



HANDS-ON

A morphologically-detailed neural network simulation library
for contemporary high performance computing architectures

12TH DECEMBER 2018 | ANNE KÜSTERS & ALEXANDER PEYSER

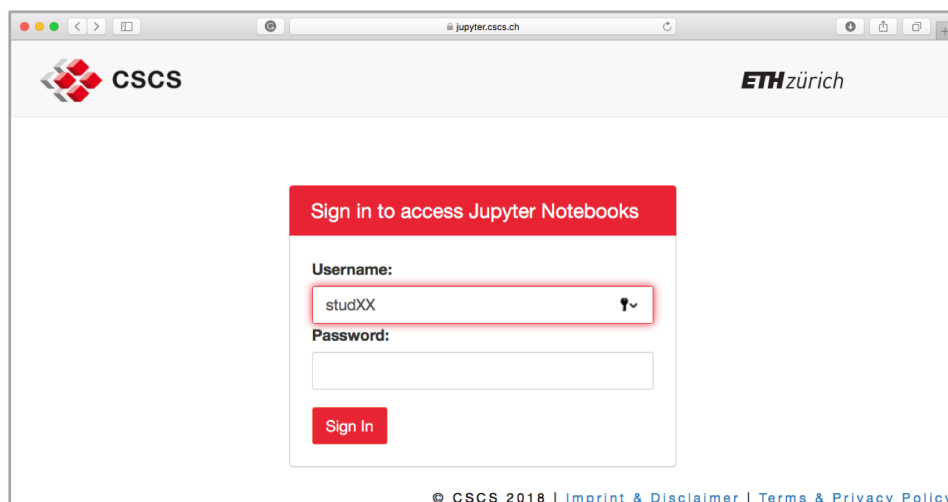


Co-funded by
the European Union



PREPARATION

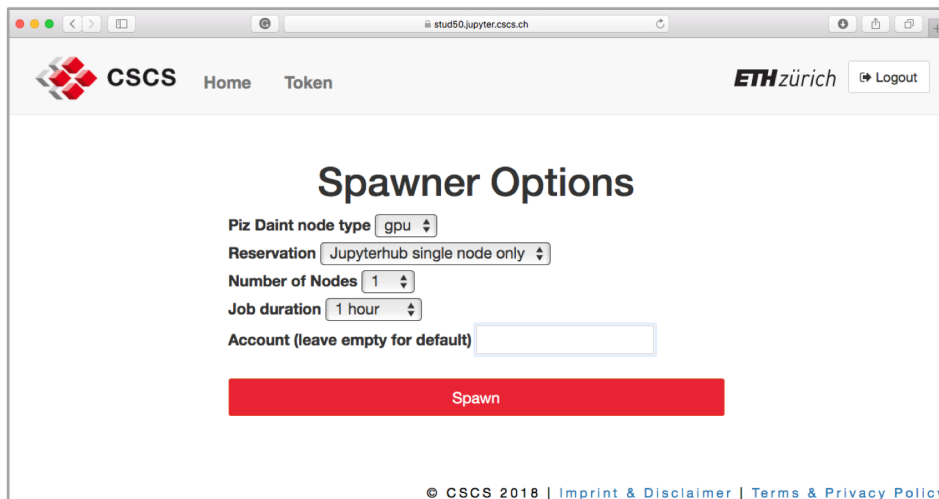
Sign in to <https://jupyter.cscs.ch> with your training username and password



The screenshot shows a web browser window with the address bar displaying `https://jupyter.cscs.ch`. The page header includes the CSCS logo on the left and the ETH zürich logo on the right. The main content area features a red-bordered box with the title "Sign in to access Jupyter Notebooks". Inside this box, there are two input fields: "Username:" with the text "studXX" and a dropdown arrow, and "Password:" with an empty field. Below these fields is a red "Sign In" button. At the bottom of the page, there is a footer with the text "© CSCS 2018 | [Imprint & Disclaimer](#) | [Terms & Privacy Policy](#)".

PREPARATION

Spawn a GPU with reservation of one node for one hour



ORGANIZATION AND WORK-FLOW

4 tasks, each dependent on the previous

Structure:

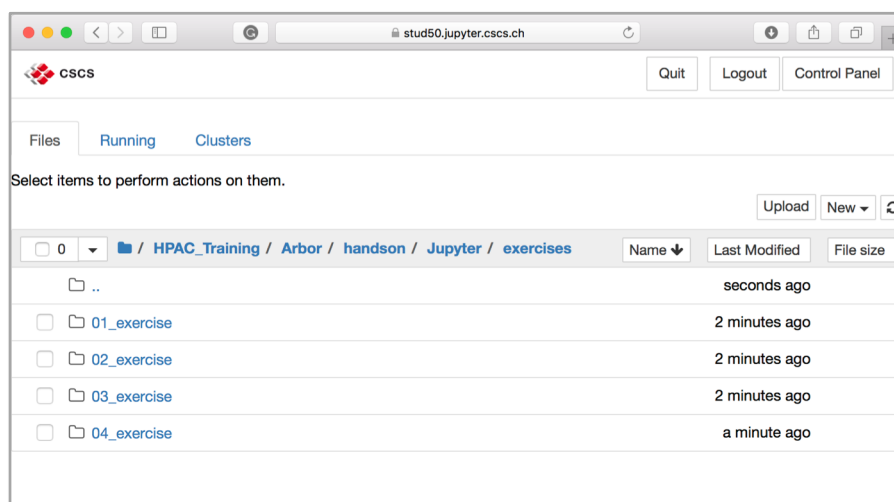
- On your training account, go to:
 - [HPAC_training](#) → [Arbor](#)
 - [handson](#) → [Jupyter](#) → [exercises](#)
 - [0X_exercise](#) (where X is the exercise number 1-4)
- Here, you find a python script with your `#TODO#`
- Each exercise builds up on the previous exercise
 - Work on your TODO`s and save your solution to the next exercise folder
 - In case your solution does not work (after really trying and asking tutors), use provided script in next exercise



Source of picture: flaticon.com

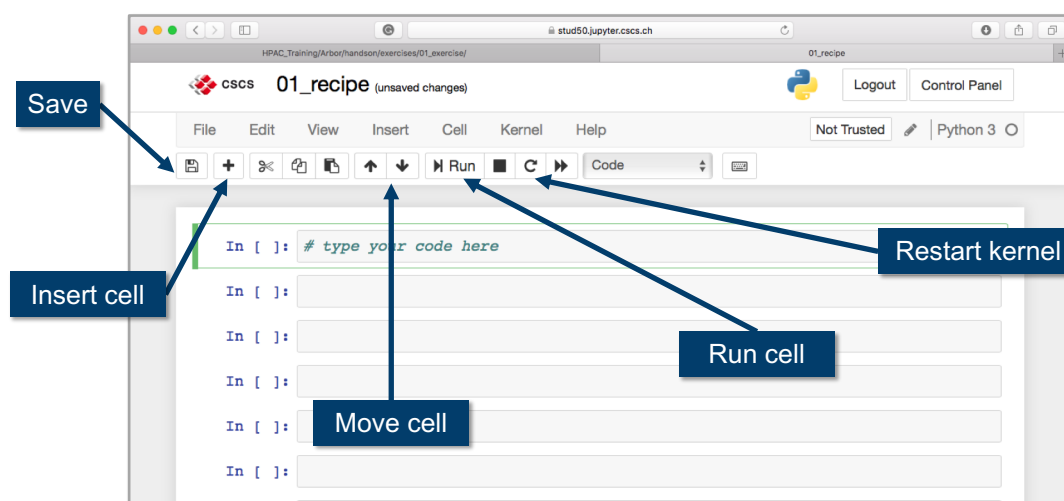
ORGANIZATION AND WORK-FLOW

4 tasks, each dependent on the previous



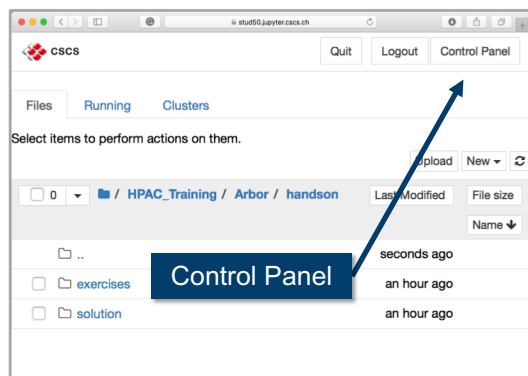
SHORT INTRODUCTION OF JUPYTER NOTEBOOKS

Easy to use



SHORT INTRODUCTION OF JUPYTER NOTEBOOKS

End your session via “Control Panel – Stop My Server” after completing the tasks



OVERVIEW OF EXERCISES

Exercises follow steps to setup and run a simulation with Arbor's python frontend



1. Describe neuron model by defining a **recipe**



2. Get the **resources**, create the parallel **execution context** and part the work into segments by partitioning the **load balance**



3. **Initiate the simulation** over the distributed system and **run** the simulation



4. (OPTIONAL:
Set up measurement meters, get spike **recorder and its spikes** to print meter report and spike times of the cells)

Source of pictures: flaticon.com

RECIPE

Distinction between the description of a model, and the execution of a model (simulation)

A **recipe** is a description of a model. The recipe is queried during the model building phase to provide cell information, such as:

- the *number of cells* in the model as input argument (`num_cells`)
- the *type* of a cell (`kind`)
- a *description* of a cell with soma, synapses, detectors, stimuli (`cell_description`)
- optionally, e.g.:
 - the number of spike *targets* (`num_targets`)
 - the number of spike *sources* (`num_sources`)
 - incoming network *connections* on a cell (`connections_on`)



TASK 1: CREATE A RECIPE FOR RING NETWORK

- a) Make a soma cell with Arbor's `make_soma_cell()`
- b) Add a synapse to the cell with the cell's function `add_synapse()` which takes a location, here at segment 0 position 0.5, using Arbor's `segment_location(segment, position)`
- c) Add a detector to the cell with the cell's function `add_detector(location, threshold)` with a threshold of 20 mV
- d) Add a stimulus to the cell with `gid 0` at $t_0=0$ ms for a period of 20 ms with weight 0.01 nA using the cell's function `add_stimulus(location, start, duration, weight)`
- e) Define the source as the previous cell with `gid-1`, bearing in mind that the cell with `gid 0` has the last cell in the ring network with `num_cells` as source



HARDWARE RESOURCES

Arbor supports running on systems from laptops and workstations to large distributed HPC clusters

Therefore, Arbor uses distributed **contexts** to

- Describe the computer system that a simulation is to be distributed over and run on.
- Perform collective operations over the distributed system.
- Query information about the distributed system.

The global context is determined at run time taking the **computational resources** as input. Further, the user can choose between using a non-distributed (local) context, or a distributed MPI context (if available).

An execution context is created by a user before building and running a simulation. This context is then used to perform domain decomposition and initialize the simulation.



DOMAIN DECOMPOSITION AND LOAD BALANCING

Arbor is performance portable

A **domain decomposition** is a description of the distribution of the model over the available computational resources. The description partitions the cells in the model as follows:

- Group the cells into cell groups of the same kind of cell.
- Assign each cell group to either a CPU core or GPU on a specific MPI rank.

A **load balancer** is a distributed algorithm that

- Determines the domain decomposition to balance the work over threads (and GPU accelerators if available).
- Uses the model recipe and a description of the available computational resources as inputs.



TASK 2: CREATE PARALLEL EXECUTION CONTEXT



- a) Get all available local hardware resources by using Arbor's `proc_allocation()`.
- b) Create a context by using Arbor's `context()` that uses the local resources, and an MPI communicator for distributed communication.
- c) Initiate the `recipe()` defined in task 1 with 100 cells.
- d) Partition the simulation over the distributed system by using Arbor's `partition_load_balance()` that uses the recipe and the context handle.

FROM RECIPE TO SIMULATION



A simulation needs a recipe, a context and a domain decomposition

To build a **simulation** the following are needed:

- ✓ • A recipe that describes the cells and connections in the model.
- ✓ • A context used to execute the simulation.
- ✓ • Thereof, a domain decomposition describing the distribution of the model over the local and distributed hardware resources.

The workflow to build a simulation is, thus, to first generate a domain decomposition, then initiate the simulation that is the executable form of a model. A simulation is initiated from a recipe, the decomposition and the context. Then the simulation is used to update and monitor the model state.



TASK 3: BUILD AND RUN THE SIMULATION



- a) Initiate the simulation by using Arbor's `simulation()` which takes the recipe, the domain decomposition and the context.
- b) Run the simulation using the simulation's command `run()` for a simulated time of 2000 ms with a time stepping size of 0.025 ms.

OPTIONAL: METER MEASUREMENT AND SPIKE RECORD

To measure memory and (if available) energy consumption

For measuring memory (and energy) consumption Arbor's **meter manager** can be used. First the meter manager needs to be initiated, then the metering started and checkpoints set, wherever the manager should report the meters. The measurement starts from the start to the first checkpoint and then in between checkpoints.

Checkpoints are defined by a string describing the process to be measured. The **meter report** finally collects all meters.

A **spike recorder** monitors all spikes recorded by a detector on the cells during the simulation.



OPTIONAL TASK 4: MEASURE



- a) Initiate Arbor's `meter_manager()` and start measuring by using the meters' `start()` function that takes the context as input.
- b) Set checkpoints using the meters' `checkpoint()` function which takes a string and the context as input
 - i. „recipe create“ after initiating the recipe
 - ii. „load balance“ after partitioning the simulation
 - iii. „simulation init“ after initiating the simulation
 - iv. „simulation run“ after running the simulation



OPTIONAL TASK 4: GET RESULTS



- c) Between initiating and running the simulation, build the spike recorder using Arbor's `make_spike_recorder()` which takes the simulation handle as input
- d) Make and print a meter report by using Arbor's `make_meter_report()` function that takes the meters and the context as input.
- e) Get the recorder's spikes
- f) Play around with parameters and see what happens.

SUMMARY

Using Arbor's python frontend

We learned how ...

- to describe a neuron model **using a recipe**;
- to get **resources**, create a **parallel execution context** and partition the **load balance**;
- **initiate the simulation** over the distributed system and run the simulation;
- set up **measurement meters**, get spikes **recorded** and print a meter report along with the spiking times of the cells.

Further, we learned how to use and create Jupyter notebooks on CSCS' JupyterHub.

LITERATURE AND LINKS

- *Arbor – a morphologically-detailed neural network simulation library for contemporary high-performance computing architectures*, High Performance Computing for Neuroscience, PDP2019
- <https://github.com/arbor-sim/arbor>
- <https://arbor.readthedocs.io>