

# PyGeNN: A Python library for GPU-enhanced neural networks

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## 2 ABSTRACT

- 3 More than half of the Top 10 supercomputing sites worldwide use GPU accelerators and they
- 4 are becoming ubiquitous in workstations and edge computing devices. GeNN is a C++ library for
- 5 generating efficient spiking neural network simulation code for GPUs. However, until now, the full
- 6 flexibility of GeNN could only be harnessed by writing model descriptions and simulation code in
- 7 C++. Here we present PyGeNN, a Python package which exposes all of GeNN's functionality to
- 8 Python with minimal overhead. This provides an alternative, arguably more user-friendly, way
- 9 of using GeNN and allows modellers to use GeNN within the growing Python-based machine
- 10 learning and computational neuroscience ecosystems. In addition, we demonstrate that, in both
- 11 Python and C++ GeNN simulations, the overheads of recording spiking data can strongly affect
- 12 runtimes and show how a new spike recording system can reduce these overheads by up to
- Tartained and show now a new spine recording system dan reduce these overneads by up to
- $10 \times 10 \times 10$ . Using the new recording system, we demonstrate that by using PyGeNN on a modern GPU, we can simulate a full-scale model of a cortical column faster even than real-time neuromorphic
- we dan simulate a fair seale model of a control column laster even than real time neuromorphic
- 15 systems. Finally, we show that long simulations of a smaller model with complex stimuli and a
- 16 custom three-factor learning rule defined in PyGeNN can be simulated up to  $72\times$  faster than
- 17 real-time.
- 18 Keywords: GPU, high-performance computing, parallel computing, benchmarking, computational neuroscience, spiking neural
- 19 networks, Python

## 1 INTRODUCTION

- 20 A wide range of spiking neural network (SNN) simulators are available, each with their own application
- 21 domains. NEST (Gewaltig and Diesmann, 2007) is widely used for large-scale point neuron simulations
- 22 on distributed computing systems; NEURON (Carnevale and Hines, 2006) and Arbor (Akar et al., 2019)
- 23 specialise in the simulation of complex multi-compartmental models; NeuroKernel (Givon and Lazar, 2016)
- 24 is focused on emulating fly brain circuits using Graphics Processing Units (GPUs); and CARLsim (Chou
- et al., 2018), ANNarchy (Vitay et al., 2015), NeuronGPU (Golosio et al., 2020) and GeNN (Yavuz et al.,
- 26 2016) use GPUs to accelerate point neuron models. For performance reasons, many of these simulators are
- 27 written in C++ and, especially amongst the older simulators, users describe their models either using a
- 28 Domain-Specific Language (DSL) or directly in C++. For programming language purists, a DSL may be an

elegant way of describing an SNN network model and, for simulator developers, not having to add bindings 30 to another language is convenient. However, both choices act as a barrier to potential users. Therefore, with both the computational neuroscience and machine learning communities gradually coalescing towards a 31 Python-based ecosystem with a wealth of mature libraries for scientific computing (Hunter, 2007; Van Der 32 Walt et al., 2011; Millman and Aivazis, 2011), exposing spiking neural network simulators to Python seems 33 a pragmatic choice. NEST (Eppler et al., 2009), NEURON (Hines et al., 2009) and CARLsim (Balaji et al., 34 2020) have all taken this route and now all offer Python interfaces. Furthermore, newer simulators such as 35 Arbor and Brian2 (Stimberg et al., 2019) have been designed from the ground up with a Python interface. 36

Our GeNN simulator can already be used as a backend for the Python-based Brian2 simulator (Stimberg 37 et al., 2019) using the Brian2GeNN interface (Stimberg et al., 2020) which modifies the C++ backend 38 "cpp\_standalone" of Brian 2 to generate C++ input files for GeNN. As for cpp\_standalone, initialisation of 39 simulations is mostly done in C++ on the CPU and recording data is saved into binary files and re-imported 40 into Python using Brian 2's native methods. While we have recently demonstrated some very competitive 41 performance results (Knight and Nowotny, 2018, 2020) using GeNN in C++, and through the Brian2GeNN 42 interface (Stimberg et al., 2020), GeNN could so far not be used directly from Python and it is not possible 43 to expose all of GeNN's unique features through the Brian2 API. Specifically, GeNN not only allows users 44 to easily define their own neuron and synapse models but, also 'snippets' for offloading the potentially 45 costly initialisation of model parameters and connectivity onto the GPU. Additionally, GeNN provides a 46 lot of freedom for users to integrate their own code into the simulation loop. In this paper we describe the 47 implementation of PyGeNN – a Python package which aims to expose the full range of GeNN functionality 48 with minimal performance overheads. Unlike in the majority of other SNN simulators PyGeNN allows 49 defining bespoke neuron and synapse models directly from Python without requiring users to extend the 50 underling C++ code. Below, we demonstrate the flexibility and performance of PyGeNN in two scenarios 51 where minimising performance overheads is particularly critical. 52

- In a simulation of a large, highly-connected model of a cortical microcircuit (Potjans and Diesmann, 2014) with small simulation timesteps. Here the cost of copying spike data off the GPU from a large number of neurons every timestep can become a bottleneck.
- In a simulation of a much smaller model of Pavlovian conditioning (Izhikevich, 2007) where learning 56 57 occurs over 1 h of biological time and stimuli are delivered – following a complex scheme – throughout the simulation. Here any overheads are multiplied by a large number of timesteps and copying stimuli 58 to the GPU can become a bottleneck. 59
- Using the facilities provided by PyGeNN, we show that both scenarios can be simulated from Python with only minimal overheads over a pure C++ implementation. 61

#### **MATERIALS AND METHODS** 2

#### 2.1 **GeNN** 62

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GeNN (Yavuz et al., 2016) is a library for generating CUDA (NVIDIA et al., 2020) code for the simulation of spiking neural network models. GeNN handles much of the complexity of using CUDA directly and 64 automatically performs device-specific optimizations so as to to maximize performance. GeNN consists of a main library – implementing the API used to define models as well as the generic parts of the code generator – and an additional library for each backend (currently there is a reference C++ backend for generating CPU code and a CUDA backend. An OpenCL backend is under development). Users describe their model by implementing a modelDefinition function within a C++ file. For example, a model consisting of 4 Izhikevich neurons with heterogeneous parameters, driven by a constant input current might be defined as follows:

```
void modelDefinition(ModelSpec &model)
72
73
74
       model.setDT(0.1);
75
       model.setName("izhikevich");
76
77
       NeuronModels::IzhikevichVariable::VarValues popInit(
78
           -65.0, -20.0, uninitialisedVar(), uninitialisedVar(),
79
           uninitialisedVar(), uninitialisedVar());
80
81
       model.addNeuronPopulation<NeuronModels::IzhikevichVariable>(
82
            "Pop", 4, {}, popInit);
83
84
       model.addCurrentSource<CurrentSourceModels::DC>(
85
           "CS", "Pop", {10.0}, {});
86 }
```

The *genn-buildmodel* command line tool is then used to compile this file; link it against the main GeNN library and the desired backend library; and finally run the resultant executable to generate the source code required to build a simulation dynamic library (a .dll file on Windows or a .so file on Linux and Mac). This dynamic library can then either be statically linked against a simulation loop provided by the user or dynamically loaded by the user's simulation code. To demonstrate this latter approach, the following example uses the **SharedLibraryModel** helper class supplied with GeNN to dynamically load the previously defined model, initialise the heterogenous neuron parameters and print each neuron's membrane voltage every timestep:

```
#include "sharedLibraryModel.h"
 95
 96
 97 int main()
 98 {
99
         SharedLibraryModel<float> model("./", "izhikevich");
100
        model.allocateMem();
101
        model.initialize();
102
         float *aPop = model.getScalar<float>("aPop");
103
         float *bPop = model.getScalar<float>("bPop");
         float *cPop = model.getScalar<float>("cPop");
104
105
         float *dPop = model.getScalar<float>("dPop");
         aPop[0] = 0.02; bPop[0] = 0.2; cPop[0] = -65.0; dPop[0] = 8.0;
106
107
         aPop[1] = 0.1; \quad bPop[1] = 0.2; \quad cPop[1] = -65.0; \quad dPop[1] = 2.0;
108
        aPop[2] = 0.02; bPop[2] = 0.2; cPop[2] = -50.0; dPop[2] = 2.0;
109
         aPop[3] = 0.02; bPop[3] = 0.2; cPop[3] = -55.0; dPop[3] = 4.0;
110
        model.initializeSparse();
111
112
         float *vPop = model.getScalar<float>("VPop");
113
        while (model.getTime() < 200.0f) {</pre>
114
             model.stepTime();
115
             model.pullVarFromDevice("Pop", "V");
```

```
116
             printf("%f, %f, %f, %f, %f\n",
117
                    t, VPop[0], VPop[1], VPop[2], VPop[3]);
118
         }
119
         return EXIT_SUCCESS;
120
```

#### **SWIG** 2.2

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In order to use GeNN from Python, both the model creation API and the SharedLibraryModel 122 functionality need to be 'wrapped' so they can be called from Python. While this is possible using 123 the API built into Python itself, wrapper functions would need to be manually implemented for each GeNN 124 function to be exposed which would result in a lot of maintenance overhead. Instead, we chose to use 125 SWIG (Beazley, 1996) to automatically generate wrapper functions and classes. SWIG generates Python 126 modules based on special interface files which can directly include C++ code as well as special 'directives' 127 which control SWIG. For example, the following SWIG interface file would wrap the C++ code in test.h in 128 a Python module called test\_module within a Python packages called test\_package: 129

```
130
    %module(package="test_package") test_module
131
    %include "test.h"
```

The module directive sets the name of the generated module and the package it will be located in and 132 the **%include** directive parses and automatically generates wrapper functions for the C++ header file. We 133 134 use SWIG in this manner to wrap both the model building and SharedLibraryModel APIs described in section 2.1. However, key parts of GeNN's API such as the ModelSpec::addNeuronPopulation method 135 employed in section 2.1, rely on C++ templates which are not directly translatable to Python. Instead, valid 136 137 template instantiations need to be given a unique name in Python using the \*template SWIG directive:

```
%template(addNeuronPopulationLIF) ModelSpec::addNeuronPopulation<NeuronModels::LIF>;
```

Having to manually add these directives whenever a model is added to GeNN would be exactly the sort 139 of maintenance overhead we were trying to avoid by using SWIG. Therefore, when building the Python 140 wrapper, we instead search the GeNN header files for the macros used to declare models in C++ and 141 automatically generate SWIG %template directives. 142

As previously discussed, a key feature of GeNN is the ease with which it allows users to define their own neuron and synapse models as well as 'snippets' defining how variables and connectivity should be initialised. Beneath the syntactic sugar described in our previous work (Knight and Nowotny, 2018), new models can be defined in C++ by defining a new class derived from, for example, the NeuronModels::Base 146 class. The ability to extend this system to Python was a key requirement of PyGeNN and, by using SWIG 'director', C++ classes can be made inheritable from Python using a single SWIG directive:

149 %feature("director") NeuronModels::Base;

#### 2.3 PyGeNN 150

While GeNN *could* be used from Python via the wrapper generated using SWIG, the resultant code 151 would be unpleasant to use directly. For example, rather than being able to specify neuron parameters 152 using a native Python types such as lists or dictionaries, one would have to use a wrapped type such as 153 DoubleVector([0.25, 10.0, 0.0, 0.0, 20.0, 2.0, 0.5]). Therefore, in order to provide a more 154 user-friendly and pythonic interface, we have built PyGeNN on top of the wrapper generated by SWIG. 155

PyGeNN combines the separate model building and simulation stages of building a GeNN model in C++ into a single API, likely to be more familiar to users of existing Python-based model description 157 languages such as PyNEST (Eppler et al., 2009) or PyNN (Davison et al., 2008). By combining the two 158 159 stages together, PyGeNN can provide a unified dictionary-based API for initialising homogeneous and heterogeneous parameters as shown in this re-implementation of the previous example: 160

```
161
    from pygenn import genn_wrapper, genn_model
162
163
    model = genn_model.GeNNModel("float", "izhikevich")
164
    model.dT = 0.1
165
    izk_init = {"V": -65.0,
166
167
                 "U": -20.0,
168
                 "a": [ 0.02,
                                 0.1,
                                        0.02,
                                                0.02],
169
                 "b": [ 0.2,
                                 0.2,
                                        0.2,
                                                0.21,
                 "c": [-65.0, -65.0, -50.0, -55.0],
170
171
                 "d": [ 8.0,
                                 2.0,
                                        2.0,
172
173
    pop = model.add_neuron_population("Pop", 4, "IzhikevichVariable",
                                        {}, izk_init)
174
175
    model.add_current_source("CS", "DC", "Pop",
176
                               {"amp": 10.0}, {})
177
178 model.build()
179
    model.load()
180
181
    v = pop.vars["V"].view
182
    while model.t < 200.0:
183
        model.step_time()
184
        model.pull_state_from_device("Pop")
185
        print("%f, %f, %f, %f, %f"
                % (model.t, v[0], v[1], v[2], v[3]))
186
```

Initialisation of variables with homogeneous values – such as the neurons' membrane potential – is 187 188 performed by initialisation kernels generated by GeNN and initialisation of variables with heterogeneous 189 values – such as the a, b and c parameters – are copied to the GPU by PyGeNN after the model is loaded. While the PyGeNN API is more pythonic and, hopefully, more user-friendly than the C++ interface, it 190 still provides users with the same low-level control over the simulation. Furthermore, by using SWIG's 191 numpy (Van Der Walt et al., 2011) interface, the host memory allocated by GeNN can be accessed directly 192 from Python using the pop.vars["V"].view syntax meaning that no potentially expensive additional 193 copying of data is required. 194

195 As illustrated in the previously-defined model, for convenience, PyGeNN allows users to access GeNN's built-in models. However, one of PyGeNN's most powerful features is that it enables users to easily 196 define their own neuron and synapse models from within Python. For example, an Izhikevich neuron 197 model (Izhikevich, 2003) can be defined using the create\_custom\_neuron\_class helper function which 198 provides some syntactic sugar over directly inheriting from the SWIG director class:

```
200
    izk_model = genn_model.create_custom_neuron_class(
201
         "izk",
```

199

```
202
         param_names=["a", "b", "c", "d"],
203
         var_name_types=[("V", "scalar"), ("U", "scalar")],
204
         sim code=
             .....
205
206
             (V) += 0.5*(0.04*(V)*(V)+5.0*(V)+140.0-(U)+(Isyn))*DT;
207
             (V) += 0.5*(0.04*(V)*(V)+5.0*(V)+140.0-(U)+(Isyn))*DT;
             (U) += (a) * (b) * (V) - (U) * DT;
208
209
210
         threshold_condition_code="$(V) >= 30.0",
211
         reset_code=
             11 11 11
212
213
             (V) = (c);
214
             $(U) += $(d);
215
```

The param\_names list defines the real-valued parameters that are constant across the whole population of neurons and the var\_name\_types list defines the model state variables and their type (the scalar type is an alias for either single or double-precision floating point, depending on the precision passed to the Gennmodel constructor). The behaviour of the model is then defined using a number of code strings. Unlike in tools like Brian 2 (Stimberg et al., 2019), these code strings are specified in a C-like language rather than using differential equations. This allows expert users to choose their own solver for models described in terms of differential equations and to programatically define models such as spike sources. For example, in the model presented above, we chose to implement the neuron using the idiosyncratic forward Euler integration scheme employed by Izhikevich (2003). Finally, the threshold\_condition\_code expression defines when the neuron will spike whereas the reset\_code code string defines how the state variables should be reset after a spike.

# 2.4 Spike recording system

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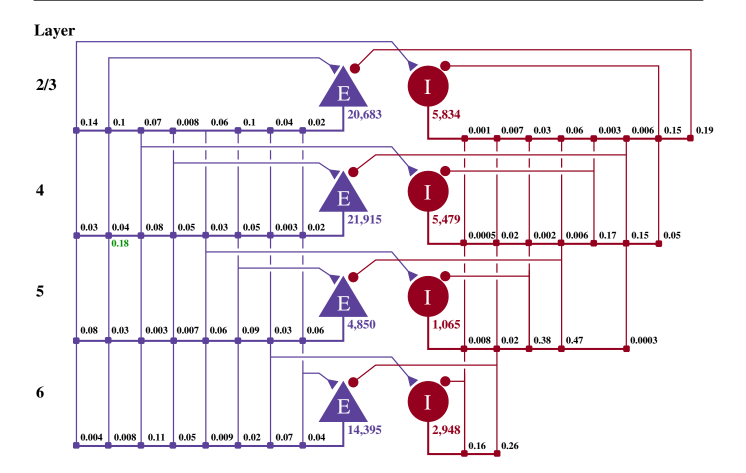
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Internally, GeNN stores the spikes emitted by a neuron population during one simulation timestep in an array containing the indices of the neurons that spiked alongside a counter of how many spikes have been emitted overall. Previously, recording spikes in GeNN was very similar to the recording of voltages shown in the previous example code – the array of neuron indices was simply copied from the GPU to the CPU every timestep. However, especially when simulating models with a small simulation timestep, such frequent synchronization between the CPU and GPU is costly – especially if a slower, interpreted language such as Python is involved. Furthermore, biological neurons typically spike at a low rate (in the cortex, the average firing rate is only around 3 Hz (Buzsáki and Mizuseki, 2014)) meaning that the amount of spike data transferred every timestep is typically very small. To address both of these sources of inefficiency, we have added a new data structure to GeNN which stores spike data for many timesteps on the GPU. To reduce the memory required for this data structure and to make its size independent of neural activity, the spikes emitted by a population of N neurons in a single simulation timestep are stored in a Nbit bitfield where a '1' represents a spike and a '0' the absence of one. Spiking data over multiple timesteps is then represented by a circular buffer of these bitfields. Using this approach, even the spiking output of relatively large models, running for many timesteps can be stored in a small amount of memory. For example, the spiking output of a model with  $100 \times 10^3$  neurons running for  $10 \times 10^3$  simulation timesteps, required less than 120 MB – a small fraction of the memory on a modern GPU. While efficiently handling spikes stored in a bitfield is a little trickier than working with a list of neuron indices, GeNN provides an efficient



**Figure 1.** Illustration of the microcircuit model. Blue triangles represent excitatory populations, red circles represent inhibitory populations and the number beneath each symbol shows the number of neurons in each population. Connection probabilities are shown in small bold numbers at the appropriate point in the connection matrix. All excitatory synaptic weights are normally distributed with a mean of  $0.0878\,\mathrm{nA}$  (unless otherwise indicated in green) and a standard deviation of  $0.008\,78\,\mathrm{nA}$ . All inhibitory synaptic weights are normally distributed with a mean of  $0.3512\,\mathrm{nA}$  and a standard deviation of  $0.03512\,\mathrm{nA}$ .

C++ helper function for saving the spikes stored in a bitfield to a text file and a numpy-based method for decoding them in PyGeNN.

## 248 2.5 Cortical microcircuit model

Potjans and Diesmann (2014) developed the cortical microcircuit model of  $1 \,\mathrm{mm}^3$  of early-sensory cortex illustrated in figure 1. The model consists of 77 169 LIF neurons, divided into separate populations representing the excitatory and inhibitory population in each of 4 cortical layers (2/3, 4, 5 and 6). The membrane voltage  $V_i$  of each neuron i is modelled as:

$$\tau_{\rm m} \frac{dV_i}{dt} = (V_{\rm rest} - V_i) + R_{\rm m} (I_{\rm syn_i} + I_{\rm ext_i}), \tag{1}$$

where  $\tau_{\rm m}=10\,{\rm ms}$  and  $R_{\rm m}=40\,{\rm M}\Omega$  represent the time constant and resistance of the neuron's cell membrane,  $V_{\rm rest}=-65\,{\rm mV}$  defines the resting potential,  $I_{\rm syn_i}$  represents the synaptic input current and  $I_{\rm ext_i}$  represents an external input current. When the membrane voltage crosses a threshold  $V_{\rm th}=-50\,{\rm mV}$  a spike is emitted, the membrane voltage is reset to  $V_{\rm rest}$  and updating of V is suspended for a refractory period  $\tau_{\rm ref}=2\,{\rm ms}$ . Neurons in each population are connected randomly with numbers of synapses derived

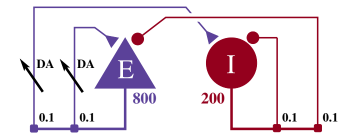


Figure 2. Illustration of the balanced random network model. The blue triangle represents the excitatory population, the red circle represents the inhibitory population, and the numbers beneath each symbol show the number of neurons in each population. Connection probabilities are shown in small bold numbers at the appropriate point in the connection matrix. All excitatory synaptic weights are plastic and initialised to 1 and all inhibitory synaptic weights are initialised to -1.

from an extensive review of the anatomical literature. These synapses are current-based, i.e. presynaptic spikes lead to exponentially-decaying input currents  $I_{\text{syn}_i}$ 

$$\tau_{\text{syn}} \frac{dI_{\text{syn}_i}}{dt} = -I_{\text{syn}_i} + \sum_{j=0}^n w_{ij} \sum_{t_j} \delta(t - t_j), \tag{2}$$

where  $\tau_{\rm syn}=0.5~{\rm ms}$  represents the synaptic time constant,  $w_{ij}$  represents the synaptic weight and  $t_j$  are the arrival times of incoming spikes from n presynaptic neurons. Within each synaptic projection, all synaptic strengths and transmission delays are normally distributed using the parameters presented in Potjans and Diesmann (2014, table 5) and, in total, the model has approximately  $0.3\times10^9$  synapses. As well as receiving synaptic input, each neuron in the network also receives an independent Poisson input current, representing input from neighbouring not explicitly modelled cortical regions. The Poisson input is delivered to each neuron via  $I_{\rm ext_i}$  with

$$\tau_{\text{syn}} \frac{dI_{\text{ext}_i}}{dt} = -I_{\text{ext}_i} + w_{\text{ext}} \text{Poisson}(\nu_{\text{ext}} \Delta t), \tag{3}$$

where  $\nu_{\rm ext}$  represents the mean input rate and  $w_{\rm ext}$  represents the weight. The ordinary differential Eq. 1, (2) and (3) are solved with an exponential Euler algorithm. For a full description of the model parameters, please refer to Potjans and Diesmann (2014, tables 4 and 5) and for a description of the strategies used by GeNN to parallelise the initialisation and subsequent simulation of this network, please refer to Knight and Nowotny (2018, section 2.3). This model requires simulation using a relatively small timestep of  $0.1 \, \mathrm{ms}$ , making the overheads of copying spikes from the GPU every timestep particularly problematic.

# 2.6 Pavlovian conditioning model

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The cortical microcircuit model described in the previous section is ideal for exploring the performance of short simulations of relatively large models. However, the performance of longer simulations of smaller models is equally vital. Such models can be particularly troublesome for GPU simulation as, not only might they not offer enough parallelism to fully occupy the device but, each timestep can be simulated so quickly that the overheads of launching kernels etc can dominate. Additional overheads can be incurred when models require injecting external stimuli throughout the simulation. Longer simulations are particularly useful when exploring synaptic plasticity so, to explore the performance of PyGeNN in this scenario, we

simulate a model of Pavlovian conditioning using a three-factor Spike-Timing-Dependent Plasticity (STDP) learning rule (Izhikevich, 2007).

## 266 2.6.1 Neuron model

The model illustrated in figure 2 consists of an 800 neuron excitatory population and a 200 neuron inhibitory population, within which, each neuron i is modelled using the Izhikevich model (Izhikevich, 2003) whose dimensionless membrane voltage  $V_i$  and adaption variables  $U_i$  evolve such that:

$$\frac{dV_i}{dt} = 0.04V_i^2 + 5V_i + 140 - U_i + I_{\text{syn}_i} + I_{\text{ext}_i}$$
 (4)

$$\frac{dU_i}{dt} = a(bV_i - U_i) \tag{5}$$

When the membrane voltage rises above 30, a spike is emitted and  $V_i$  is reset to c and d is added to  $U_i$ . Excitatory neurons use the regular-spiking parameters (Izhikevich, 2003) where a=0.02, b=0.2, c=-65.0, d=8.0 and inhibitory neurons use the fast-spiking parameters (Izhikevich, 2003) where a=0.1, b=0.2, c=-65.0, d=2.0. Again,  $I_{\text{syn}_i}$  represents the synaptic input current and  $I_{\text{ext}_i}$  represents an external input current. While there are numerous ways to solve Eq. 4 and 5 (Humphries and Gurney, 2007; Hopkins and Furber, 2015; Pauli et al., 2018), we chose to use the idiosyncratic forward Euler integration scheme employed by Izhikevich (2003) in the original work (Izhikevich, 2007). Under this scheme, Eq. 4 is first integrated for two 0.5 ms timesteps and then, based on the updated value of  $V_i$ , Eq. 5 is integrated for a single 1 ms timestep.

# 276 2.6.2 Synapse models

The excitatory and inhibitory neural populations are connected recurrently, as shown in figure 1, with instantaneous current-based synapses:

$$I_{\operatorname{syn}_{i}}(t) = \sum_{j=0}^{n} w_{ij} \sum_{t_{j}} \delta(t - t_{j}), \tag{6}$$

where  $t_j$  are the arrival times of incoming spikes from n presynaptic neurons. Inhibitory synapses are static with  $w_{ij} = -1.0$  and excitatory synapses are plastic. Each plastic synapse has an eligibility trace  $C_{ij}$  as well as a synaptic weight  $w_{ij}$  and these evolve according to a three-factor STDP learning rule (Izhikevich, 2007):

$$\frac{dC_{ij}}{dt} = -\frac{C_{ij}}{\tau_c} + \text{STDP}(\Delta t)\delta(t - t_{\text{pre/post}})$$
(7)

$$\frac{dw_{ij}}{dt} = -C_{ij}D_j \tag{8}$$

where  $\tau_c=1000\,\mathrm{ms}$  represents the decay time constant of the eligibility trace and  $STDP(\Delta t)$  describes the magnitude of changes made to the eligibility trace in response to the relative timing of a pair of pre and postsynaptic spikes with temporal difference  $\Delta t=t_{post}-t_{pre}$ . These changes are only applied to the trace at the times of pre and postsynaptic spikes as indicated by the Dirac delta function  $\delta(t-t_{pre/post})$ . Here, a

double exponential STDP kernel is employed such that:

$$STDP(\Delta t) = \begin{cases} A_{+} \exp\left(-\frac{\Delta t}{\tau_{+}}\right) & \text{if } \Delta t > 0\\ A_{-} \exp\left(\frac{\Delta t}{\tau_{-}}\right) & \text{if } \Delta t < 0\\ 0 & \text{otherwise} \end{cases}$$
 (9)

where the time constant of the STDP window  $\tau_+ = \tau_- = 20 \,\mathrm{ms}$  and the strength of potentiation and depression are  $A_+ = 0.1$  and  $A_- = 0.15$  respectively. Finally, each excitatory neuron has an additional variable  $D_j$  which describes extracellular dopamine concentration:

$$\frac{D_j}{t} = -\frac{D_j}{\tau_d} + \text{DA}(t) \tag{10}$$

where  $\tau_d = 200 \, \mathrm{ms}$  represents the time constant of dopamine uptake and DA(t) the dopamine input over time.

# 279 2.6.3 PyGeNN implementation of three-factor STDP

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282 283 The first step in implementing this learning rule in PyGeNN is to implement the STDP updates and decay of  $C_{ij}$  using GeNN's event-driven plasticity system, the implementation of which was described in our previous work (Knight and Nowotny, 2018). Using a similar syntax to that described in section 2.3, we first create a new 'weight update model' with the learning rule parameters and the  $w_{ij}$  and  $C_{ij}$  state variables:

We then instruct GeNN to record the times of current and previous pre and postsynaptic spikes. The current spike time will equal the current time if a spike of this sort is being processed in the current timestep whereas the previous spike time only tracks spikes which have occurred *before* the current timestep:

```
is_pre_spike_time_required=True,
is_post_spike_time_required=True,
295
296
is_prev_pre_spike_time_required=True,
297
is_prev_post_spike_time_required=True,
```

Next we define the 'sim code' which is called whenever presynaptic spikes arrive at the synapse. This code first implements Eq. 6 – adding the synaptic weight  $(w_{ij})$  to the postsynaptic neuron's input  $(I_{\text{syn}_i})$  using the (addToInSyn, x) function.

Within the sim code we also need to calculate the time that has elapsed since the last update of  $C_{ij}$  using the spike times we previously requested that GeNN record. Within a timestep, GeNN processes presynaptic

spikes before postsynaptic spikes so the time of the last update to  $C_{ij}$  will be the latest time either type of 306 spike was processed in previous timesteps: 307

```
308
             const scalar tc = fmax($(prev_sT_pre),
309
                                     $(prev_sT_post));
```

Using this time, we can now calculate how much to decay  $C_{ij}$  using the closed-form solution to Eq. 7: 310

```
311
             const scalar tagDecay = exp(-($(t) - tc) / $(tauC));
312
             scalar newTag = $(c) * tagDecay;
```

To complete the sim code we calculate the depression case of Eq. 9 (here we use the *current* postsynaptic 313 spike time as, if a postsynaptic and presynaptic spike occur in the same timestep, there should be no 315 update).

```
316
             const scalar dt = $(t) - $(sT_post);
317
             if (dt > 0) {
318
                 newTag -= ($(aMinus) * exp(-dt / $(tauMinus)));
319
320
             (c) = newTag;
321
             """,
```

Finally we define the 'learn post code' which is called whenever a postsynaptic spike arrives at the synapse. 322 323 Other than implementing the potentiation case of Eq. 9 and using the *current* presynaptic spike time when calculating the time since the last update of  $C_{ij}$  – in order to correctly handle presynaptic updates made in 324 the same timestep – this code is very similar to the sim code: 325

```
326
         learn_post_code=
327
328
             const scalar tc = fmax($(sT_pre),
329
                                      $ (prev_sT_post));
330
331
             const scalar tagDecay = exp(-($(t) - tc) / $(tauC));
332
             scalar newTag = $(c) * tagDecay;
333
334
             const scalar dt = $(t) - $(sT_pre);
335
             if (dt > 0) {
                 newTag += ($(aPlus) * exp(-dt / $(tauPlus)));
336
337
             }
338
             (c) = newTag;
339
             """)
```

Adding the synaptic weight  $w_{ij}$  update described by Eq. 8 requires two further additions to the model. 340 As well as the pre and postsynaptic spikes, the weight update model needs to receive events whenever 341 342 dopamine is injected via DA. GeNN supports such events via the 'spike-like event' system which allows 343 events to be triggered based on an expression evaluated on the presynaptic neuron. In this case, this expression simply tests an injectDopamine flag which gets set by the dopamine injection logic in our presynaptic neuron model:

event threshold condition code="injectDopamine",

344

345

346

In order to extend our event-driven update of  $C_{ij}$  to include spike-like events we need to instruct GeNN to record the times at which they occur:

```
is_pre_spike_event_time_required=True,
is_prev_pre_spike_event_time_required=True,
```

351 The spike-like events can now be handled using a final 'event code' string:

After updating the previously defined calculations of tc in the sim code and learn post code in the same way to also include the times of spike-like events, all that remains is to update  $w_{ij}$ . Mikaitis et al. (2018) showed how Eq. 8 could be solved algebraically, allowing  $w_{ij}$  to be updated in an event-driven manner with:

$$\Delta w_{ij} = \frac{C(t_c^{last})D(t_d^{last})}{-\left(\frac{1}{\tau_c} + \frac{1}{\tau_d}\right)} \left(e^{-\frac{t - t_c^{last}}{\tau_c}}e^{-\frac{t - t_d^{last}}{\tau_d}} - e^{-\frac{t_w^{last} - t_c^{last}}{\tau_c}}e^{-\frac{t_w^{last} - t_d^{last}}{\tau_d}}\right)$$
(11)

where  $t_c^{last}$ ,  $t_w^{last}$  and  $t_d^{last}$  represent the last times at which  $C_{ij}$ ,  $W_{ij}$  and  $D_j$  respectively were updated. Because we will always update  $w_{ij}$  and  $C_{ij}$  together when presynaptic, postsynaptic and spike-like events occur,  $t_c^{last} = t_w^{last}$  and Eq. 11 can be simplified to:

$$\Delta w_{ij} = \frac{C(t_c^{last})D(t_d^{last})}{-\left(\frac{1}{\tau_c} + \frac{1}{\tau_d}\right)} \left(e^{-\frac{t_c t_c^{last}}{\tau_c}}e^{-\frac{t_c t_d^{last}}{\tau_d}} - e^{-\frac{t_c^{last}}{\tau_d}}\right)$$
(12)

and this update can now be added to each of our three event handling code strings to complete the implementation of the learning rule.

# 360 2.6.4 PyGeNN implementation of Pavlovian conditioning experiment

To perform the Pavlovian conditioning experiment described by Izhikevich (2007) using this model, we chose 100 random groups of 50 neurons (each representing stimuli  $S_1...S_{100}$ ) from amongst the two neural populations. Stimuli are presented to the network in a random order, separated by intervals sampled from U(100,300)ms. The neurons associated with an active stimulus are stimulated for a single 1 ms simulation timestep with a current of 40.0 nA, in addition to the random background current of U(-6.5,6.5)nA, delivered to each neuron via  $I_{\text{ext}_i}$  throughout the simulation.  $S_1$  is arbitrarily chosen as the Conditional Stimuli (CS) and, whenever this stimuli is presented, a reward in the form of an increase in dopamine is delivered by setting DA(t) = 0.5 after a delay sampled from U(0,1000)ms. This delay period is large enough to allow a few irrelevant stimuli to be presented which act as distractors. The simplest way to implement this stimulation regime is to add a current source to the excitatory and inhibitory neuron populations which adds the uniformly-distributed input current to an externally-controllable per-neuron current. In PyGeNN, the following model can be defined to do just that:

```
373 stim noise model = create custom current source class(
```

361

362

363

364

365

366

367

368

369

370

371

372

```
374
        "stim_noise",
375
        param_names=["n"],
376
        var_name_types=[("iExt", "scalar", VarAccess_READ_ONLY)],
377
        injection_code=
378
379
             $(injectCurrent, $(iExt) + ($(gennrand_uniform) * $(n) * 2.0) - $(n));
380
```

where the n parameter sets the magnitude of the background noise, the \$(injectCurrent, I) function injects a current of InA into the neuron and  $(gennrand\_uniform)$  samples from U(0,1) using the 382 'XORWOW' pseudo-random number generator provided by cuRAND (NVIDIA Corporation, 2019). Once 383 a current source population using this model has been instantiated and a memory view to iExt obtained 384 in the manner described in section 2.3, in timesteps when stimulus injection is required, current can be 385 injected into the list of neurons contained in stimuli input set with: 386

```
387
    curr_ext_view[stimuli_input_set] = 40.0
388 curr_pop.push_var_to_device("iExt")
```

381

The same approach can then be used to zero the current afterwards. However, as almost 20 000 stimuli will 389 be injected over the course of a 1 h simulation, we can offload the stimulus delivery entirely to the GPU in 390 391 order to reduce potential overheads, using the following slightly more complex model:

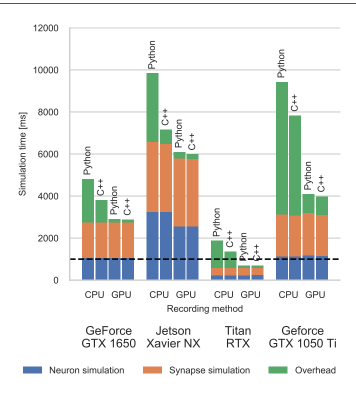
```
392
    stim_noise_model = create_custom_current_source_class(
393
        "stim_noise",
394
        param_names=["n", "stimMagnitude"],
        var_name_types=[("startStim", "unsigned int"),
395
396
                          ("endStim", "unsigned int", VarAccess_READ_ONLY)],
397
        extra_global_params=[("stimTimes", "scalar*")],
398
        injection_code=
             0.00
399
400
             scalar current = ($(gennrand_uniform) * $(n) * 2.0) - $(n);
401
             if($(startStim) != $(endStim) && $(t) >= $(stimTimes)[$(startStim)]) {
402
                current += $(stimMagnitude);
403
                $ (startStim) ++;
404
             }
405
             $(injectCurrent, current);
             """)
406
```

This model retains the same logic for generating background noise but adds a stimTimes array which contains the times at which each neuron should have current injected. This array is an 'extra global 408 parameter' – a read-only memory area that can be allocated and populated from PyGeNN, in this case by 409 'stacking' together a list of lists of spike times:

```
curr_pop.set_extra_global_param("stimTimes", np.hstack(neuron_stimuli_times))
```

412 The startStim and endStim variables are then used to point to the subset of the stimTimes array 413 corresponding to each neuron. Once the simulation time (\$(t)) passes the time at stimTimes[startStim],

414 current is injected and startStim is advanced.



**Figure 3.** Simulation times of the microcircuit model running on various GPU hardware for 1s of biological time. 'Overhead' refers to time spent in simulation loop but not within CUDA kernels. The dashed horizontal line indicates realtime performance

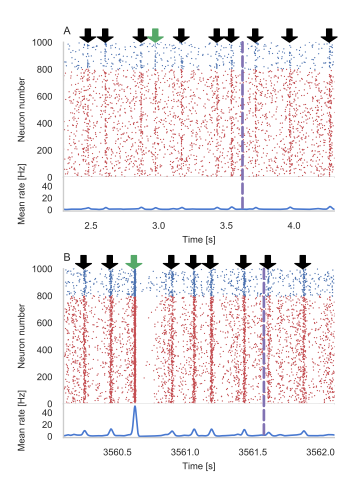
# 3 RESULTS

415 In the following subsections we will analyse the performance of the models introduced in 416 sections 2.5 and 2.6 on a representative selection of NVIDIA GPU hardware:

- Jetson Xavier NX a low-power embedded system with a GPU based on the Volta architecture with
   8 GB of shared memory.
- GeForce GTX 1050Ti a low-end desktop GPU based on the Pascal architecture with 4 GB of dedicated memory.
- GeForce GTX 1650 a low-end desktop GPU based on the Turing architecture with 4 GB of dedicated memory.
- Titan RTX a high-end workstation GPU based on the Turing architecture with 24 GB of dedicated memory.
- All of these systems run Ubuntu 18 apart from the system with the GeForce 1050 Ti which runs Windows 10.

# 427 3.1 Cortical microcircuit model performance

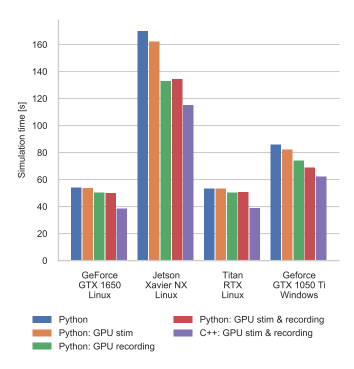
Figure 3 shows the simulation times for the full-scale microcircuit model. We measured the total simulation time by querying the std::chrono::high\_resolution\_clock in C++ and the time.perf\_counter in Python before and after the simulation loop; and used CUDA's own event timing system (NVIDIA Corporation, 2021, Section 3.2.6.7) to record the time taken by the neuron and synapse kernels. As one might predict, the Jetson Xavier NX is slower than the three desktop GPUs but, considering



**Figure 4.** Results of Pavlovian conditioning experiment. Raster plot and spike density function (SDF) (Szücs, 1998) showing activity centred around first delivery of Conditional Stimulus (CS) during initial (A) and final (B)  $50\,\mathrm{s}$  of simulation. Downward green arrows indicate times at which CS is delivered and downward black arrows indicate times when other, un-rewarded stimuli are delivered. Vertical dashed lines indicate times at which dopamine is delivered. The population SDF was calculated by convolving the spikes with a Gaussian kernel of  $\sigma=10\,\mathrm{ms}$  width.

that it only consumes a maximum of 15 W compared to 75 W or 320 W for the GeForce cards and Titan RTX respectively, it still performs impressively. The time taken to actually simulate the models ('Neuron simulation' and 'Synapse simulation') are the same when using Python and C++ as all GeNN optimisation options are exposed to PyGeNN. Interestingly, when simulating *this* model, the larger L1 cache and architectural improvements present in the Turing-based GTX 1650 do not result in significantly improved performance over the Pascal-based GTX 1050Ti. Instead, the slightly improved performance of the GTX 1650 can probably be explained by its additional 128 CUDA cores.

Without the recording system described in section 2.4, the CPU and GPU need to be synchronised after every timestep to allow spike data to be copied off the GPU and stored in a suitable data structure. The 'overheads' shown in figure 3 indicate the time taken by these processes as well as the unavoidable overheads of launching CUDA kernels etc. Because Python is an interpreted language, updating the spike data structures is somewhat slower and this is particularly noticeable on devices with a slower CPU such as the Jetson Xavier NX. However, unlike the desktop GPUs, the Jetson Xavier NX's 8 GB of memory is shared between the GPU and the CPU meaning that data does not need to be copied between their memories and can instead by accessed by both. While, using this shared memory for recording spikes



**Figure 5.** Simulation times of the Pavlovian Conditioning model running on various GPU hardware for 1 h of biological time. 'GPU stim' indicates simulations using the GPU stimulus delivery model and 'GPU recording' indicates simulations where the new recording system is employed.

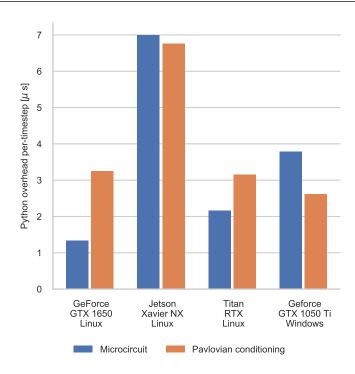
reduces the overhead of copying data off the device, because the GPU and CPU caches are not coherent, caching must be disabled on this memory which reduces the performance of the neuron kernel. Although the Windows machine has a relatively powerful CPU, the overheads measured in both the Python and C++ simulations run on this system are extremely large due to additional queuing between the application and the GPU driver caused by the Windows Display Driver Model (WDDM). When small – in this case 0.1 ms – simulation timesteps are used, this makes per-timestep synchronisation disproportionately expensive.

However, when the spike recording system described in section 2.4 is used, spike data is kept in GPU memory until the end of the simulation and overheads are reduced by up to  $10\times$ . Because synchronisation with the CPU is no longer required every timestep, simulations run approximately twice as fast on the Windows machine. Furthermore, on the high-end desktop GPU, the simulation now runs faster than real-time in both Python and native C++ versions – significantly faster than other recently published GPU simulators (Golosio et al., 2020) and even specialised neuromorphic systems (Rhodes et al., 2020).

# 3.2 Pavlovian conditioning performance

Figure 4 shows the results of an example simulation of the Pavlovian conditioning model. At the beginning of each simulation (Figure 4A), the neurons representing every stimulus respond equally. However, after 1 h of simulation, the response to the CS becomes much stronger (Figure 4B) – showing that these neurons have been selectively associated with the stimulus even in the presence of the distractors and the delayed reward.

In figure 5, we show the runtime performance of simulations of the Pavlovian conditioning model, running on the GPUs described above using PyGeNN with and without the recording system described in section 2.4 and the optimized stimulus-delivery described in section 2.6. These PyGeNN results are



**Figure 6.** Comparison of the duration of individual timestep in Python and C++ simulation in microcircuit and Pavlovian conditioning experiments. Times are taken from averages calculated over 5 runs of the fastest version of each model i.e. the microcircuit using the GPU recording system and the Pavlovian conditioning model using the GPU recording system and the GPU stimulus delivery.

compared to a C++ simulation using both optimizations. Because each simulation timestep only takes a few  $\mu$ s, the overhead of using CUDA timing events significantly alters the performance so, for this model, we only measure the duration of the simulation loop using the approaches described in the previous section. Although we only record the spiking activity during the first and last  $50\,\mathrm{s}$ , using the recording system still significantly improves the overall performance on all devices – especially on the Jetson Xavier NX with its slower CPU. Interestingly the Titan RTX and GTX 1650 perform identically in this benchmark with speedups ranging from  $62\times$  to  $72\times$  real-time. This is because, as discussed previously, this model is simply not large enough to fill the 4608 CUDA cores present on the Titan RTX. Therefore, as the two GPUs share the same Turing architecture and have very similar clock speeds (1350 MHz–1770 MHz for the Titan RTX and  $1485\,\mathrm{MHz}$ – $1665\,\mathrm{MHz}$  for the GTX 1650), the two GPUs perform very similarly. Similarly to the simulations of the microcircuit model, the Jetson Xavier NX performs rather slower than the desktop GPUs but still achieves speedups of up to  $31\times$ .

Interestingly, unlike in the simulations of the microcircuit model, here the GTX 1050 Ti performs rather differently. Although the clock speed of this device is approximately the same as the other GPUs (1290 MHz–1392 MHz) and it has a similar number of CUDA cores to the GTX 1650, its performance is significantly worse. The difference in performance across all configurations is likely to be due to architectural differences between the older Pascal; and newer Volta and Turing architectures. Specifically, Pascal GPUs have one type of Arithmetic Logic Unit (ALU) which handles both integer and floating point arithmetic whereas, the newer Volta and Turing architectures have equal numbers of dedicated integer and floating point ALUs as well as significantly larger L1 caches. As discussed in our previous work (Knight and Nowotny, 2018), these architectural features are particularly beneficial for SNN simulations with STDP where a large amount of floating point computation is required to update the synaptic state *and* 

491 additional integer arithmetic is required to calculate the indices into the sparse matrix data structures.

- 492 Furthermore, due to the additional synchronisation overheads caused by the Windows Display Driver
- 493 Model (WDDM) which we discussed in the previous section, offloading stimulus delivery to the GPU
- 494 improves the performance significantly on the Windows machine. However, on the other devices, it only
- 495 provides a minimal improvement.
- The difference between the speeds of the Python and C++ simulations of the Pavlovian conditioning
- 497 model (figure 5) appear much larger than those of the microcircuit model (figure 3). However, as figure 6
- 498 illustrates, for individual timesteps the excess time due to overheads is approximately the same for both
- 499 models and consistent with the cost of a small number of Python to C++ function calls (Apache Crail,
- 500 2019). Depending on the size and complexity of the model as well as the hardware used, this overhead may
- or may not be important. For example, when simulating the microcircuit model for 1 s on the Titan RTX,
- 502 the overhead of using Python is less than 0.2% but, when simulating the Pavlovian conditioning model on
- 503 the same device, the overhead of using Python is almost 31%.

# 4 DISCUSSION

- In this paper we have introduced PyGeNN, a Python interface to the C++ based GeNN library for GPU accelerated spiking neural network simulations.
- 303 accelerated spiking neural network simulations.
- 506 Uniquely, the new interface provides access to all the features of GeNN, without leaving the comparative
- 507 simplicity of Python and with, as we have shown, typically negligible overheads from the Python
- 508 bindings. PyGeNN also allows bespoke neuron and synapse models to be defined from within Python,
- 509 making PyGeNN much more flexible and broadly applicable than, for instance, the Python interface
- 510 to NEST (Eppler et al., 2009) or the PyNN model description language used to expose CARLsim to
- 511 Python (Balaji et al., 2020).
- In many ways, the new interface resembles elements of the Python-based Brian 2 simulator (Stimberg
- et al., 2019) (and it's Brian2GeNN backend (Stimberg et al., 2020)) with two key differences. Unlike in
- 514 Brian 2, bespoke models in PyGeNN are defined with 'C-like' code snippets. This has the advantage of
- 515 unparalleled flexibility for the expert user, but comes at the cost of more complexity as the code for a
- 516 timestep update needs to include a suitable solver as well as merely differential equations. The second
- 517 difference lies in how data structures are handled. Whereas simulations run using the C++ or Brian2GeNN
- 518 Brian 2 backends use files to exchange data with Python, the underlying GeNN data structures are directly
- 519 accessible from PyGeNN meaning that no disk access is involved.
- As we have demonstrated, the PyGeNN wrapper, exactly like native GeNN, can be used on a variety
- 521 of hardware from data centre scale down to mobile devices such as the NVIDIA Jetson. This allows for
- 522 the same codes to be used in large-scale brain simulations and embedded and embodied spiking neural
- 523 network research. Supporting the popular Python language in this interface makes this ecosystem available
- 524 to a wider audience of researchers in both Computational Neuroscience, bio-mimetic machine learning and
- 525 autonomous robotics.
- The new interface also opens up opportunities to support researchers that work with other Python based
- 527 systems. In the Computational Neuroscience and Neuromorphic computing communities, we can now build
- 528 a PyNN (Davison et al., 2008) interface on top of PyGeNN and, infact, a prototype of such an interface is
- 529 in development. Furthermore, for the burgeoning spike-based machine learning community, we can use
- 530 PyGeNN as the basis for a spike-based machine learning framework akin to TensorFlow or PyTorch for
- 531 rate-based models. A prototype interface of this sort called mlGeNN is in development and close to release.

- In this work we have introduced a new spike recording system for GeNN and have shown that, using
- 533 this system, we can now simulate the Potjans microcircuit model (Potjans and Diesmann, 2014) faster
- 534 than real-time and, to the best of our knowledge, faster than any other systems. Finally, the excellent
- 535 performance we have demonstrated using low-end Turing architecture GPUs is very exciting in terms of
- 536 increasing the accessibility of GPU accelerated Computational Neuroscience and SNN machine learning
- 537 research.

# **CONFLICT OF INTEREST STATEMENT**

- 538 The authors declare that the research was conducted in the absence of any commercial or financial
- 539 relationships that could be construed as a potential conflict of interest.

# **AUTHOR CONTRIBUTIONS**

- 540 JK and TN wrote the paper. TN is the original developer of GeNN. AK was the original developer of
- 541 PyGeNN. JK is currently the primary developer of both GeNN and PyGeNN and was responsible for
- 542 implementing the spike recording system. JK performed the experiments and the analysis of the results that
- 543 are presented in this work.

# **FUNDING**

- 544 This work was funded by the EPSRC (Brains on Board project, grant number EP/P006094/1), the European
- 545 Union's Horizon 2020 research and innovation program under Grant Agreement 945539 (HBP SGA3) and
- 546 a Google Summer of Code grant to AK.

# **ACKNOWLEDGMENTS**

- 547 We would like to thank Malin Sandström and everyone else at the International Neuroinformatics
- 548 Coordinating Facility (INCF) for their hard work running the Google Summer of Code mentoring
- 549 organisation every year. Without them, this and many other exciting Neuroinformatics projects would not
- 550 be possible.

## DATA AVAILABILITY STATEMENT

- 551 All models, data and analysis scripts used for this study can be found in https://github.com/
- 552 BrainsOnBoard/pygenn\_paper.

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