SpiNNaker Human Brain Project Portal Introduction

This document describes how to use the EBRAINS portal to run PyNN scripts on SpiNNaker machines.

1. Getting Started

This section describes the current state of the SpiNNaker machine hosted by the HBP Portal and the software stack's limitations.

1.1 SpiNNaker Machine

Before starting to run PyNN scripts on SpiNNaker machines via the portal, be aware that the total SpiNNaker machine capacity is a 10 cabinet machine, comprising 1,036,800 cores, as shown in Figure 1.



Figure 1: A 10 cabinet machine, wired together (Note: the 11th cabinet in the picture contains servers for handling jobs sent to the machine).

1.2 Software Limitations

To understand how this relates to a PyNN script, be aware that SpiNNaker stores all the data needed to execute a PyNN model within its on-board memory (SDRAM). This SDRAM is limited in size, as each SpiNNaker chip contains only 128MB of memory, split dynamically between the 18 cores (including the operating system core) that reside on the chip. On average, every core has available approximately 8MB to store the neuron parameters as well as the synaptic matrix of the atoms¹ it is executing. Most models, to date, are limited to 256 atoms per core², and therefore the connectivity between these 256 atoms and the rest of the simulation must fit within these constraints.

Due to these limitations, the maximum possible number of atoms that can be simulated by a PyNN script on the largest SpiNNaker machine available on the HBP portal is 250,675,200.

¹ Atoms here represent the atomic element which each core models. In the case of PyNN, these are neurons from a given population.

² This limitation may be lifted in the future, but to date, this is due the way that synaptic delays are implemented on SpiNNaker.

This sounds rather large, but when it is factored in that each atom can only have a maximum incoming fixed number of connections of 8000, and that any more than this will result in a smaller number of neurons per core; therefore a smaller total neurons being simulated.

To make matters worse, currently, if a PyNN Projection between two Populations has delays greater than 16 machine time steps³ then each core that simulates these neurons will have a delay model attached to it, which in practice reduces the maximum number of simulated neurons by half.

Finally, if the PyNN model includes plastic connections, these require more memory than fixed connections and therefore will further reduce the number of neurons per core. A summary of the rest of the SpiNNaker back end limitations can be found here: <u>https://spinnakermanchester.github.io/latest/spynnaker_limitations.html</u>

1.3 EBRAINS Portal Limitations

The EBRAINS portal executes its PyNN scripts in a batch mode process. This means certain capabilities supportable by the SpiNNaker software stack (sPyNNaker) are not currently avilable by the EBRAINS portal. These are:

- 1. Real time visualisation of an executing PyNN script to a standalone visualization kit.
- 2. Closed loop simulation with real robotics.
- 3. Closed loop simulation with robots via the virtual robotic environment.

1.4 Script Limitations

This section describes some basic limitations that the PyNN script can experience when running on the EBRAINS Portal.

- 1. Figures generated by the script must be stored in file format, no figures will be presented to the end user in an interactive mode.
- 2. A PyNN script running on the SpiNNaker backend needs to adhere to the subset of PyNN that is supported by sPyNNaker⁴, a list of what is not implemented can be found here:

https://spinnakermanchester.github.io/latest/spynnaker_limitations.html

1.5 Recommendations

It is recommended that you test your simulation with the sPyNNaker front end in virtual mode before attempting to run it through the EBRAINS portal. Instructions on how to install the sPyNNaker front end can be found here:

https://spinnakermanchester.github.io/latest/PyNNOnSpinnakerInstall.html

³ This is 16 milliseconds in a simulation that runs at 1ms time steps, or 1.6 milliseconds in a simulation running at 0.1 ms time steps

⁴ This is the SpiNNaker software stack's front end for simulating PyNN neuron models. See <u>https://www.frontiersin.org/articles/10.3389/fnins.2018.00816/full</u> for more details.

and instructions on how to operate the tool chain in virtual mode can be found here:

https://spinnakermanchester.github.io/latest/VirtualMode.html

2. Running a PyNN script on the EBRAINS portal

This section describes how an end user can run a PyNN simulation on the EBRAINS Portal / collaborator, both through the Neuromorphic Computing Job Manager (section 2.2) and through the SpiNNaker Jupyter service (section 2.3). We suggest that if you are testing out PyNN scripts / networks, then you use the Jupyter service to interact with the machine; then, if you wish to run more jobs once you are happy with your network and wish to get some results (e.g. to do parameter sweeps), use the NMC Job Manager to submit scripts.

2.1 Getting an EBRAINS account

To acquire an EBRAINS account, go to <u>https://ebrains.eu/register</u> and follow the instructions.

2.2 Creating a Job on the EBRAINS portal/collaboratory.

1. First you must log onto the EBRAINS collaboratory via the link in the top right of the page at:

https://wiki.ebrains.eu/bin/view/Main/

and you may then also find it useful to read the "Getting Started" documentation at

https://wiki.ebrains.eu/bin/view/Collabs/the-collaboratory/Getting%20Started/

Collaboratory Collabor	ocumentation Support Forum Q 🌲 🌅 😁 Log-out
Collaboratory Collaboratory	comentation support Forum Q at the Log-out
Warning: Collaboratory 1 will be shutdown on Wednesday September 8th at 18-00 CEST (my timezone #). Please migrate your data before then. Read more.	
Collaborate. Create reproducible science. Discover EBRAINS services at work. From anywhere. Getting started	
Search word in all wilk pages Q	
Highlighted collabs View all	a shared a shared a shared a shared a shared a shared a shared a shared a shared a shared a s

- 2. To run jobs, you need to have your own collab; if you don't yet have one then to do this click on "Collabs" in the top-right menu, and then on the subsequent page click on the "Create a collab" button. On this page, you will be required to give your collab a Title, a Name, a Description, and declare whether it is private or public (note: making it public makes it easier for others to see and help with your work should you need it).
- 3. Once you have created your collab you will be able to edit the associated page appropriately. It will look something like the following image (with less menu items on the left-hand menu to start with).

	Collabs	Documentation	Support	Forum Q	A [() Log	-out
Warning: Collaboratory 1 will be shutdown on Wednesday Septem	nber 8th at 18:00 CEST (my timezone#). Please migrate your data before then. Read more.						
	# ~ / Collabs ~ / Test SpNNakar jobs ADG ~ Test SpiNNaker jobs ADG Last modified by Andrew Gait on 202100/01 12:18		♥ 0	Settings	ℰ Edit	+ Create	I
 Lab SpiNkaker Jupyter testing SpiNkaker testing Team 	Test SpiNNaker jobs A collab for running test jobs via the NMPI submission queue to old data from collab v1	SpiNNaker; a	Iso store	s			
	 What can I find here? The "SpiNNaker testing" item in the left-hand menu is what I use to access spiNNaker from this colab v2 (created using the "EBRAINS Neuromorphic Computing Service: Job Manager" Community App from the ~ Create' button). You can also access SpiNNaker utpref and the SBAINS account directly via our JupyterLab at http://spiNnaker_Jupyter setsing" item. Migrated data from my old collabs in collab v1 Whon bas accesss? Igenerally used this in v1 to check problems other users were having with submitting jobs to NMC. If anyone is interested in a SpiNNaker-taer feater date me would suggest starting arew one youres! faan dating user (SpiNNaker). 	Contents • Test SpiNi • What can • Who has a	I find here?				ihow tour

4. In this instance, "SpiNNaker testing" links to the portal submission process that we want to use, and "SpiNNaker Jupyter testing" links directly to a Jupyterlab that can be run using your EBRAINS account. To create a portal submission link in this menu, click on the "+ Create" button in the top right to get the following page:

		Collabs Documentation Support Forum Q 🌢 🏹 😕 Log-out
Collaboratory Colla	Second	Collabs Documentation Support Forum Q ▲ Q Image: Select the kind of page that you want to create Select the kind of page that you want to create Type to filter Default (1) Default (Nr page Standard empty page Templates (3) Article with table of content Image that table of content Image that a Community App Image that a Community App in your collab
		0.03980

5. Give your new page a Title, then highlight "Community App" and click "Create"

			Collabs Documentation St	upport Forum Q 🌲 <table-cell> 😁 Log-</table-cell>
Public Member	# = / Collabs = / Test SpiNNaker jobs ADG = / Te	st =		
est SpiNNaker jobs ADG	Test			♥0 ✔Edit +Create
rive ab piNNaker Jupyter testing piNNaker testing eam				
	PSP-Validation-Dev	Dummy App	**test**	wp6ws_messines_workshop
	Brain Simulation Platform	Collaboratory Basics	Development	Development
		A dummy app to demo how to integrate applications within the Collaboratory.	test	My collaboratory App
	More O Select	More O Select	More O Select	More O Select
	See.			1000 C
	Nutil	morph-viewer-test	Neuromorphic Computing Job Manager	E-mail Signature
	Neuroinformatics		Neuromorphic	Development
	NeuroImaging Utilities		This app provides a graphical user interface for remote access to the BrainScaleS and SpINNaker neuromorphic computing systems. It allows you to co	Simple e-mail signature generator
	More O Select	More O Select	More O Select	More O Select

6. On this Community App screen select the "Neuromorphic Computing Job Manager" and click "Save & View"

Collaboratory					Co	ollabs Do	cumentation	Support	Forum	a 🔺	e Log
Warning: Collaboratory 1 will be shutdown on Wednesday September	r 8th at 18:00 CEST	(my timezor	ne≌). Please migrat	e your data before I	hen. Read more.						
📽 Public 🔹 Member	n - / Collabs - /	Test SpiNNak	er jobs ADG = / SpiNN	laker testing ~							
 ✓ Test SpiNNaker jobs ADG Bucket Orive 	SpiNNa Last modified by An								•	0 🖋 Edit	+ Create
 Lab SpiNNaker Jupyter testing SpiNNaker testing Team 	EBRAINS Neu	uromorphic	c Computing Ser	vice: Job Manage	ər					oL	obs Quotas
	• C	ID	🛷 Status	System	<> Code			🕒 Submitte	d on	🕒 Subm	nitted by
	٩	161154	inished	SpiNNaker	# write your code here import spynnaker8 as sim i.		:	021/08/18 10	:31:11	andrewga	iit
	۹	161153	Ø finished	SpiNNaker	# write your code here import spynnaker8 as sim i.		:	2021/08/18 10	:25:16	andrewga	iit
	Last updated: 1	/9/2021 @ 14	4:42:40								

7. You should end up with something like this (without the completed jobs). In order to run some jobs using the NMC Job Manager, you will need to request a quota; click on Quotas (on the right of the black bar) and then "New Request" to get to

Warning: Collaboratory 1 will be shutdown on Wednesday Septembe	ar 8th at 18:00 CEST (my timezoned). Please migrate your data before then. Read more.	
	# - Cotilates - / Test SpRNkaker jobs ADG - / SpRNkaker testing - SpinNaker testing Last modified by Andrew Get on 2021/08/18 12:15	♥0 ≠Edit +Create I
	EBRAINS Neuromorphic Computing Service: Job Manager	X Jobs Quotas +
	The The A dark tet quote The • Not or related Full • Not or related Full description of project Full description of project Full description of project Full description is not required for full quoted Full description of project Full description is Not required for full quoted Safe CHANGES	
	Tags: test simple [+]	#1 Deport this document

where you should fill in the Title & Summary (& Description) accordingly. Once your quota has been approved it will show up when you again click "Quotas".

8. Once your quota has been approved you are now ready to start running jobs. Click on the white plus sign inside a blue circle at the top of the list of jobs, which will take you to the following page:

		Collabs	Documentation	Support I	Forum Q	• []	🕒 Log-out
Warning: Collaboratory 1 will be shutdown on Wednesday Septen	nber 8th at 18:00 CEST (my timezone #). Please migrate your data before then. Read more.						
∎² Public 🚳 Member	<pre># - / Collabs - / Test SpINNaker jobs ADG - / SpINNaker testing -</pre>						
 ✓ Test SpiNNaker jobs ADG Bucket Drive 	SpiNNaker testing Last modified by Andrew Gait on 2021/08/18 12:15				♥ 0	₽ Edit ◀	Create 1
 Lab SpiNNaker Jupyter testing SpiNNaker testing Team 	EBRAINS Neuromorphic Computing Service: Job Manager					Jobs	X Quotas +
- Team	New job Hardware Platform Hardware Prease choose a simulation platform Code EDTOR FROM GIT REPOSITORY OR ZIP ARCHIVE 1 # write your code here	FROM THE DF	RIVE		graphical MC	> DDEL BUILDER	
	Command Command Command Optional: exectly the path to the main Python script, with any command-free arguments. Hardware Configuration						

- 9. On this page, under "Hardware Platform" select "SpiNNaker".
- 10. There are now two ways in which you can submit a PyNN script; either by writing / pasting a script into the editor directly, or directly from a git repository or zip archive file.
- 11. If you submit a job via a git repository or archive, your script that you wish to run either needs to be called "run.py", or you need to specify it in the "Command" box along with any (optional) command line arguments.

 Drive 		
 Lab 		
 SpiNNaker Jupyter testing 		×
 SpiNNaker testing 		
 Team 	EBRAINS Neuromorphic Computing Service: Job Manager	Jobs Quotas +
	1 # write your code here	
	Command	
	Command:	
	Optional: specily the path to the main Python script, with any command-line arguments.	
	Optional: speciny me pain to the main Python script, with any command-line arguments.	
	Hardware Configuration	
	r Hardware config	
	Please type a JSON-formatted object. See the Guidebook for more details	
	Tags	
	idys _ top=	
	Tag1,Tag2;This is Tag3	
	Please type job tags, separated by a comma, or semicolon. Tags can have spaces.	
	CANCEL SUBMIT >	
	Tags: test simple (+)	Report this document
		to report this document
	Comments (0) Attachments (0) History Information	
	No comments for this page	
	Comment	

12. The "Hardware Configuration" box does not need to be filled in to use the SpiNNaker backend, but gives you some options as to which version of the software you would like to use and whether your script requires any additional installs; for more details see

https://electronicvisions.github.io/hbp-sp9-guidebook/using_the_platform.html#hardw are-platform-configuration

- 13. You can additionally put an optional number of tags on your job.
- 14. Once you are happy with what you have filled in on this page, you can click "Submit".
- 15. This will return you to the job list: your Job will start in the "submitted" stage, then "running" stage, then either "finished" or "error".

					Collabs	Documentation Support Forum	Q 🌲 🌅 😁 Log
Warning: Collaboratory 1 will be shutdown on Wednes	day September 8th at 18:00 CE	ST (my timezo	ne≓). Please migrat	e your data before th	en. Read more.		
🚽 Public 🔹 Member	# - / Collabs	+ / Test SpiNNa	ker jobs ADG - / SpiNN	laker testing ~			
Volue V			esting			•	0 🖋 Edit 🔸 Create
SpiNNaker Jupyter testing SpiNNaker testing Team	EBRAINS	√euromorphi	c Computing Ser	vice: Job Manage	r		Jobs Quotas
	• C	ID	√ Status	🚍 System	<> Code	() Submitted on	e Submitted by
	٩	161407	submitted	SpiNNaker	# write your code here import spynnaker8 as sim i	2021/09/01 13:59:39	andrewgait
	٩	161154	inished	SpiNNaker	<pre># write your code here import spynnaker8 as sim i # write your code here import spynnaker8 as sim i</pre>	2021/08/18 10:31:11 2021/08/18 10:25:16	andrewgait
	Last update	d: 1/9/2021 @ 1	4:59:39				

- 16. You are able to view more about your job at any stage by clicking on the magnifying glass symbol to the left of the row detailing your job. In particular, while the job is in the "running" state you can see the current state of the log at the bottom of the page.
- 17. Once your job has reached the finished state, you can then download various files associated with your job:

Collaboratory		Collabs	Documentation Support Forum Q	🌲 🔽 🖙 Log-out
Warning: Collaboratory 1 will be shutdown on Wednesday September	er 8th at 18:00 CEST (my timezone≅). Please migrate your data before then. Rea	ad more.		
	# - / Collabs - / Test SpiNNaker jobs ADG - / SpiNNaker testing - SpiNNaker testing Last modified by Andrew Galt on 2021/08/18 12:15		•0	Edit + Create I
e Team	EBRAINS Neuromorphic Computing Service: Job Manager Job 161407 Interes Submitted on 2021/09/01 13:59:39 by andrengalt to Spikhtaker Comeleted on 2021/09/01 14:03:43 Output files			Jobs Quotas +
	http://spinaker.cs.man.ac.uk/services/reti/output/test-spinaker/obs- adg/161407/output_3646536388262161624.txt Code	https://spinnaker.cs.man.ac.uk/services/cet/output/test spinnaker-jobs-adg/161407/reports.zip	https://spinnaker.cs.man.ac.uk/services/re jabs-adg/161407/simple_example.png	st/output/test-spinnaker-
	<pre># write your cade here import spynakerd as im import pMA.utility.plating as plat import matplatLib.pyplat as plt sim.set_number_of_neurons_per_core(sim.If_curr_exp, 100) pop_l = sim.Population() sim.If_curr_exp(), tabelurpop_l") imput = sim.Population() is no SpideSorderStyplicit_imstel[] imput_proj = sim.Projection(imput, pop_l, sim.OperOpdeSonector)</pre>	r(),		

Note that compared to a "normal" local run of a PyNN script, in order to plot anything using matplotlib you need to use plt.savefig(filename) rather than plt.show(). Anything plotted in this manner will be saved as one of the files you can download, along with a directory of provenance reporting on your job as well as the output log

file of your job. Congratulations, you have now run a PyNN script job via the EBRAINS portal.

2.3 Using the SpiNNaker JupyterLab via EBRAINS

1. It is possible to also use your collab to access SpiNNaker using Jupyter. You may find this an easier and more interactive way to test your PyNN scripts on the machine before running them using the NMC Job Manager described in the previous section. In order to do this, from your collab main page, click on "+ Create", then from the subsequent page, give what you are creating a "Title", select "Use Community App", click "Create", then on the CommunityApp screen find and highlight the "SpiNNaker Jupyter" app. Once highlighted, click "Save & View", and this should then log you directly into a SpiNNaker JupyterLab instance as shown in the following image:

Collaboratory		Collabs Documentation Support Forum Q 🌲 🌅 🏵 Log-out
Warning: Collaboratory 1 will be shutdown on Wednesday Septemb	er 8th at 18:00 CEST (my timezone 2). Please migrate your data before then, Read more.	
🔊 Public 🔹 Member	₩ + / Collabs + / Test SpiNNaker jobs ADG + / SpiNNaker Jupyter testing +	
 Test SpiNNaker jobs ADG Bucket Drive 	SpiNNaker Jupyter testing	♥0 / Edit +Create :
 Lab SpiNNaker Jupyter testing SpiNNaker testing 	🞽 File Edit View Run Kernel Tabs Settings Help	🖉 documentation d 💥
• Team	Clander Clander	
L	Simple 🛄 1 🔟 0 🛞	Launcher

For more information about our Jupyter setup see: <u>https://spinnakermanchester.github.io/latest/jupyter.html</u>

Note that you can also use the Jupyter Notebook instance as detailed in the above document by simply doing File - Log Out, then once you log in again from the subsequent screen it will give you the option of using the Notebook or Lab interfaces.