputs are: <split> or <combined></combined></split>
v. labels voxel size/Resolution in um (default: 10)
accepted inputs are: 10, 25 or 50
 input Allen labels to warp (default: annotation_hemi_combined_10um_nii.gz)
input labels could be at a different depth than default labels
If l. is specified (m & v cannot be specified)
b. olfactory bulb included in brain, binary option (default: \emptyset -> not included)
s. skull strip or not , binary option (default: 1 -> skull-strip)
f. FSL skull striping fractional intensity (default: 0.3), smaller values give larger
→brain outlines
n. No orientation needed (input image in "standard" orientation), binary option
→(default: 0 -> orient)

3.6 Stats

This module consists of multiple sub-modules to apply statistical and correlation analysis to the data.

Note that not all functions have tutorials yet...we are working on it!!!

3.6.1 ACE Cluster Only

Apply cluster-wise analysis including a cluster-wise TFCE test and correlation analysis on voxelized and warped segmentation maps.

Main Inputs

Control and Treated directories, containing voxelized and warped segmentation maps for each group.

CLI

To get more information about the workflow and its required arguments use the following command on the CLI:

```
$ miracl stats ace -h
```

Example usage (link to sample data):

```
$ miracl stats ace \
    -c ./ctrl/ \
    -e ./treated/ \
    -sao ./output_dir \
    -n 1000 \
    -a ./atlas/ \
    -r 25 \
    -sfwhm 3 \
    -start 0.05 \
    -step 5 \
    -h 2 \
    -e 0.5
```

Flag	Parameter	Туре	Description	Default	t
-c,control	CON- TROL_BASE_DIR	str	path to base control directory	None quired)	(re-
-e,experiment	EXPERI- MENT_BASE_DIR	str	path to base experiment directory	None quired)	(re-
-sao, sa_output_folder	SA_OUTPUT_FOLDE	str	path to output directory	None quired)	(re-
-n,num_perm	NUM_PERM	int	number of permutations	100	
-a,atlas_dir	ATLAS_DIR	str	path to atlas directory	miracl	_home
-r,img_resolution	IMG_RESOLUTION	int	image resolution (atlas resolution 10 or 25 um)	25	
-sfwhm, smoothing_fwhm	SMOOTH- ING_FWHM	int	fwhm of Gaussian kernel in pixel	3	
-start,tfce_start	TFCE_START	float	tfce threshold start	0.01	
-step,tfce_step	TFCE_STEP	float	tfce threshold step	10	
-h,tfce_h	TFCE_H	float	tfce H power	2	
-e,tfce_e	TFCE_E	float	tfce E power	0.5	
-c,cpu_load	CPU_LOAD	float	percent of cpus used for parallelization	0.9	
-p,step_down_p	STEP_DOWN_P	float	step down p value	0.3	
-m,mask_thr	MASK_THR	int	percentile to be used for binarizing differ- ence of the mean	95	

Jupyter notebook

An accompanying Jupyter notebook for this tutorial can be found *here*.

3.7 Segmentation

Functions to segment tiff image files.

Note that not all functions have tutorials yet...we are working on it!!!

3.7.1 ACE Segmentation Function

Cutting edge vision transformer and CNN-based DL architectures trained on very large LSFM datasets to map cFos brain-wide.

CLI

To look at the arguments that need to be provided to the function, invoke the help menu using:

\$ miracl seg ace -h

The following menu will be printed to the terminal:

```
usage: miracl ace [-h] -sai SA_INPUT_FOLDER -sao SA_OUTPUT_FOLDER -sam
                  {unet,unetr,ensemble} [-sas height width depth]
                  [-sar X-res Y-res Z-res] [-saw SA_NR_WORKERS]
                  [-sac SA_CACHE_RATE] [-sasw SA_SW_BATCH_SIZE] [-samc] [-sav]
                  [-sau]
AI-based Cartography of Ensembles (ACE) segmentation method
optional arguments:
  -h, --help
                        show this help message and exit
  -sai SA_INPUT_FOLDER, --sa_input_folder SA_INPUT_FOLDER
                        path to raw tif/tiff data folder
  -sao SA_OUTPUT_FOLDER, --sa_output_folder SA_OUTPUT_FOLDER
                        path to output file folder
  -sam {unet,unetr,ensemble}, --sa_model_type {unet,unetr,ensemble}
                        model architecture
  -sas height width depth, --sa_image_size height width depth
                        image size (type: int; default: fetched from image
                        header)
  -sar X-res Y-res Z-res, --sa_resolution X-res Y-res Z-res
                        voxel size (type: _validate_vox_res)
  -saw SA_NR_WORKERS, --sa_nr_workers SA_NR_WORKERS
                        number of cpu cores deployed to pre-process image
                        patches in parallel (type: int; default: 4)
  -sac SA_CACHE_RATE, --sa_cache_rate SA_CACHE_RATE
                        percentage of raw data that is loaded into cpu during
                        segmentation (type: float; default: 0.0)
  -sasw SA_SW_BATCH_SIZE, --sa_sw_batch_size SA_SW_BATCH_SIZE
```

(continues on next page)

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```
number of image patches being processed by the model
in parallel on gpu (type: int; default: 4)
-samc, --sa_monte_dropout
use Monte Carlo dropout (default: False)
-sav, --sa_visualize_results
visualizing model output after predictions (default:
False)
-sau, --sa_uncertainty_map
enable map (default: False)
```

Flag	Parameter	Туре	Description	Default
-sai, sa_input_folder	SA_INPUT_FOI	str	path to raw tif/tiff data folder	None (required)
-sao, sa_output_folder	SA_OUTPUT_F	str	path to output file folder	None (required)
-sam, sa_model_type	{unet,unetr,enser	str	model architecture	None (required)
-sas, sa_image_size	height width depth	int	image size; provided as three arguments	fetched from image header
	X-res Y-res Z- res	int	voxel resolution; provided as three arguments	None (required)
-saw, sa_nr_workers	SA_NR_WORKI	int	number of cpu cores deployed to pre-process image patches in parallel	4
-sac, sa_cache_rate	SA_CACHE_RA	floa [.]	percentage of raw data that is loaded into cpu during segmentation	0.0
-sasw, sa_sw_batch_size	SA_SW_BATCH	int	number of image patches being processed by the model in parallel on gpu	4
-samc, sa_monte_dropout	True/False	bool	use Monte Carlo dropout	False
-sav, sa_visualize_resul	True/False	bool	visualizing model output after predictions	False
-sau, sa_uncertainty_ma	True/False	bool	enable map	False

Note: The -sa in the flag part stands for segmentation ACE.

Example usage:

```
$ miracl seg ace \
   -sai ./walking/subject_01/cells/ \
   -sao ./output_dir \
   -sam unet
```

3.8 Utilities

A collection of utility functions.

Note that not all functions have tutorials yet...we are working on it!!!

3.8.1 Intensity correction

Intensity correction for data with inhomogeneity issues. Performs correction on CLARITY tiff data in parallel using N4.

- 1. Creates a downsampled nifti from the tiff data
- 2. Runs N4 'bias field'/intensity correction on the nifti file
- 3. Up-samples the output bias field and applies it to the tiff data

Command-line

Usage:

Example:

```
$ miracl utils int_corr -f tiff_folder -od bias_corr_folder
```

Required arguments:

Flags	Description	Default Default	
-f, –folder	Input CLARITY TIFF folder/dir	No default. Input folder must be provided by user.	

Optional arguments:

Flags	Description	Default
-od,outdir	Output folder name	int_corr_tiffs
-cn, channum	Chan # for extracting single channel from multiple channel	data 1
-cp, chanprefix	Chan prefix (string before channel number in file name). Example: C00	None
-ch, channame	Output chan name	AAV
-on,outnii	Output nii name (script will append downsample ratio & channel info to given name)	
-vx,resx	Original resolution in x-y plane in	um 5
-vz,resz	Original thickness (z-axis resolution/spacing between slices) in um	5
-m,maskimg	Mask images before	correction 1
-s,segment	Perform level-set seg using brain mask to get a dilated	one 0
-d,down	Downsample/shrink factor to run bias corr on downsampled	data 5
-n,noise	Noise parameter for histogram sharpening - deconvolution	0.005
-b,bins	Histogram bins	200
-k,fwhm	FWHM for histogram sharpening - deconvolution	0.3
-l,levels	Number of levels for	convergence 4
-it,iters	Number of iterations per level for convergence	50
-t,thresh	Threshold per iteration for	convergence Ø
-p,mulpower	Use the bias field raised to a power of p to enhance its effects	1.0

3.9 HPC/SLURM clusters

MIRACL was built with HPC/SLURM clusters in mind. We recommend **Singularity** as it is well suited to run in a cluster environment. We provide a **Singularity** container of **MIRACL's** latest version that can be pulled to a node directly from our online repo.

We provide tutorials on how to use **MIRACL** on Compute Canada and Sherlock (supercomputer at Stanford university) but the principles explained here will be similar to other SLURM clusters.

Note: If you would like to add a tutorial for a particular cluster that you are working with that is missing here, we invite you to add it to this section by submitting a PR (note that we write Sphinx documentation in .rst format) through our official GitHub.

3.9.1 Running MIRACL on Compute Canada

This tutorial highlights the registration workflow but a similar approach applies to other commands.

When using Compute Canada, **MIRACL** can be used as a **Singularity** container. The following instructions are based on the steps provided on the Compute Canada Wiki.

Copy your data to Compute Canada

For example, to copy a folder called input_clar containing tiff files you want to register to the Allen Atlas use:

\$ scp -r input_clar niagara.computecanada.edu:/scratch/<username>/.

or

\$ rsync -avPhz input_clar <username>@niagara.computecanada.edu:/scratch/<username>/.

Log in to Compute Canada server

Log in to the Compute Canada server you copied your data to:

\$ ssh -XY <username>@niagara.computecanada.edu

Setting up and using MIRACL

Load the specific Singularity module you would like to use (e.g. Singularity 3.5):

\$ module load singularity/3.5

Note: Not specifying the version will load the latest version available on your node

Since **MIRACL** will take up a significant amount of space, it is recommended to download and work with the **MIRACL Singularity** container in the scratch directory. First, navigate there:

\$ cd \$SCRATCH

Then pull (download) the **Singularity** container:

\$ singularity pull miracl_latest.sif library://aiconslab/miracl/miracl:latest

Attention: singularity pull requires **Singularity** version 3.0.0 or higher. Please refer to our *Troubleshoot-ing* section ("Q: Can I build a Singularity container from the latest MIRACL image on Docker Hub") if you are using an older version of Singularity.

Note: If you have a particular Singularity container of **MIRACL** that you want to use on Compute Canada, just copy it to the servers directly using e.g. scp or rsync instead of pulling (downloading) the latest version of **MIRACL** from the **Singularity** registry

Start the **MIRACL Singularity** container with the default folders mounted:

```
$ singularity shell miracl_latest.sif bash
```

Singularity will automatically mount your scratch folder to your container. If you need to mount a specific directory into a specific location, use the following:

Once you are logged in to the container, load the GUI from the shell:

\$ miraclGUI

Note: Please consult our *Troubleshooting* section on **Singularity** if you experience problems with opening **MIRACL's** GUI on Compute Canada

Or use **MIRACL** from the command line. For example, run **MIRACL's** CLARITY registration workflow on the folder that you copied over previously:

\$ miracl flow reg_clar -f input_clar -n "-d 5 -ch autofluo" -r "-o ARS -m combined -v 25"

Note: If you have a particular Singularity container of **MIRACL** that you want to use on Compute Canada, just copy it to the servers directly using e.g. scp or rsync instead of pulling (downloading) the latest version of **MIRACL** from the **Singularity** registry

Jupyter notebook

An accompanying Jupyter notebook for this tutorial can be found *here*.

3.9.2 Running MIRACL commands on Sherlock (Stanford supercomputer)

This tutorial highlights the registration workflow but a similar approach applies to other commands.

Setting up MIRACL (first time)

Log in to Sherlock:

\$ ssh -Y username@sherlock.stanford.edu

Start an interactive session:

\$ sdev

Move to your scratch folder:

\$ cd SCRATCH

Pull (download) Singularity container:

\$ singularity pull miracl_latest.sif library://aiconslab/miracl/miracl:latest

Attention: singularity pull requires **Singularity** version 3.0.0 or higher. Please refer to our Troubleshooting section ("Q: Can I build a **Singularity** container from the latest MIRACL image on Docker Hub") if you are using an older version of **Singularity**.

Tip: If you have a particular **Singularity** container of **MIRACL** that you want to use on Sherlock, just copy it to the servers directly using e.g. **scp** or **rsync** instead of pulling (downloading) the latest version of **MIRACL** from the **Singularity** registry

Copying your data to Sherlock

Copy a folder called, e.g. input_clar with tiff files that you want to register to the Allen Atlas using scp:

\$ scp -r input_clar sherlock.stanford.edu:/scratch/users/<username>/clarity_registration/ ...

or **rsync**:

Attention: Make sure to replace <username> with your Sherlock username

Running MIRACL in an interactive session

For quick jobs that don't require much resources you can login to Sherlock:

\$ ssh -Y username@sherlock.stanford.edu

Move to your scratch folder:

\$ cd SCRATCH

Start interactive session:

\$ sdev

Start **Singularity** with binded data:

\$ singularity shell miracl_latest.sif bash

Within the shell, load the GUI:

\$ miraclGUI

Or use the command-line:

\$ miracl lbls stats -h

Note: Please consult our Troubleshooting section if you experience problems with opening MIRACL's GUI on Sherlock

Running SBATCH jobs

If you want to run jobs with specific resources for larger, longer jobs (e.g. running the registration workflow) you can do the following:

First get the data orientation (please check the registration tutorial for setting orientation):

```
$ miracl conv set_orient
```

After setting the orientation, a file called ort2std.txt will be created that might look like this:

```
$ cat ort2std.txt
tifdir=/scratch/users/username/clarity_registration/input_clar
ortcode=ARS
```

Use that orientation code (ARS) in your registration workflow.

First check the workflow arguments:

```
$ miracl flow reg_clar -h
```

Assuming you wanted to run this command with the following arguments, for example on your data:

\$ miracl flow reg_clar -f input_clar -n "-d 5 -ch autofluo" -r "-o ARS -m combined -v 25"

Create an sbatch script named, for example reg_job.sbatch and paste the following lines:

```
#!/bin/bash
#SBATCH --job-name=clar_reg
#SBATCH --ntasks=1
#SBATCH --time=05:00:00
#SBATCH --cpus-per-task=12
#SBATCH --mem=32G
module load singularity
singularity exec ${SCRATCH}/miracl_latest.sif miracl flow reg_clar -f ${SCRATCH}/clarity_
oregistration/input_clar -n "-d 5 -ch autofluo" -r "-o ARS -m combined -v 25"
```

Attention: Note that the miracl function call comes after invoking the **Singularity** call singularity exec **\${SCRATCH}/miracl_latest.sif** and that full file paths were used for the .sif container and the input data

This sample job (called: clar_reg) asks for 5 hours, 12 cpus and 32G of memory on one node. Adjust the requested resources based on the job you are submitting.

Next submit the sbatch script:

\$ sbatch reg_job.sbatch

To check on the status of your submitted job use:

\$ squeue -u \$USER

See also:

For more resources on SLURM sbatch jobs check Stanford's tutorials on submitting and running jobs on Sherlock

Jupyter notebook

An accompanying Jupyter notebook for this tutorial can be found *here*.

FOUR

JUPYTER NOTEBOOKS

ACE STA Registration Regional statistics and visualization Installing MIRACL on Windows Singularity on Compute Canada and Sherlock

FIVE

TROUBLESHOOTING

Choose an troubleshooting issue from the sidebar menu or TOC.

If you cannot find an answer to your problem here, please open an issue on our offical GitHub page.

5.1 Docker

5.1.1 MIRACL's GUI (miracIGUI) is not working

Access control might be enabled on your host machine. Change your **xhost** access control on your host machine (not within your **Docker** container). Exit the running **Docker** container and type:

xhost +

Log back in to your container. Reset xhost after you are done with using the MIRACL GUI using:

xhost -

The above will enable/disable access for all users. If this is not the desired behavior, access can be granted in a more fine grained manner.

Example:

xhost +SI:localsuer:<yourusername>

Note: Replace <yourusername> with the actual user name of your host system

5.1.2 A The GUI worked before but does not work anymore

Navigate to the directory that contains your docker-compose.yml and restart the container with docker compose down followed by docker compose up -d. The GUI should work again.

5.1.3 **I** cannot run X or Y with Docker because of permission denied errors

If you have not set up a **Docker** user you might need to run **Docker** commands with sudo. While this should work, setting up a **Docker** user is the prefered.

5.1.4 Processes that require TrackVis or Diffusion Toolkit are not working

Because of their respective licenses, we could not include **TrackVis** or **Diffusion Toolkit** in our **Docker** image directly. Please download and install them on you host machine using their installation guide. After they have been successfully installed, mount a volume to your **MIRACL Docker** container that contains the binary folder for **TrackVis** and **Diffusion Toolkit** and add the binaries to your **\$PATH** within your **MIRACL Docker** container using the mounted volume.

5.1.5 **STA** workflow fails when trying to create tracts

Make sure that the **TrackVis** and **Diffusion Toolkit** binaries are available to **MIRACL**. See the previous question for details.

5.1.6 I need to install or make changes to apps in the MIRACL container but my user is not authorized to do so

The user in the container is the user of your host machine. This is done to avoid issues with the **MIRACL** GUI and X11. If you need to make changes that require sudo privileges, just log out of your container and log back in as root:

\$ docker exec -it -u root miracl bash

After making your changes, log out and log back in with your regular user:

\$ docker exec -it miracl bash

Attention: Always remembert that changes to the container are not persistent. If you need to make permanent changes as sudo, add them to the Dockerfile instead.

5.1.7 MIRACL's GUI (miraclGUI) does not work anymore after my ssh connection has been broken

Assuming you logged back in with ssh and are in the container, exit the container and restart it using docker compose down and docker compose up -d.

Shell back into the container and the GUI should work again.

Note: Note that the **Docker Compose** syntax is different if you installed it using the standalone method. **Compose standalone** uses the -compose syntax instead of the current standard syntax compose. The above command would thus be docker-compose up -d when using **Compose standalone**.

5.1.8 **A** I do not want to create the image using the provided script

You can build the image yourself, not using the script we provide. However, the build script makes sure that the GUI version of **MIRACL** works with **Docker** and it is therefore recommended to use it. Build the image with:

\$ docker build -t mgoubran/miracl .

Note: Do not forget the . at the end of the command. It is required to point docker build to the Dockerfile.

To run the container use:

\$ docker run -it mgoubran/miracl bash

Attention: If you make changes in the container that are not stored on a volume, make sure to use the same container the next time you run **MIRACL** as changes made to a container will only apply to this specific container. If you run **MIRACL** again from the same image using docker run -it mgoubran/miracl bash, a new container will be created that does not contain the changes you made to the first container.

Warning: The **MIRACL** GUI will be unlikely to work out-of-the-box but you can try the troubleshooting steps in the following section to make it work.

5.1.9 1 get either or both of the following errors whenever I try to run the GUI from within a Docker container that was build without the provided build script

Authorization required, but no authorization protocol specified qt.qpa.xcb: could **not** connect to display :1

Note: The number for the display could be different in your case

```
qt.qpa.plugin: Could not load the Qt platform plugin "xcb" in "" even though it was_

→found.

This application failed to start because no Qt platform plugin could be initialized._

→Reinstalling the application may fix this problem.
```

Available platform plugins are: eglfs, linuxfb, minimal, minimalegl, offscreen, vnc, wayland-egl, wayland, wayland-xcomposite-egl, wayland-xcomposite-glx, webgl, xcb.

Exit your running **Docker** container and run the following to mount an X11 socket from the host system in a new **Docker** container:

docker run -it -e DISPLAY=\$DISPLAY -v /tmp/.X11-unix:/tmp/.X11-unix mgoubran/miracl bash

If you still receive the above error, you may have to change your xhost access control. See previous troubleshooting step above.

5.2 Singularity

5.2.1 Can I build a Singularity container from the latest MIRACL image on Docker Hub

Absolutely! To do so, however, you will need to grab a development node after logging in to the cluster. If you try pulling from the login node, you will use a ton of memory building the SIF image, and the process will be killed (in other words, it won't work).

\$ sdev

or

\$ salloc

Once you have your node, you can then build the container:

```
$ cd $SCRATCH
$ singularity build miracl_latest.sif docker://mgoubran/miracl:latest
```

5.2.2 A Processes that require TrackVis or Diffusion Toolkit are not working

Because of their respective licenses, we could not include **TrackVis** or **Diffusion Toolkit** in our **Docker** image directly. Please download and install them on you host machine using their installation guide. After they have been successfully installed, mount a volume to your **MIRACL Docker** container that contains the binary folder for **TrackVis** and **Diffusion Toolkit** and add the binaries to your **\$PATH** within your **MIRACL Docker** container using the mounted volume.

5.2.3 **A** I get the following error whenever I try to run the GUI from within the Singularity container on Compute Canada

qt.qpa.plugin: Could not load the Qt platform plugin "xcb" in "" even though it was. → found. This application failed to start because no Qt platform plugin could be initialized. → Reinstalling the application may fix this problem. Available platform plugins are: eglfs, linuxfb, minimal, minimalegl, offscreen, vnc,.

→wayland-egl, wayland, wayland-xcomposite-egl, wayland-xcomposite-glx, webgl, xcb.

We do not recommend trying to make X11 forwarding work directly from the terminal. You should use VNC instead. Follow the instructions here. Once you are connected to your login or compute node with VNC, you will see a desktop environment. Open a terminal there and follow *our tutorials* on how to use **MIRACL** with **Singularity** on clusters.

If you for some reason need to run the **MIRACL** GUI directly in the terminal, using a **Singularity** container and X11, try the following workarounds:

Login Nodes

Exit your Singularity container and start a VNC server (for 3600sec or more as required) on your login node:

vncserver -MaxConnectionTime 3600

The first time the VNC server is started you will prompted for a password (do not leave this blank). Once done, check if a X11 socket is available for your username:

ls -la /tmp/.X11-unix/

Note: If no socket is available for your username, log out and log back in to your login node

Start another **Singularity** container and try to run miraclGUI again from within it.

Compute Nodes

Exit your **Singularity** container and set an environment variable on your allocated compute node:

export XDG_RUNTIME_DIR=\${SLURM_TMPDIR}

Start a VNC server:

vncserver

Start another **Singularity** container and try to run miraclGUI again from within it.

5.3 Local installation

5.3.1 A get the following error whenever I try to run the GUI: qt.qpa.plugin: Could not load the Qt platform plugin "xcb" in "{anaconda path}/envs/miracl_merge/lib/python3.7/site-packages/cv2/qt/plugins" even though it was found

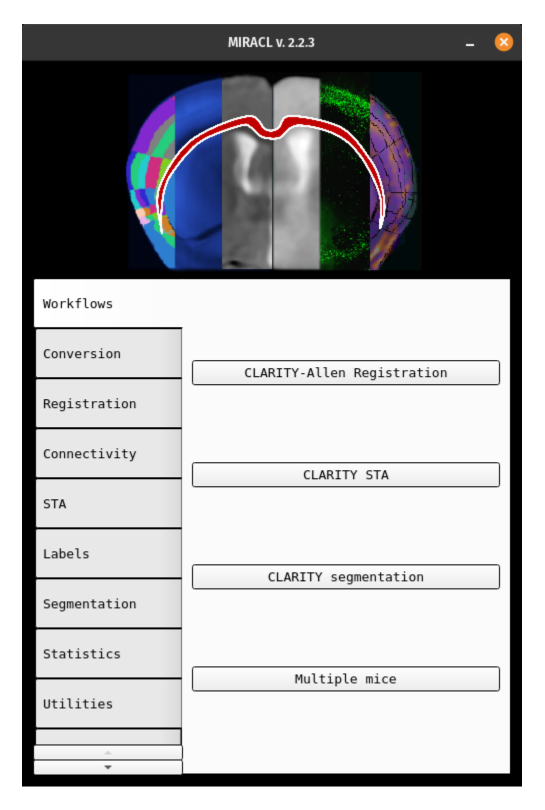
If you know the path to the environment name, try running the following line to remove the specific file in question:

SIX

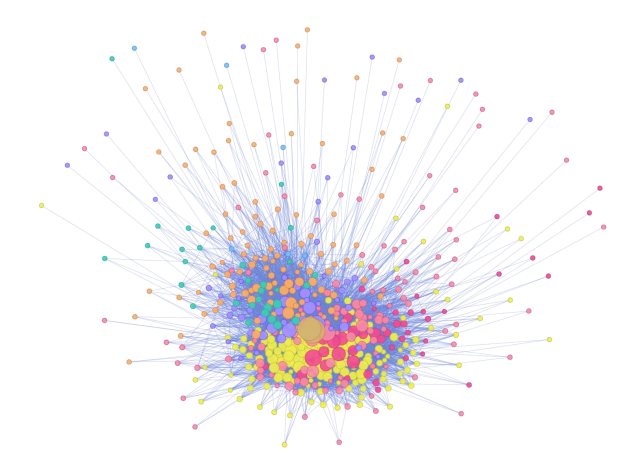
GALLERY

Here is some representative work!

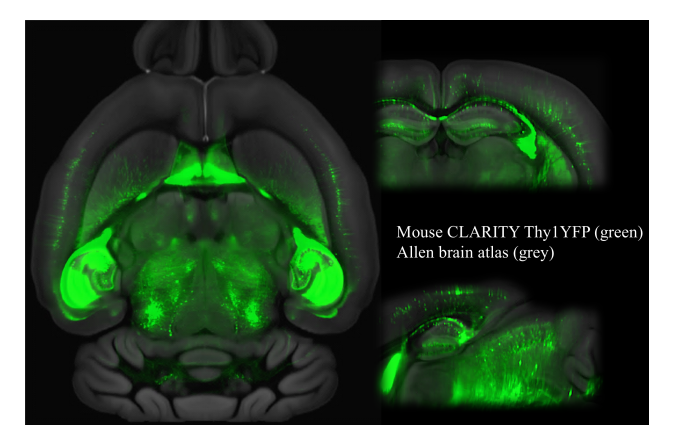
6.1 Graphical User Interface (GUI)



6.2 Brain Graph



6.3 Clarity Registration





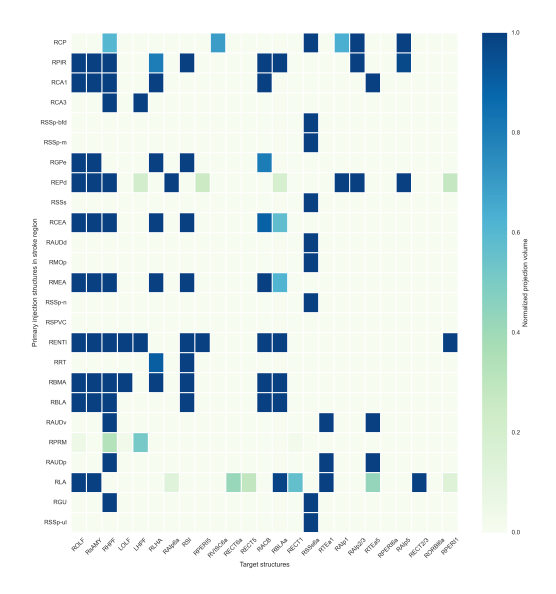


Fig. 1: Connectivity matrix heat map with 25 labels



Fig. 2: Connectogram grouped by parent ID with 50 labels

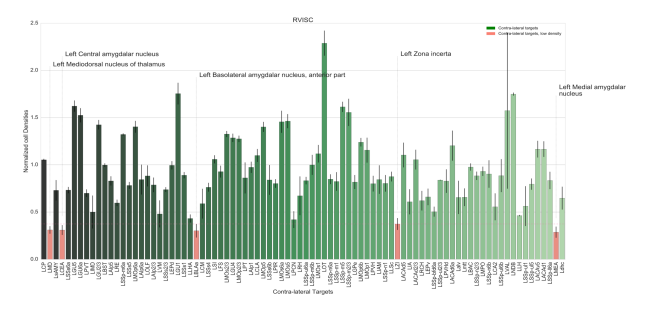
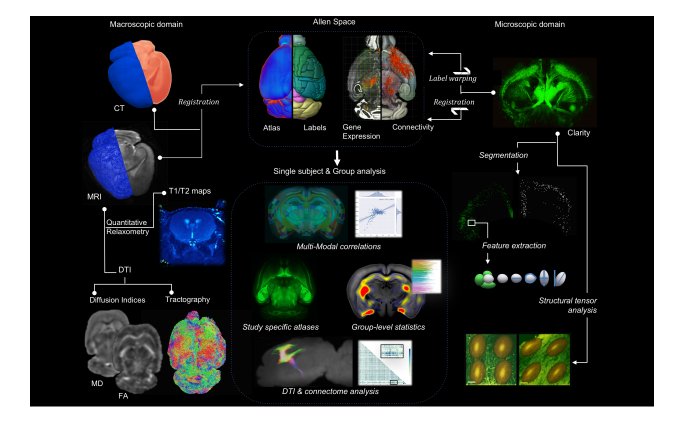


Fig. 3: Density along connectivity graph

6.4 Connectivity

6.5 Pipeline



6.6 Registration and Segmentation

RCP	RSSs5	RSNr	RSSs6a	RAIp5	RMOp5	RSSs1	RMOp2/3	RSSs2/3	RAIp2/3	RGU5 F	SSp-m6	RMOs5	RMOp6a	RMOs2/3	RAId5	RAId2/3	RSSs4	RVISC5	RGU2/3	RSSp-m	RVISC6a	RGU6a	RAlp1	RHPF	RMOp1
RPIR	ROLF	RHPF	RsAMY	RCOAp	ROT	RTR	RAON	RACB	RPAA	RBLAa	RCOAa	RSI	RBLAp	RMOB	RAIp2/3	REPv	RAAA	RENTm	RTT	RSUB	RAIp5	RPERI2/3	RBLAv	RLHA	RAIv5
RCA1	RHPF	ROLF	RSUB	RAON	RLSr	RACB	RTT	RTR	Rfi	RENTm	RBLAp	RBST	RCOAp	ROT	RsAMY	RTEa5	RLSc	RMOB	RLHA	Ralv	RILA5	REPv	RMS	RTEa6a	RECT5
RCA3	RHPF	LHPF	LCA1	LCA3	RCA2	RDG-mo	LCA2	RSUB	RDG-po	RLSc	RLSr	RDG-sg	LLSc	LLSr	Rfi	Lfi	LDG-po	LLD	RLD	LSUB	RTRS	LTRS	LDG-sg	RSF	Lalv
RSSp-bfd	Sp-bfd3	566p-bfd₽	SSp-bf@	SSp-bfd	RMOs2/3	B LCP	RSSs2/3	RMRN	RSSs6a	RSSp-bfd	1RSSs5	RVPM	RMOs5	RMOs1	RSCig	RSSs1	RPO	RSSs4	RMOs6a	RSCiw	RMOp6a	SSp-bfd6	a6Sp₊ul6a	a RZI	RAPN
RSSp-m	LCP	RMOp2/3	RMOp5	RMOp66	SSp-m2	RSSp-m6	BMOs2/3	RSSs6a	.SSp-m6	RSSp-m5	RVPM	RSSs5	RSSs2/3	RSSp-m1	RSSp-m4	RMRN	RMOp1	RAId5	RMOs1	RMOs5	RAId2/3	LSSs6a	SSp-m2/	BMOp6al	.SSp-m5
RGPe	RSNr	RMD	RPF	RSI	RLHA	RSTN	RMRN	RGPi	RVAL	RVM	RPCN	RZI F	RACAd6a	RMOs6a	ROLF	RSNc	RVTA	RsAMY	RCL	RMOs5	RPL5	RLH	RNPC	RACB	RCM
REPd	ROLF	RHPF	RCOAp	RAIp2/3	RPAA	RTR	RAIp5	REPv	RAlp6a	RsAMY	RAIp1 F	RPERI2/3	BRCOAa	RCLA	RBLAv	RBLAp	ROT	RENTm	RPERI1	RPERI5	RAIv2/3	LHPF	RBLAa	RENTmv	RAAA
RSSs F	RSSp-m6	RMOp2/3	RMOp6a	RSSs2/3	RSSs6a	RMOp S R	(SSp-m2/	3 LCP	RSSs5	RSSp-m4	RVPM P	RSSp-m5	LSSs6a	RSSs4	RSSs1	RMOs2/3	RMOs5	RMOp1	LSSs5	LSSs2/3	RMRN	RSSp-m*	RVISC5	RVISC2/3	RMOs1
RCEA	ROLF	RsAMY	RSI	RLHA	RHPF	RBST	RMRN	RCOAp	RPARN	RPAG	ROT	RIRN	RACB	RAAA	RPRNc	RTR	RSNr	RBLAp	RBLAa	RLPO	RCOAa	RPB	RPAA	RNTS	RNDB
B RAUDd	RIC	RAUDp5	AUDp2/	RAUDv5	RAUDp1	RAUDv2/:	RSSs2/3	RAUDd2/	RAUDd5	RSCig F	RAUDp6a	RSSs5	RMRN	RAUDd1	RSSs1	RSCiw	RSSp-tr2/	RSSs6R	SSp-bfd	RAUDpo:	RTEa2/3	RAUDv1	RAUDp 	SSp-bfd	1RTEa1
a stroke	LCP	RMOp2/S	RMOp6a	RMOp5	RMOs5	LMOp5F	SSp-m6	RMOs2/3	LMOp6a	RMRN	RMOs6a	RSSp-m8	LMOp2/3	LMOs5	RSSs6a	RMOp1	.SSp-m6	aRMOsf	SSp-m2/	3RVPM	RSSp-m ⁻	LMOs6a	RVAL	LIRN	RSSs5
RAUDd RAUDd I Store Legiou sturctmes I Store Legiou sturctmes I Store RMOp RMEA RSSp-n	ROLF	RsAMY	RHPF	RSI	RLHA	ROT	RCOAp	RBST	RMRN	RACB	RCOAa	RAAA	RTR	RPAG	RLPO	RPAA	RNDB	LCP	RAON	RSNr	RBLAa	RBLAp	REPv	RVTA	RNLOT1
uojto RSSp-n	RMOs2/8	RSSp-n@	SSp-n2/	RMOp2/3	RMOp6a	RMOs5	RMOs1	RSSp-n5	RSSs2/3	LCP	RVPM	RSSp-n4	RSSp-n1	RSSs5	RMOp5	RSSs6a	RMOs6a	RSSp-ul6	a RPO	RMRNR	SSp-bfd2	/BRMOp1	RSSs1	RSSp-mR	SSp-m2/3
ilui fuemind RSPVC	RSPVI	RPSV	RSPVO	RVII	RPARN	RsptV	RIRN	RMDRN	RMDRN	LGRN	LMRN	RGRN	RPGRNI	LIRN	RPB		LPRNc	RLRN	RPRNc	LPARN	LSCig	LMARN	LIO	LIC	RII
د RENTI	RHPF	ROLF	RACB	RSUB	RsAMY	ROT	RBLAa	LCP	RDG-mc	LACB	LOLF	LHPF	RBLAp	RCOAp	RAON	RENTm	RTR	RPERI2/	3 RTT	RSI	RPERI5	LOT	LENTI	RPERI1	LPIR
RRT	RVAL	RVPM	RPO	RPF	RSNr	RMD	RPCN	RVM	RVPL	RMOs6a	RLD	RMOp6a	RSI	RCL	RMOs5	RMRN	RVPMpc	RGPi	RZI	RLHA	RLP	RLH	RMOp5	RCM F	ACAd6a
RBMA	ROLF	RsAMY	RHPF	RACB	ROT	RBST	RSI	RCOAp	LCP	RLHA	LACB	RPAA	LOT	RCOAa	RAAA	RBLAp	RTR	RBLAa	LOLF	RLPO	RPA	LBST	RBLAV	REPv	RSUB
RBLA	LCP	RACB	ROT	ROLF	RsAMY	RSI	RHPF	LOT	RPL5	LACB	RMOs2/3				RBLAa		RMOs6a		RAAA			3 RPL1		RAIv5	LOLF
RAUDy		RAUDp5				RAUDp6a											RAUDv4		RIP					LTEa6a	
RPRM		RCOPY	LPYR	RPYR	RHPE		RFOTU			LCOPY	LPRM	ROLF	LNU	RSNr			LCA2		LFOTU	Lbic		LSCig	LSNr	LVISp1	LCA1
RAUDo	RIC	RALIDas		AUDp2/		RAUDp6a					RMG		RTEa2/3					-				-		RAUDv1	
		o to o po	KAOD PI		-																				
RLA						RsAMY							RECT5F				RPAA	REPv	RTEa4						
						8 RAId6a																			
RSSp-ul	RMOp2/3	2 LCP	RMOp5	RMOp6B	SSp-ul2	/3RMRN	RSSs6a	RSSp-ulB	Sp-bfde	10	RSSp-ul5	RSSs2/3	RPO 13	RVPL	RSSp-ul1	1 RMOp1	RMOs5F	18	BMOs2/3	20	21	RSCig F	23	RVPM	RSNr 25
		~		-		÷				10	Target s		rder along									**	20	2.4	

Fig. 4: Projection map along graph with 25 labels

30 Normalized projection volume

15

45

60

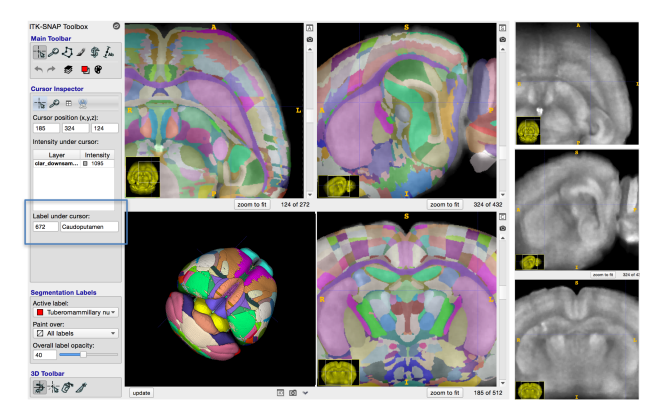


Fig. 5: Registration result visualized in ITK-SNAP

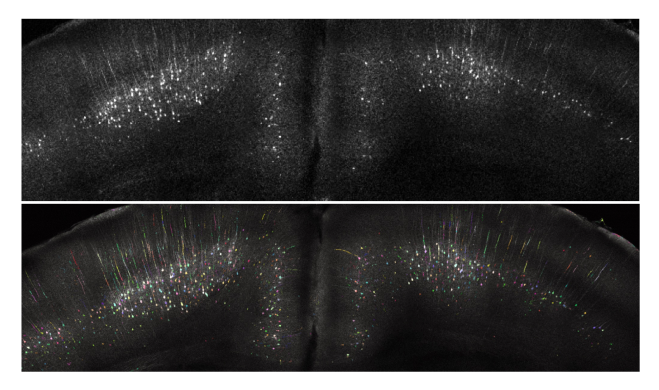


Fig. 6: Segmentation

SEVEN

DOWNLOADS

Here you can find downloads for e.g. atlasses, example data or workshop slides.

7.1 Data

Example datasets, atlases etc.

7.1.1 Example and atlases data

Sample data you can use to test MIRACL and the data for the atlases used by MIRACL can be found here:

Dropbox Data Link

The Atlases folder contains templates, annotations, histology, ontology graph info and LUT/label description of the Allen Reference Atlas (ARA).

The Data folder contains test data with example inputs and outputs for the registration and segmentation modules.

For a detailed description and input parameters please check the respective help or *tutorial* of each module.

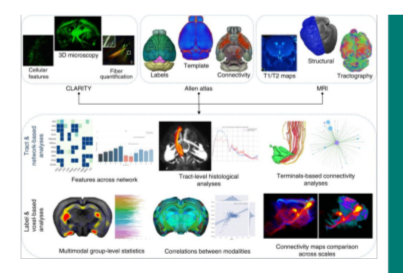
7.2 Workshops

Slides and notebooks for workshops given on MIRACL can be downloaded from here.

7.2.1 2024

March 20th: MIRACL Workshop at Stanford

Flyer



MARCH 20TH MIRACL WORKSHOP

Maged Goubran, PhD, Assistant Professor Canada Research Chair in Computational Neuroscience & AI University of Toronto

Workshop introducing the MIRACL platform, a computational platform that incorporates AI for image registration to reference atlas, atlas generation, segmentation, and novel sub-regional analyses.

MIRACL (Multi-modal Image Registration And Connectivity anaLysis) is a general-purpose AI-based pipeline for automated:

- Registration of tissue cleared and imaging data (eg. LSFM and MRI) to standard (eg. Allen Reference Atlas) or study-specific atlases
- 3D Segmentation/mapping of neuronal activity and feature extraction (eg. Whole brain c-fos)
- Tract-specific or network-level connectivity analysis (eg. Virus tracing)
- Regional and cluster-wise (sub-regional) statistical analyses & visualization

Wu Tsai Neurosciences Institute Stanford University Neuroscience Microscopy Service WHOLE BRAIN ATLAS REGISTRATION

LARGE DATA VOLUME WORKFLOWS

CELL / C-FOS SEGMENTATION AND CLUSTER ANALYSIS

TRACT TRACING & CONNECTIVITY ANALYSIS

ATLAS GENERATION

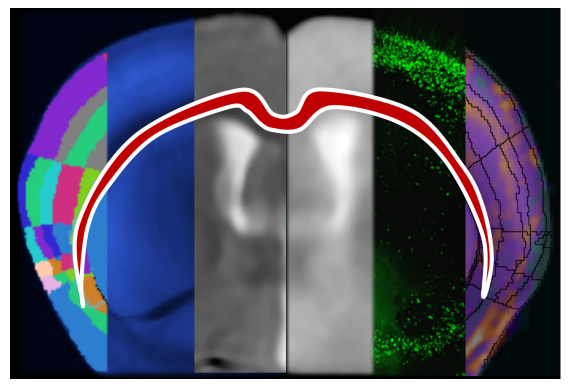
STANFORD NEUROSCIENCE BUILDING

2 90 Jane Stanford Way James Ling and Nisa Leung Seminar Room E153

March 20th, 1:30pm

Notebooks

ACE STA Registration Regional statistics and visualization Installing MIRACL on Windows Singularity on Compute Canada and Sherlock



MIRACL (Multi-modal Image Registration And Connectivity anaLysis) is a general-purpose, open-source pipeline for automated:

- 1. Registration of cleared and imaging data (ex. LSFM and MRI) to atlases (ex. Allen Reference Atlas)
- 2. 3D Segmentation and feature extraction of cleared data
- 3. Tract-specific or network-level connectivity analysis
- 4. Statistical analysis of cleared and imaging data
- 5. Comparison of dMRI/tractography, virus tracing, and connectivity atlases
- 6. Atlas generation and Label manipulation

NEW WORKFLOW/FEATURE RELEASE

We have released our AI-based Cartography of Ensembles (ACE) workflow, an end-to-end, automated pipeline that integrates cutting-edge deep learning segmentation models and advanced statistical methods to enable unbiased and generalizable brain-wide mapping of 3D alterations in neuronal activity, morphology, or connectivity at the sub-regional and laminar levels beyond atlas-defined regions.

The tutorial for using ACE can be found here.

We recommend using MIRACL with the Docker or Singularity containers we provide but it can also be installed locally. See our *installation instructions* for more information.

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