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) Instantiate a sessionfilter object to:
- A) Get general information on the brain observatory data
- B) Filter sessions by specified criteria such as brain areas, imaging depth and stimuli type
- 2) Instantiate a session object to:

Retrieve and examine kinds of experiment data as listed below of an interessed session from its Neurodata Without Borders (NWB) file:

- a) ROI masks
- b) fluorescence traces
- c) running speed
- d) motion correction
- e) eye tracking
- 3) Call function convert_fluorescence_trace_into_raster_format to:

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. Neurodata Without Borders (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

sessionfilter is a class that resembels BrainObservatoryCache in allensdk (see Python code at:https://github.com/AllenInstitute/AllenSDK/blob/master/allensdk/core/brain_observatory_cache.py).

The MATLAB sessionfilter enables you to get information and access data using three types of methods:

- A) Get general information on the brain observatory data using methods that start with "get_"
- B) Filter sessions by different criteria using methods that start with "filter_"

To beign, build a brain_observatory_cache object

1A) Get general information on the brain observatory data

There are several methods start with "get_" which summairze in the brain observatory data based on particular criteria.

To get the total number of experiment containers we can use:

275 300

```
bosf1.get_total_num_of_containers()
ans = 181
```

Note: Total number of sessions is the number of containers times 3 since there are three sessions per container.

To get all the cortical depths (um) that were ever recorded in any experiment container we can use:

```
bosf1.get_all_imaging_depths()

ans =
    175
    265
```

```
320
325
335
350
365
375
```

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

To get all the get all type of cre driver lines from all mice we can use:

```
bosf1.get_all_cre_lines()

ans = 6×1 cell array
    'Cux2-CreERT2'
    'Emx1-IRES-Cre'
    'Nr5a1-Cre'
    'Rbp4-Cre_KL100'
    'Rorb-IRES2-Cre'
    'Scnnla-Tg3-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

To get all the all brain regions that were recorded in any experiment container we can use:

```
bosf1.get_all_targeted_structures()

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

For more information on these locations, see homepage of Brain Observatoy: http://observatory.brain-map.org/visualcoding

To get all the all types of sessions that appear in any experiment container we can use:

```
bosf1.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

To get all the all type of stimuli that were used in any of the four types of sessions we can use:

```
bosf1.get_all_stimuli()

ans = 9x1 cell array
    'drifting_gratings'
    'locally_sparse_noise_4deg'
    'locally_sparse_noise_8deg'
    'natural_movie_one'
    'natural_movie_three'
    'natural_movie_two'
    'natural_scenes'
    'spontaneous'
    'static_gratings'
```

Note: For mapping between session type and stimulus type see http://alleninstitute.github.io/AllenSDK/brain_observatory.html

To get the number of experiment containers recorded in each brain region we can use:

```
bosfl.get_summary_of_containers_along_targeted_structures()

VISal 35
VISam 25
VISl 38
VISp 66
VISpm 36
VISrl 16
```

To get the number of experiment containers recorded at each cortical depth we can use:

```
bosf1.get_summary_of_containers_along_imaging_depths()
175 53
```

```
250
          1
265
          1
275
         75
300
          4
320
          1
325
          3
335
          3
350
         33
365
          1
375
         40
435
          1
```

To get the number of experiment containers recorded at each cortical depth in each brain region we can use:

bosf1.get summary of containers along depths and structures()

ans = 11×7 table							
	VISal	VISam	VISl	VISp	VISpm	VISrl	total
175	8	5	10	11	11	5	50
265	1	0	0	0	0	0	1
275	11	9	14	17	9	5	65
300	1	0	1	1	1	Θ	4
320	0	1	0	0	0	Θ	1
325	0	1	1	0	1	0	3
335	0	0	0	2	1	Θ	3
350	2	3	3	10	4	3	25
365	1	0	0	0	0	Θ	1
375	6	4	5	5	6	2	28
total	30	23	34	46	33	15	181

1B) Filter sessions by specified criteria such as brain areas, imaging depth and stimuli type

sessionfilter.filtered_session_table is an m-by-n table listing the n fields of meta data of m successful sessions. As different filter methods get called on sessiionfilter, rows/sessions in filtered_session_table are liminated to meet specified criteria, and other properies (e.g., stimuli, targeted_structure, etc) of sessionfilter, that are essentially important fields of meta data in filtered_session_table, are also updated.

Here we show an example of searching for sessions that Rorb-IRES2-Cre mice had their primary visual cortex recorded at 275 μ m deep, during where drifting gratings were shown and eye tracking went through.

To ensure an "inviolate" sessionfilter:

ans =

```
bosf1.clear_filters
```

Eliminate sessions to the ones where drifting gratings were shown:

```
bosf1.filter_sessions_by_stimuli('drifting_gratings')
```

```
eye_tracking_avail: [2×1 logical]
```

Note that several properties of bosf have changed.

Eliminate sessions to the ones that have posterior Primary Visual Cortex recordings:

Eliminate sessions to the ones that were recoreded at 275 um deep

Eliminate sessions to the ones that were operated on mice of Rorb-IRES2-Cre

```
bosf1.filter_session_by_cre_line('Rorb-IRES2-Cre')

ans =
    sessionfilter with properties:

    valid_session_table: [543×15 table]
    filtered_session_table: [5×15 table]
        stimulus: {4×1 cell}
    targeted_structure: {'VISp'}
        imaging_depth: 275
        container_id: [5×1 double]
        session_id: [5×1 double]
        session_type: [5×1 categorical]
```

```
cre_line: {'Rorb-IRES2-Cre'}
eye tracking avail: [2×1 logical]
```

Eliminate sessions to the ones that eye tracking completely went through

```
bosf1.filter_session_by_eye_tracking(1)
ans =
```

As we can see now, there are 2 sessions belonging to 2 diffrerent containers that meet all of the criteria given.

Every experiment container in the Brain Obervatory has an unique experiment container ID that was created by the Allen Institute. These container IDs are stored in the <code>experiment_container_id</code> field in <code>filtered_session_table</code>. For example, to see the experiment container IDs that met our filtering criteria we can run:

```
bosf1.filtered_session_table.experiment_container_id

ans =
   512124562
   511510989
```

If we want to find out the three sessions in the first experiment contianer, we can call:

```
bosf2 = bot.sessionfilter
```

bosf2.filter_sessions_by_container_id(bosf1.filtered_session_table.experiment_container_id(1)) ans = sessionfilter with properties:

bosf2.session id

```
ans = 512124564
512176430
512270518
```

Now let's dig into the real data.

2) Instantiate a session object to:

Retrieve and examine kinds of experiment data as listed below of an interessed session from its Neurodata Without Borders (NWB) file:

To illustrate this, let's instantiate a session object for the first session in bosf1.

Alternatively, we can instantiate an array of session objects for both sessions in bosf1 and replace bos1 with bos0(1) in the codes below.

```
bos0 = bosf1.get_filtered_sessions

bos0 =
   1×2 session array with properties:
   sSessionInfo
```

All the meta data stored in <code>bosf2.filtered_session_table</code> for this session are carried over to <code>bos1.sSessionInfo</code>

bos1.sSessionInfo

2a) ROI Masks

If you want to take a look at specific cells or all cells visually, you can extract and plot thier pixel masks. You can also pull out the maximum intensity projection of the movie for context.

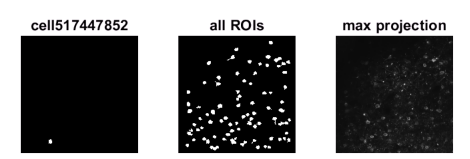
```
% get all the cell ids
cell_ids = bosl.get_cell_specimen_ids()
cell_ids = 90×1 int64 column vector
```

```
cell_ids = 90×1 int64 column vector
517447852
517447402
517447321
517447330
517447325
517447730
517447755
517447745
517447847
```

When you call this or any other method of session class, BOT automatiacally searchs for the corresponding NWB file and downloads it if it is not there yet.

Dowanloading an NWB file from their server can take about 20 min.

```
% plot the ROI mask for the first cell in the list
subplot(1,3,1)
imshow(bos1.get roi mask array(cell ids(1)))
title(['cell' num2str(cell ids(1))])
% plot the ROI masks for all celss
all mask = logical(zeros(512,512));
for iCell = 1: length(cell_ids)
    all mask = or(all mask, bos1.get roi mask array(cell ids(iCell)));
subplot(1,3,2)
imshow(all mask)
title('all ROIs')
% plot the maximum intensity projection
subplot(1,3,3)
imshow(double(bos1.get max projection)/max(max(double(bos1.get max projection))))
% colormap(gray)
title('max projection')
```



2b) Fluorescence Traces

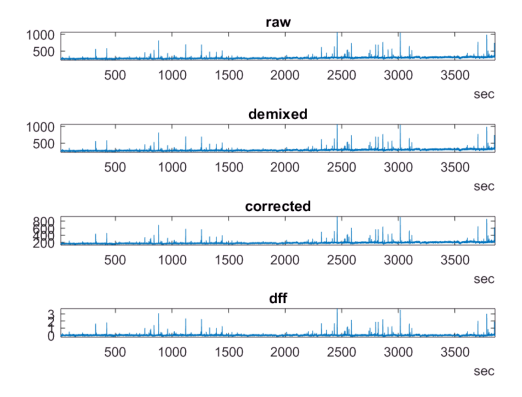
You can get the fluorescence traces of a specific cell by its index. Let's take cell 589163560 as an example.

```
[time, raw] = bos1.get_fluorescence_traces;
[time, demixed] = bos1.get_demixed_traces;
```

```
[time, neuropil] = bos1.get_neuropil_traces;
neuropil_r = bos1.get_neuropil_r;
[time, corrected] = bos1.get_corrected_fluorescence_traces;
% % compute it yourself
% corrected = demixed - neuropil * (neuropil_r .* eye(size(neuropil_r,1)));
[time, dff] = bos1.get_dff_traces;
iCell = bos1.get_cell_specimen_indices(cell_ids(1))
```

iCell = 1

```
subplot(4,1,1)
plot(time, raw (:,iCell))
axis tight
title ('raw')
subplot(4,1,2)
plot(time, demixed (:,iCell))
axis tight
title ('demixed')
subplot(4,1,3)
plot(time, corrected(:,iCell))
axis tight
title ('corrected')
subplot(4,1,4)
plot(time, dff(:,iCell))
axis tight
title ('dff')
```



You can also examine the fluorescence traces of all cells recorded in a session one cell at a time in sequence. The traces will be plotted in an external figure, hit any key to proceed to the next cell.

```
figure('Position', get(0, 'Screensize'))
set(gcf, 'Visible', 'on')

for iCell = 1:length(cell_ids)
clf

ft_to_be_plotted = {'raw', 'demixed', 'corrected', 'dff'};

for ift = 1: length(ft_to_be_plotted)
subplot(length(ft_to_be_plotted),1,ift)
cur_ft = eval(ft_to_be_plotted{ift});
plot(time, cur_ft(:,iCell))
% axis tight
ylim([0,max(max(cur_ft))])
title(ft_to_be_plotted{ift})
end

fprintf('Program paused. Press any key to continue to the next cell.\n');
pause;
end
```

Program paused. Press any key to continue to the next cell. Program paused. Press any key to continue to the next cell.

```
Program paused. Press any key to continue to the next cell.

Program paused. Press any key to continue to the next cell.

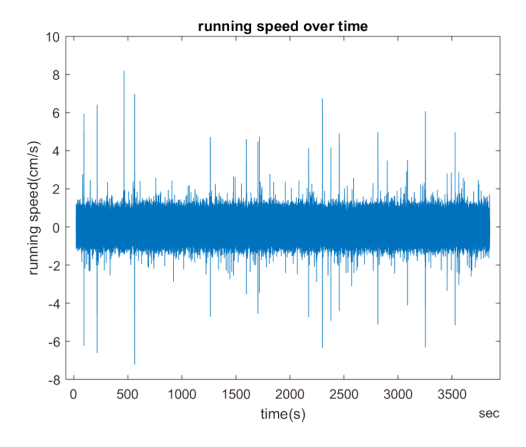
Program paused. Press any key to continue to the next cell.

close all
```

2c) Running Speed

All experiments contain running speed. Extreme outliers from the tracking have been removed and replaced with NaN, which will appear as gaps in the plotted data.

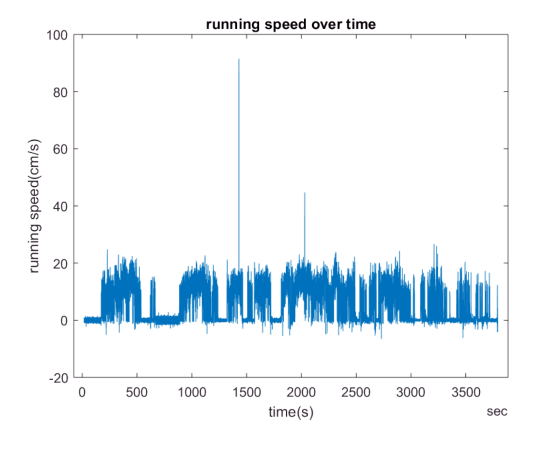
```
[time, runningSpeed] = bos1.get_running_speed;
plot(time, runningSpeed)
title('running speed over time')
xlabel('time(s)')
ylabel('running speed(cm/s)')
```



As you can see here, the mouse in this session didn't run much. Here is another session 580124131, whose running data is more exciting:

bos2.sSessionInfo

```
ans = struct with fields:
        date of acquisition: '2017-04-07T17:11:58Z'
    experiment container id: 580051757
          fail_eye_tracking: 0
                         id: 580124131
              imaging_depth: 375
                       name: '20170407_300663_3StimB'
                specimen id: 571153214
              stimulus_name: 'three_session_B'
          storage directory: '/external/neuralcoding/prod22/specimen 571153214/ophys experiment 58012413
      targeted structure id: 417
       experiment container: [1x1 struct]
           well known files: [1×1 struct]
         targeted structure: [1×1 struct]
                   specimen: [1x1 struct]
                   cre line: 'Rbp4-Cre KL100'
[time, runningSpeed] = bos2.get running speed;
plot(time, runningSpeed)
title('running speed over time')
xlabel('time(s)')
ylabel('running speed(cm/s)')
```

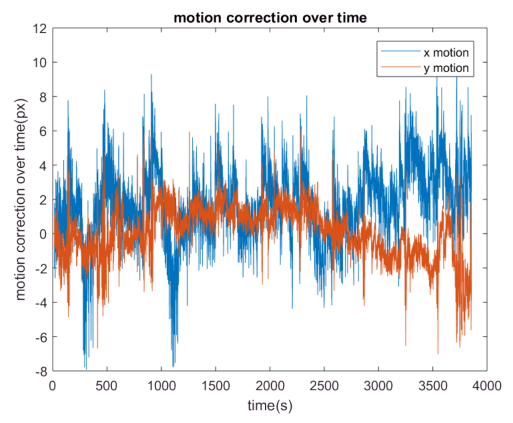


2d) Motion Correction

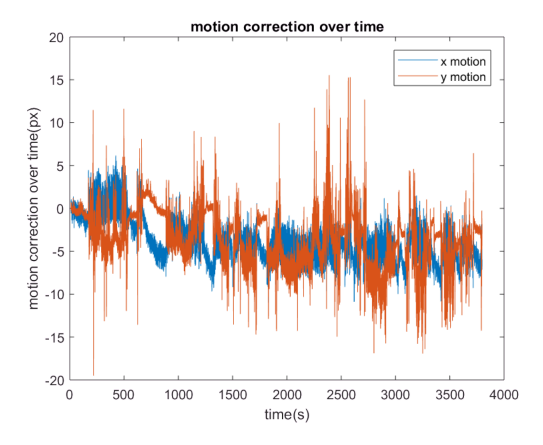
X and Y translation values in pixels required to correct for motion artifacts during the experiment are available as well.

As you can expect, since the mouse in bos2 ran more than the mouse in bos1, there was more motion correction in bos2 than in bos1.

```
close all
mc = bosl.get_motion_correction;
plot(table2array(mc(:,'timestamp')), table2array(mc(:,'x_motion')),table2array(mc(:,'timestamp'))
title('motion correction over time')
xlabel('time(s)')
ylabel('motion correction over time(px)')
legend('x motion','y motion')
xlabel('time(s)')
```



```
mc = bos2.get_motion_correction;
plot(table2array(mc(:,'timestamp')), table2array(mc(:,'x_motion')),table2array(mc(:,'timestamp'))
title('motion correction over time')
xlabel('time(s)')
ylabel('motion correction over time(px)')
legend('x motion','y motion')
xlabel('time(s)')
```

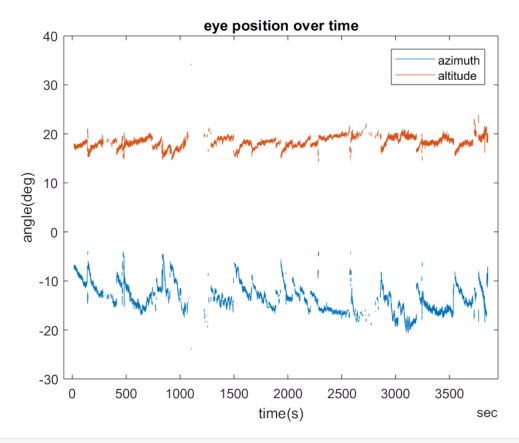


Eye Tracking

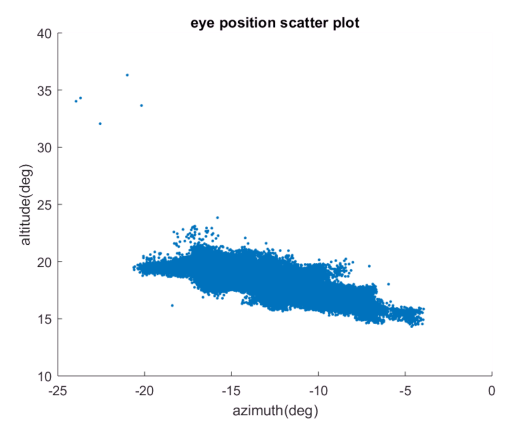
Many experiments contain pupil position and pupil size from eye tracking. Extreme outliers from the tracking have been removed and replaced with NaN, which will appear as gaps in the plotted data. If an experiment does not have eye tracking data, a NoEyeTrackingException will be raised.

As shown in bos1.sSessionInfo, the session in bos1 doesn't have eye tracking data, so let's filer out all sessions with eye tracking data and takes the first session in the list to demonstrate this functionality.

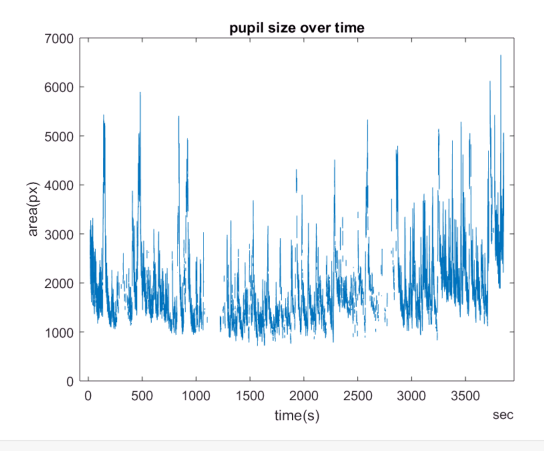
```
[time, pupilLocation] = bos1.get_pupil_location;
azimuth = pupilLocation(:,1);
altitude = pupilLocation(:,2);
plot(time, azimuth, time, altitude)
title('eye position over time')
legend ('azimuth', 'altitude')
xlabel('time(s)')
ylabel('angle(deg)')
```



```
scatter(azimuth, altitude, '.')
title('eye position scatter plot')
xlabel('azimuth(deg)')
ylabel('altitude(deg)')
```



```
[time, pupilArea] = bos1.get_pupil_size;
plot(time, pupilArea)
title('pupil size over time')
xlabel('time(s)')
ylabel('area(px)')
```



3) Call function convert_fluorescence_trace_into_raster_format to:

Convert fluorescence trace into data that is in "raster format" for further analysis

bot.util.convert_fluorescence_trace_into_raster_format('Df0verF', bos1.sSessionInfo.id,'drift

All raster files already exist in directory [raster\drifting_gratings\drifting_gratings_512270518].