# A tutorial on how to use the Brain Observatory Toolbox

This tutorial will show you how to access the Brain Observatory data using MATLAB. In particular, this tutorial will show you how to:

- 1) Download meta information from brain observatory api and save it as MATLAB datatype
- 2) Build a brain\_observatory\_cache object to:
- A) Get general information on the brain observatory data
- B) Filter sessions by specified criteria in different aspects such as brain areas, imaging depth and stimuli type
- C) Download NWB files of filtered sessions
- 3) Extract imaging data from NWB files to:
- a) Get and plot fluorescence traces
- b) Convert fluorescence trace into data that is in "raster format" for further analysis

#### **Organization of the Brain Observatory Data**

Before we start, we first want to describe the organization of the Brain Observatory Data.

An experiment container contains three sessions (also called experiments or ophys\_experiments) where recordings were made on a singe mouse, in a single brain region and at a particular imaging depth. Each of these three sessions consists of a series of "subexperiments" where a particular stimulus set was shown. NWB files downloaded from the Allen Institute API each consist of data from a single session. As described below, all data that we extract into "raster format" consists of data from a single "subexperiment" where a particular stimulus set was shown.

Note: within an experiment container the same stimuli might be repeated in different sessions (i.e., different sessions can have the same "subexperiment type"). For example, Natural Movie 1 is shown in all three sessions in an experiment container. For more information see: http://alleninstitute.github.io/AllenSDK/brain\_observatory.html

# 1) Download meta information from brain observatory api and save it as MATLAB datatype

To begin, set up the path to Brain Observatory Toolbox:

```
bot_dir_name = 'Brain-Observatory-Toolbox/';
```

Then, add it to MATAB's search path

```
addpath(bot_dir_name)
```

Next we will download the meta information which is stored in manifest files on the Allen API. We can do this using the function get\_manifests\_info\_from\_api(). This function creates a MATLAB structure array with three tables and saves this structure array in a file called manifests.mat. Manifests.mat will be taken by the brain\_observatory\_cache as shown in the following section.

```
get_manifests_info_from_api()

Elapsed time is 0.000814 seconds.

load('manifests')
```

There are three tables in manifest.mat: container\_manifest contains metadata of all experiment containers (m containers by n aspects of meta data), session\_manifest contains metedata of all sessions (m sessions by n aspects of meta data), and cell\_id\_mapping contains mapping between new cell ids and the old cell id of all cells (m cells by number of ids).

As quoted from AllenSDK 's release note (0.13.2): "The cross-session alignment algorithm has been updated and re-run, so all cell specimen IDs have changed".

# 2) Build a brain\_observatory\_cache object

Brain\_observatory\_cache is a class that resembels BrainObservatoryCache in allensdk (https://github.com/AllenInstitute/AllenSDK/blob/master/allensdk/core/brain\_observatory\_cache.py).

It takes manifests and helps you fanthom and access Brain Observatory Data through three stages:

Stage A, it returns general information on the brain observatory data from different aspects through correpsonding methods whose names all start with "get\_";

Stage B, it filters sessions by different criteria through correpsonding methods with generic name pattern "filter\_...by\_...";

Stage C, it downloads NWB file(s) of filtered session(s) through the method named "download nwb".

To beign, build a brain observatory cache object

```
stimuli: []
targeted_structure: []
imaging_depth: []
container_id: []
session_id: []
session_type: []
```

As you can see here, manifests.mat is stored in property manifest, table session\_manifest from manifest.mat is stored in property session\_table, and table container\_manifest from manifest.mat is stored in property container\_table.

The other properties will shine more light during the Stage B: filtering sessions.

### 2A) Get general information on the brain observatory data

We implemented several methods whose names all start with "get\_" to summairze the brain observatory data from different aspects.

Here shown total number of containers

```
boc.get_total_num_of_containers()
ans = 199
```

Note: Total number of sessions should be whatever this returns \* 3

Here shown all cortical depths that were ever recorded in any experiment container

Note: Recording didn't take place equally among these depths, only popular depths are shown on homepage of Brain Observatory: http://observatory.brain-map.org/visualcoding

Here shown all type of cre driver lines of all mice participated

```
boc.get_all_cre_lines()
```

```
ans = 6×1 cell array
'Cux2-CreERT2'
'Emx1-IRES-Cre'
'Nr5al-Cre'
'Rbp4-Cre_KL100'
'Rorb-IRES2-Cre'
'Scnn1a-Tq3-Cre'
```

Note: All mice had the same reporter line: Ai93 and tTA driver line: Camk2a-tTA

For more information about transgenic lines: http://observatory.brain-map.org/visualcoding/transgenic

Here shown all brain regions that were ever recorded in any experiment container

```
boc.get_all_targeted_structures()

ans = 6×1 cell array
    'VISal'
    'VISam'
    'VISp'
    'VISpm'
    'VISprl'
```

For their locations, see homepage of Brain Observatoy: http://observatory.brain-map.org/visualcoding

Here shown all types of sessions that ever appeared in any experiment container

```
boc.get_all_session_types()

ans = 4×1 cell array
    'three_session_A'
    'three_session_B'
    'three_session_C'
    'three_session_C2'
```

Note: There are always three sessions in each container: session A, session B, and session C or session C2

Here shown all type of stimuli that were ever used in any of the four types of sessions

```
boc.get_all_stimuli()
ans =
[]
```

Note: For mapping between session type and stimulus type, see http://alleninstitute.github.io/AllenSDK/brain\_observatory.html

Here shown the numer of experiment containers recorded in each brain region

```
boc.get_summary_of_containers_along_targeted_structures()
```

VISal 33 VISam 25 VISl 36 VISp 54 VISpm 35 VISrl 16

Here shown the numer of experiment containers recorded at each cortical depth

boc.get summary of containers along imaging depths()

Here shown the number of experiment containers recorded at each cortical depth in each brain region

boc.get\_summary\_of\_containers\_along\_depths\_and\_structures()

ans = $12 \times 7$	table						
	VISal	VISam	VISl	VISp	VISpm	VISrl	total
175	8	5	10	11	11	5	50
265	1	Θ	0	0	0	Θ	1
275	12	9	15	20	11	5	72
300	1	0	1	1	1	0	4
320	0	1	0	0	0	Θ	1
325	0	1	1	Θ	1	0	3
335	0	0	0	2	1	0	3
350	4	4	4	13	4	4	33
365	1	Θ	0	0	0	Θ	1
375	6	4	5	7	6	2	30
435	0	1	0	Θ	0	0	1
total	33	25	36	54	35	16	199

# 2B) Filter sessions by specified criteria in different aspects such as brain areas, imaging depth and stimuli type

Now we proceed to Stage B: filtering sesssions.

Build a new "clean" brain\_observatory\_cache object first and I will explain right after.

To review, manifests.mat is taken in by brain\_observatory\_cache and stored in property "manifets", the manifest of all sessions (m sessions by n aspects of meta data) from manifest.mat is stored in property "session\_table", and the manifest of all containers (m containers by n aspects of meta data) from manifest.mat is stored in property "container\_table".

We implemented several methods with generic name pattern "filter\_...by\_..." to filter sessions by criteria in different aspects. As methods get called, property filtered\_session\_table, which started with the same content as session\_table, gets eliminated to meet new criteria, and then other properties that each summarize the meta data of all filtered sessions stored in "fitered\_session\_table" from a different aspect ("meta meta data") get updated.

Here we show an example of searching for sessions that primary visual cortex was recorded at 275 mm deep and during which drifting gratings were shown.

Eliminate all sessions to sessions that only showed drifting gratings

```
boc.filter_sessions_by_stimuli('drifting_gratings')

ans =
   brain_observatory_cache with properties:

        session_table: [597×14 table]
        container_table: [199×13 table]
        manifests: [1×1 struct]
        filtered_session_table: [199×14 table]
            stimuli: {'drifting_gratings' 'natural_movie_one' 'natural_movie_three' 'spontaneone' targeted_structure: {6×1 cell}
        imaging_depth: [11×1 double]
        container_id: [199×1 double]
        session id: [199×1 double]
```

```
session_type: {'three_session_A'}
```

Eliminate current filtered sessions to sessions that have posterior Primary Visual Cortex recoreded

Eliminate current filtered sessions to sessions that were recoreded at 275 mm deep

As we can see now, there are 20 sessions that met all of the criteria given.

manifests: [1×1 struct]

filtered\_session\_table: [1×14 table]

Let's just get one random session out of the 20 sessions, so we don't wait forever when I show you how to download real data at the next stage

```
stimuli: {'drifting_gratings' 'natural_movie_one' 'natural_movie_three' 'spontaned
targeted_structure: 'VISp'
   imaging_depth: 275
   container_id: 527550471
      session_id: 527745328
   session_type: {'three_session_A'}
```

Take a look at metedata of the final filtered session to convince you how friendly my property list ("meta meta data") is

boc.filtered_session_table				
ans = 1×14 table	ovneriment container id	fail ove tracking	; d	imaging don't
date_of_acquisition 	experiment_container_id	Tait_eye_tracking	id 	imaging_dept
'2016-07-08T15:59:05Z'	5.2755e+08	true	5.2775e+08	275

Hint: In future days, when you get lots NWB files downloaded and you lose track of who the heck they are, you can use brain\_observatory\_cache to look up their meta data and "meta meta data".

# 2C) Download NWB files of filtered session(s)

Finally comes to Stage C: downading real data stored in NWB files

For more info about NWB,

See its github page: http://neurodatawithoutborders.github.io/

See its publication: http://www.sciencedirect.com/science/article/pii/S0896627315009198

First, set up the path to NWB directory.

```
nwb_dir_name = [bot_dir_name,'nwb_files/'];
```

Then, call the method named "download\_nwb" to dowload all NWB file(s) corresponding to filtered session(s) stored in brain\_observatory\_cache and save it under your NWB directory.

```
boc.download_nwb(nwb_dir_name);
```

desired nwb file already exists

Note: The size of a nwb file is at the scale of 100 MB, so you definielty want to move around instead of sitting here and waiting. Once brain\_observatory\_cache gets the file downloaded, it will spit you a message, so no worries.

## 3) Import imaging data from nwb files

Now we have acquired an NWB file, we can extract the florescence traces from the NWB file and convert them to data in "raster format" (n trials by t times for every cell), which is easier to analyze.

For more info about raster format, see <a href="http://www.readout.info/toolbox-design/data-formats/raster-format/">http://www.readout.info/toolbox-design/data-formats/raster-format/</a>

To start, add the NWB file to MATLAB searching path.

```
% add path to nwb files addpath([bot_dir_name, 'nwb_files/'])
```

## 3a) Get and plot fluorescence traces

Before we start converting data format, you might just want to plot some fluorescence traces to make sure they look normal or for fun.

To do this you can either

- i) get fluorescence traces of all cells in one session and plot them in your own way or
- ii) use the built-in plot function to plot fluorescence traces of one cell in one session

Here we show an example of i):

Extract and compute all four types fluorescence traces available in the NWB file of session 517745328 and save them to the worksapce.

```
session_id = 527745328;
[raw,demixed,neuropil_corrected,DfOverF] = get_fluorescence_traces(session_id);
```

```
Elapsed time is 4.392045 seconds.
```

Note: each type of trace was stored as h5read table (k dimensions of cells by t dimesions of times) and saved as n dimesions of sampling points by k dimensions of cells matlab matrix

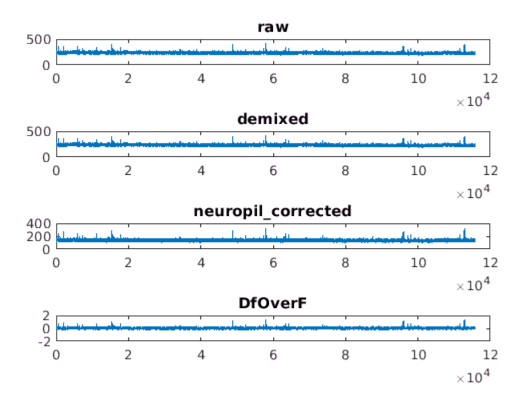
Alternatively, this is an example of ii):

Plot fluorescence traces of a random cell 529022196 in session 527745328 directly

Caution: "plot\_fluorecence\_traces" takes one new cell id

```
session_id = 527745328;
cell_id = 529022196;
plot_fluorecence_traces(session_id,cell_id);
```

Warning: MATLAB has disabled some advanced graphics rendering features by switching to software OpenGL. For more information, click here.



### 3b) Convert fluorescence trace into data in "raster format"

To start, set up the path to raster directory

```
raster_dir_name = [bot_dir_name, 'raster/'];
```

Tell the "convert" function the session you are working on, the type of stimuli you are interested in, and the type of fluorescence trace you want to use, and then set it off.

```
session_id = 527745328;
stimuli = 'drifting_gratings';
fluorescence_trace = 'DfOverF';

current_raster_dir_name = convert_fluorescenece_trace_into_raster_format(fluorescence_trace,...
    session_id, stimuli,raster_dir_name);
```

Brain-observatory-lootbox/raster/drifting_gratings_52//45328/	atready	exists
Note: This doesn't take long.		

4)	Get	Masks
71		MIGSKS