3- Networked Hoc in BMTK

Instructions

This document assumes you have completed the necessary steps in 02-Single Cell Hoc BMTK

In the previous guide you developed a single cell network using a single half-center oscillator cell. In this guide we'll take it a step further by adding a second cell and networking them together.

- 1. To keep our previous model intact we'll create a whole new set of configuration files.
- 2. Start by creating a new build_network1.py script and paste the following code into that file:

```
build network1.pv
2
   from bmtk.builder.networks import NetworkBuilder
 3
 4
   net1 = NetworkBuilder('hco net')
 5
   net1.add nodes(N=1,
                  cell name='HCOCell1',
 6
 7
                  model type='biophysical',
 8
                  model template='hoc:HCOcell',
 9
                  morphology='blank.swc'
10
11
12
   net1.add nodes(N=1,
13
                  cell name='HCOCell2',
                  model type='biophysical',
14
15
                  model template='hoc:HCOcell',
                  morphology='blank.swc'
16
17
18
19
   net1.add edges(source={'cell name': 'HCOCell1'},
20
   target={'cell name':'HCOCell2'},
21
                  connection rule=1,
22
                  syn weight=40.0e-02,
                  dynamics_params='GABA InhToInh.json',
23
24
                  model template='Exp2Syn',
25
                  delay=0.0,
26
                  target sections=["soma"],
27
                  distance range=[0,999])
28
   net1.add edges(source={'cell name': 'HCOCell2'},
29
   target={'cell name':'HCOCell1'},
30
31
                  connection rule=1,
32
                  syn weight=40.0e-02,
                  dynamics params='GABA InhToInh.json',
33
34
                  model template='Exp2Syn',
35
                  delay=0.0,
                  target sections=["soma"],
36
                  distance range=[0,999])
37
```

```
38
39   net1.build()
40   net1.save_nodes(output_dir='network')
41
42   net1.build()
43   net1.save_edges(output_dir='network')
44
```

- 3. Things to note about this new file:
 - a. Line 4: 'hco_net' will be used for network component files later, be descriptive
 - b. The cell name 'hco1' and 'hco2' have been assigned to nodes that have been added to keep the cells separate
 - c. Line 12: We can call add nodes more than once
 - d. Line 19: The add edges method creates connections between cells
 - i. Line 19: Specify the source cells via parameters of that cell, set in the previous add nodes call. You can specify multiple conditions for that cell to connect.
 - ii. Line 20: Target cells can be set in the same way. When using multiple networks, specify the target (or source) in the following way:

```
target=otherNetwork.nodes(cell_name="poolosyn_cell")
```

iii. Line 23: dynamics_params specifies a file located in \biophys components\synaptic models

That file will contain information in the following way, and can be duplicated or modified: For now, this file exists and does not need to be modified.

```
GABA_InhToInh.json

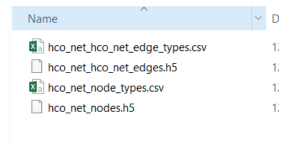
1 {
2    "level_of_detail": "exp2syn",
3    "tau1": 0.2,
4    "tau2": 8.0,
5    "erev": -70.0
6  }
7
```

- e. Line 42 and 43. Just as the nodes were saved to the network directory, we must also save the edges.
- f. We can specify a different output folder for our network components for multiple simulations, but we won't here.
- 4. Now you should be ready to build your network. In your Anaconda Prompt, in the root of your directory execute the following command to build your network:

```
python build network.py
```

A successful run may not have any output.

5. Visit your network directory (network) You should have several files for nodes and edges. Make note of the files listed:



6. It's now time to create a new circuit config file, create a file named circuit_config1.json. Copy the contents from your previous circuit_config.json file and update the node/edge references. The files referenced here are from the previous step. Your file should like to the following:

```
circuit config1.json
 1
 2
 3
      "manifest": {
 4
        "$BASE DIR": ".",
 5
        "$COMPONENTS DIR": "$BASE DIR/biophys components",
        "$NETWORK DIR": "$BASE DIR/network"
 6
 7
 8
      "components": {
 9
        "morphologies dir": "$COMPONENTS DIR/morphologies",
        "synaptic models dir": "$COMPONENTS DIR/synaptic models",
10
        "mechanisms dir": "$COMPONENTS DIR/mechanisms",
11
12
        "biophysical neuron models dir":
13
    "$COMPONENTS DIR/biophysical neuron templates",
        "point neuron models dir": "$COMPONENTS DIR/point neuron templates",
14
15
        "templates dir": "$COMPONENTS DIR/hoc templates"
16
      "networks": {
17
        "nodes": [
18
19
            "nodes file": "$NETWORK DIR/hco net nodes.h5",
20
21
            "node types file": "$NETWORK DIR/hco net node types.csv"
22
23
        "edges": [
24
25
            "edges file": "$NETWORK DIR/hco net hco net edges.h5",
26
            "edge types file": "$NETWORK DIR/hco net hco net edge types.csv"
27
28
29
30
31
32
33
```

- 7. Things to note about circuit config1.json:
 - a. When using multiple networks and multiple edges between networks you will need to have multiple blocks of node_file/node_type_file enclosed by brackets, separated by commas.
- 8. Next we'll set up our simulation by creating a new simulation configuration file (simulation_config1.json) You can either copy your old configuration file into this and edit it or copy the following:

```
simulation config1.json
1
 2
      "manifest": {
 3
        "$BASE DIR": ".",
        "$OUTPUT DIR": "$BASE DIR/output"
 4
 5
 6
      "target simulator": "NEURON",
 7
      "run": {
 8
        "tstop": 1000.0,
        "dt": 0.1,
 9
       "dL": 20.0,
10
        "spike threshold": -15.0,
11
        "nsteps block": 5000
12
13
      "conditions": {
14
15
        "celsius": 34.0,
16
        "v init": -50.0
17
      },
18
19
      "node sets": {
        "hco1": {
20
21
          "cell name": "HCOCell1"
22
23
      },
24
25
      "inputs": {
      "current clamp": {
26
          "input type": "current clamp",
27
          "module": "IClamp",
28
          "node set": "hcol",
29
30
          "amp": 1.0,
31
          "delay": 50.0,
32
          "duration": 10.0
33
34
      },
35
      "output": {
36
        "log file": "log.txt",
        "output dir": "${OUTPUT DIR}",
37
        "spikes file": "spikes.h5",
38
        "spikes file csv": "spikes.csv",
39
        "overwrite output dir": true
40
41
      } ,
42
      "reports": {
43
          "membrane report": {
          "module": "membrane report",
44
```

```
45
          "cells": "all",
          "variable name": [
46
47
             "cai",
             " 77 "
48
49
50
          "file name": "cell vars.h5",
51
          "sections": "soma"
52
53
      "network": "./circuit config1.json"
54
55
56
```

- 9. Things to note about simulation config1.json
 - a. Line 8: tstop has been modified to make the simulation last longer
 - b. Line 9: dt has been increased to make the simulation run faster in less detail
 - c. Line 16: v_init has been set to a value that will not cause the HCO cell to spike spontaneously as it did previously
 - d. Line 19: A "node_set" has been defined. This set can be used later in the file and is based on properties of individual cells. In this case we want to target HCO Cell 1 and only provide an input to it initially. The "node_sets_file":

 "\$BASE_DIR/node_sets.json" line must be deleted from the file because the node set has already been defined.

e.

- f. Line 26: A current clamp input has been created to target the previously mentioned node set "hco1" input parameters are set here directly.
- g. Line 54: The network configuration file has been updated to our new circuit built in the last step.
- 10. You should now be ready to run your network and view the output. Execute the following command to specify your new simulation config1.json file.

```
python run bionet.py simulation config1.json
```

A successful run will output the following

```
(clean) C:\Users\Tyler\Desktop\BMTK Morpho Work\bmtk-howto\my_bmtk_model>python run_bionet.py simulation_config1.json 2018-12-31 18:47:04,286 [INFO] Created log file 2018-12-31 18:47:04,397 [INFO] Building cells. 2018-12-31 18:47:04,420 [INFO] Building recurrent connections
```

```
2018-12-31 18:47:04,431 [INFO] Running simulation for 1000.000 ms with the time step 0.100 ms
2018-12-31 18:47:04,431 [INFO] Starting timestep: 0 at t_sim:
0.000 ms
2018-12-31 18:47:04,435 [INFO] Block save every 5000 steps
2018-12-31 18:47:06,205 [INFO] step:5000 t_sim:500.00 ms
2018-12-31 18:47:07,786 [INFO] step:10000 t_sim:1000.00 ms
2018-12-31 18:47:07,834 [INFO] Simulation completed in 3.403 seconds
```

If you receive "PermissionError: [WinError 5] Access is denied: './output'" just run the network again.

11. Run python test plot.py to view your new results.

