Building a Spiking Neural Network Model of the Basal Ganglia on SpiNNaker

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Abstract—We present a biologically-inspired and scalable model of the Basal Ganglia (BG) simulated on the SpiNNaker machine, a biologically-inspired low-power hardware platform allowing parallel, asynchronous computing. Our BG model consists of six cell populations, where the neuro-computational unit is a conductance-based Izhikevich spiking neuron; the number of neurons in each population is proportional to that reported in anatomical literature. This model is treated as a single-channel of action-selection in the BG, and is scaled-up to three channels with lateral cross-channel connections. When tested with two competing inputs, this three-channel model demonstrates action-selection behaviour. The SpiNNaker-based model is mapped exactly on to SpinML running on a conventional computer; both model responses show functional and qualitative similarity, thus validating the usability of SpiNNaker for simulating biologically-plausible networks. Furthermore, the SpiNNaker-based model simulates in real time for time-steps $\geq 1$ ms; power dissipated during model execution is $\approx 1.8$ W.

I. INTRODUCTION

The aim of this work is to build a biologically-inspired, scalable, spiking neural network model of the Basal Ganglia on the SpiNNaker machine [1]. The Basal Ganglia (BG) are a set of subcortical nuclei, which are evolutionarily very old and appear in all vertebrates, enabling them to make decisions and take subsequent actions [2]. The information on which the decision needs to be made, i.e., the environmental circumstance, constitutes the input to the BG, and is available via the Thalamus and Cortex. Output from the BG is the specific action that is decided upon, referred to as ‘action-selection’ [3], and is relayed to the motor pathway for execution via the Thalamus, Cortex and other sub-cortical structures. The objective of this work is to build a computational framework that will provide a ‘basic building block’ for further testing and development of automated decision-making tools in low-power, real time hardware such as the SpiNNaker machine [4], [5].

The neurotransmitter dopamine lies at the heart of the decision-making/action-selection functions of the BG. Seminal modelling work by Gurney et al [3], [6] introduces the concept of ‘selection-control’ pathways in the BG, a deviation from the more common nomenclature of ‘direct-indirect’ pathways associated with how dopamine controls and executes the action-selection mechanism. Subsequently, the model was also demonstrated as a computational tool to study brain disorders [7], [8], [9], as well as to form a conceptual understanding of the action-selection mechanism adopted by the BG and implemented in robots [10], [11].

The BG model presented in this work is based on the ‘selection-control’ BG model proposed by Gurney et al [3], and its later extension by Humphries et al [9]. This circuitry consists of six cell populations viz. the Subthalamic Nucleus (STN), Globus Pallidus externa (GPe), Substantia Nigra pars reticulata (SNr), and the Striatal Medium Spiny Neurons (Str-MSN). In addition, we have included the Striatal Fast Spiking Interneurons (Str-FSI) along the lines presented in [12] (sans the gap junction connections). Furthermore, our BG network model comprises several recurrent connections that are based on literature reporting anatomical data. The basic computational unit in our model is implemented using Izhikevich’s conductance-based spiking neurons, supported by SpiNNaker’s underlying software toolchain sPyNNaker [13]. Our choice of the single spiking neuron model is inspired by a similar implementation in the BG models presented by Liu et al [14] and Thibeault et al [15]. Inputs to the model are simulated with Poisson distributed spike trains generated separately for each simulation run, and are provided to the Str-MSN, Str-FSI, and STN cell populations. The response of the SNr cell population is the model output. At first, we built a macroscopic ‘channel’-like columnar model capturing neural information pathways in the BG, and parameterised this model to produce base firing rates as reported in [9], [15]. Next, to emulate arbitration by the BG of parallel macroscopic information channels representing competing sources, the single-channel model is used as a basic building block to scale up to three channels. Our results demonstrate selection of a competing action by the three-channel BG model, and are in agreement with previous model-based research [9], [15].

To compare and validate the SpiNNaker-based model outputs with those obtained using a conventional computer, we have implemented the same BG circuit using SpinML [16], an XML-based format for the specification of networks of point-neuron models. To create the SpinML-based model, we made use of SpineCreator, a graphical editor that is designed to provide an easy-to-use and flexible interface for building and visualising neuronal models [17], [18]. The model output dynamics show functional and qualitative similarity on both platforms i.e. SpiNNaker and SpinML, indicating the...
SpiNNaker machine as a viable platform for implementing spiking neural networks. A stringent bootstrapped t-test [19] (see Appendix) performed on the total number of spikes generated by each population over a period of 6 s shows that \( p < 0.05 \), implying statistically significant numerical differences between the model spike counts. This difference is due to the stochastic nature of the model inputs, replicating the numerical differences between data recorded from different animals for the same behavioural task, and is aligned with our expectations.

The underlying SpiNNaker architecture is designed to run in real time for time-steps \( \geq 1 \) ms. However, we solve the Izhikevich neuron models with a time-step of 0.1 ms to ensure solution accuracy. Thus, all simulations of the BG model on SpiNNaker ran in 10 s real time for 1 s simulation time, i.e. slowed down by a factor of 10. That said, a performance analysis indicates that the model is guaranteed to execute in this time, which lends a reliability factor e.g. for real time implementations. In addition, the run time is unaffected by scaling up the model, i.e. 10 s model simulation time is guaranteed to execute in 100 s real time for both single- and three-channel models. The power dissipation during model execution, measured using equipment built in-house [20], is \( \approx 0.8 \) W and 1.8 W for the single- and three-channel model respectively.

In Sect. II, we present the model design and implementation methods. In Sect. III, we present the model simulation methods and results. A comparison study with simulation of the same model on the SpineML platform running on a conventional computer is presented in Sect. IV-A; a performance analysis in terms of simulation time and power dissipation on SpiNNaker is presented in Sect. IV-B. We discuss the results and conclude the paper in Sect. V.

II. MODEL DESIGN AND IMPLEMENTATION

![Schematic diagram of the single-channel Basal Ganglia (BG) model. An overview of the biological basis of the model layout is provided in Sect. II-A. A pool of 25 Poisson distributed spike trains provide input to the Str-MSN and Str-FSI populations, while a separate pool consisting of 2 Poisson spike trains provide input to the STN population. Model output is the average firing rate of all neurons in the SNr population.](image)

Table I shows the total number of neurons in each population of the BG model and presents the proportional weightings of each model component. The numerical differences between data recorded from different animals are due to the stochastic nature of the model inputs, replicating the numerical differences between data recorded from different animals for the same behavioural task, and is aligned with our expectations.

### Table I

<table>
<thead>
<tr>
<th>Population</th>
<th>Total number of neurons (reported)</th>
<th>Total number of neurons (model)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STR (Str-MSN + Str-FSI) ((N_{str}))</td>
<td>( \times 2790 \times 10 )</td>
<td>( 2790 )</td>
</tr>
<tr>
<td>Str-MSN ((N_{msn}))</td>
<td>( 90% \times N_{str} )</td>
<td>( 2511 )</td>
</tr>
<tr>
<td>Str-MSN-D1/D2</td>
<td>( 50% \times N_{msn} )</td>
<td>( 1255 )</td>
</tr>
<tr>
<td>Str-FSI</td>
<td>( 3% \times N_{str} )</td>
<td>( 84 )</td>
</tr>
<tr>
<td>STN</td>
<td>( 13560 )</td>
<td>( 14 )</td>
</tr>
<tr>
<td>GPe</td>
<td>( 49960 )</td>
<td>( 46 )</td>
</tr>
<tr>
<td>SNr</td>
<td>( 26320 )</td>
<td>( 27 )</td>
</tr>
</tbody>
</table>

A. Biological background

The basic BG model circuitry simulated on SpiNNaker is shown in Fig. 1. The Striatum forms the main input structure of the BG and receives excitatory glutamatergic synapses from both the cortex and the thalamus. Studies on the BG cells of the rat brain [21] report that around 90 – 95% of the cells of the Striatum are of the Str-MSN variety. The remaining 5 – 10% of the cells constitute the interneurons of the Striatum. While there are three known varieties of interneurons, the predominant inhibitory influence on the Str-MSN is thought to be from the \( \gamma \)-aminobutyric acid (GABA)-ergic Str-FSI, which constitute around 2 – 5% of the cells of the Striatum [22], [23]. In this work, we model the Str-FSI population constituting around 3% of the cells of the Striatum.

A core feature of the BG is the modulation of population behaviour by dopamine released by the Substantia Nigra pars compacta (SNC; not modelled here). The Str-MSNs receive major dopaminergic input from the SNC and are modulated selectively by two types of dopamine receptors, classified broadly as D1 and D2. The D1 receptors are known to facilitate N-methyl-D-aspartate (NMDA) and GABA\(_A\) mediated synapses [24], [25], [26], while the D2 receptor types suppress the \( \alpha \)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) and GABA\(_A\) mediated synapses [27], [28].

The Str-MSN cells that are modulated by the D1 receptors (Str-MSN-D1) make major inhibitory axonal projections to the SNr. The Str-MSN cells that are modulated by the D2 receptors (Str-MSN-D2) project to the GPe. The GPe cells project inhibitory efferents to the STN, SNr, Str-FSI [29], as well as on themselves [30]. Both Str-MSN-D1 and Str-MSN-D2 have recurrent inhibitory connections on themselves in addition to laterally inhibiting one another [28]. The Str-FSI s make feed-forward inhibitory synapses on both Str-MSN-D1 and Str-MSN-D2 [31], as well as within the population. We note that the STN are the only excitatory cell population in the BG; all other cells are inhibitory. The STN cells receive major excitatory inputs from the thalamus and cortex [32], and project excitatory efferents to the GPe and SNr populations. The SNr forms the output structure of the BG and projects inhibitory efferents to the ventral thalamus and the brainstem reticular formation. In addition, the SNr cells make inhibitory projections on other cells within the population [33].

Table I shows the total number of cells in each population...
of a rat BG as reported by Oorschot [21]. However, in this preliminary endeavour to model the BG circuit on SpiNNaker, our objective is to build a single-channel columnar architecture that will serve as the basic building block towards building multi-channel BG models; the justification towards such an approach is to build a scalable framework. Towards this, we scale down the number of neurons in each population by a factor of 10³. While there are a myriad chemical neurotransmissions in the BG, we have implemented only two types of synapses in this work, viz. those mediated by the AMPA and GABA\_A neurotrans-receptors corresponding to glutamatergic (excitatory) and GABA-ergic (inhibitory) neurotransmitters. Further details on synaptic layout and parameterisation of the network are mentioned in the following sections.

B. Single neuron models and spiking patterns

Each computational unit in the BG network is a conductance-based form of Izhikevich’s spiking neuron model [34] implemented on the SpiNNaker software toolchain sPyNNaker [13], and defined in Equations (1) – (3).

\[
\begin{align*}
\frac{dv(t)}{dt} &= 0.04v^2(t) + 5v(t) + 140 - u(t) + I_{dc} + I_{syn}(t) \\
\frac{du(t)}{dt} &= a(bv(t) - u(t)) \\
\text{If } v(t) > 30 \text{ then } v(t) &\leftarrow c; u(t) \leftarrow u(t) + d
\end{align*}
\]

where, \(a, b, c, d\) are parameters that define the dynamic behaviour of the model and can be tuned to obtain various spiking patterns as observed in biology; \(v(t)\) is the membrane potential and \(u(t)\) is a membrane recovery variable; \(I_{dc}\) is the DC bias current that is built into the model definition in sPyNNaker; \(I_{syn}(t)\) is the post-synaptic current corresponding to the synaptic processes mediated by the neuro-receptors \(syn \in \{AMPA,GABA_A\}\) and is defined in Equations (4) – (6).

\[
\begin{align*}
I_{syn}(t) &= g_{syn}(t) \cdot (E_{syn} - v(t)), \\
g_{syn}(t) &= g_{syn}(t_0) \cdot e^{-(t-t_0)/\tau_{syn}}, \\
g_{syn}(t_0) &= g_{syn}(t_0 - \Delta t) + n \hat{g}_{syn},
\end{align*}
\]

where \(E_{syn}\) and \(g_{syn}(t)\) are the membrane reversal potential and membrane conductance respectively of the post-synaptic neuron; \(\tau_{syn}\) is the decay time constant of the synapse; \(g_{syn}(t_0)\) is the instantaneous conductance after the most recent afferent spike; \(n\) is the total number of spikes incident at the synapse in the time-step (\(\Delta t\)) before \(t_0\) (implemented on the ring buffer of sPyNNaker (see Sect. II-D)); \(g_{syn}\) is the conductance increment per afferent spike (the ‘synaptic weight’).

Initially, each population of the BG is simulated on SpiNNaker with neither any inter- or intra-population connectivities nor any extrinsic model input \(I_{syn}(t) = 0\). The DC bias current \(I_{dc}\) corresponding to a population \(X\) forms part of the neuron definition on SpiNNaker, and is present as an intrinsic input stimulus to all neurons in \(X\) from the start of simulation time. In this state, a single simulation run of the BG model will generate spiking behaviour from all neurons in the model, where each neuron is responding to \(I_{dc}\) only, and is otherwise acting independent of all other neurons in the model. Such a set-up allows us to specify a ‘base state’ spiking pattern for each population of the model; specifically, we aim to emulate the spike patterns demonstrated by Thibeault et al [15] (see Figure 1) generated on a conventional computer. The spiking patterns generated on SpiNNaker are shown in Fig. 2. The total duration of simulation is 5 s, where \(I_{dc}\) is varied after every 1 s of simulation time; this demonstrates the spike response characteristics of each population in terms of increasing or decreasing frequency corresponding to changes in \(I_{dc}\). The base (reference) state parameters of the single neuron models (with the exception of \(I_{dc}\)) for each cell population in our...
TABLE II

(A) The base (reference) state parameters of single neuron models in each population of the BG model defined in Equations (1) – (3) [15], [14], [35]. Readers may note that the base values for \( I_{leak} \) are set during the simulation of the whole network discussed in Sect. III-A. (B) Base state parameter values for synaptic and dopaminergic modulation attributes defined in Equations (4) – (9).

<table>
<thead>
<tr>
<th>(A) Izhikevich Neuron Parameters</th>
<th>Basal Ganglia</th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>d</th>
<th>( u_{init} ) (mV)</th>
<th>( I_{leak} ) (nA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Str-MSN</td>
<td>0.02</td>
<td>0.2</td>
<td>-65</td>
<td>8</td>
<td>-80</td>
<td>-16</td>
<td>-10</td>
</tr>
<tr>
<td>Str-FSI</td>
<td>0.1</td>
<td>0.2</td>
<td>-65</td>
<td>8</td>
<td>-70</td>
<td>-14</td>
<td>-10</td>
</tr>
<tr>
<td>GPc</td>
<td>0.005</td>
<td>0.585</td>
<td>-65</td>
<td>4</td>
<td>-70</td>
<td>-40.95</td>
<td>2</td>
</tr>
<tr>
<td>SNr</td>
<td>0.005</td>
<td>0.265</td>
<td>-65</td>
<td>2</td>
<td>-60</td>
<td>-15.9</td>
<td>5</td>
</tr>
</tbody>
</table>

(B) Synaptic Parameters

<table>
<thead>
<tr>
<th>Neurotransmitter</th>
<th>Parameters</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>AMPA</td>
<td>( \tau_{ampa} )</td>
<td>6 ms</td>
</tr>
<tr>
<td></td>
<td>( g_{ampa} )</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>( \phi_{ampa} )</td>
<td>0 mV</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>2.75</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>3.75</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>0.2</td>
</tr>
<tr>
<td>GABA</td>
<td>( \tau_{gaba} )</td>
<td>4 ms</td>
</tr>
<tr>
<td></td>
<td>( g_{gaba} )</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>( E_{gaba} )</td>
<td>-80 mV</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>2.75</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>2.75</td>
</tr>
<tr>
<td></td>
<td>( \epsilon_{gaba} )</td>
<td>0.073</td>
</tr>
<tr>
<td></td>
<td>( \phi_{dop} )</td>
<td>0.2</td>
</tr>
</tbody>
</table>

work are mentioned in Table II (A), and are informed by those in [14], [15]. (Readers may note that the base values for \( I_{leak} \) mentioned in Table II (A) are set during simulation and testing of the BG network as discussed in Sect. III-A).

C. Synaptic layout and dopaminergic modulation

In a recent review [32], all cell populations in the BG are reported as expressing both AMPA and NMDA neurotransceptors corresponding to glutamatergic neurotransmitters, and both GABA\(_A\) and GABA\(_B\) neuroreceptors corresponding to GABA neurotransmitters. However, the GABA\(_B\) neuroreceptor mediated synapses are modulatory in nature, and they do not participate actively in the synaptic transmission process. Furthermore, the NMDA neuro-receptor based synapse is a function of the membrane voltage. For simplicity, we ignore the NMDA and GABA\(_B\) based neurotransmission in this work, and consider synapses mediated by the AMPA and GABA\(_A\) neuro-receptors only.

The reversal potential for the GABA\(_A\) neuro-receptor mediated synapses depends on the flow of Cl\(^-\), and its value relative to the resting potential of the cell affects the nature of the post-synaptic membrane potential [36], [37], [38]. Here, we assume the case where a GABA\(_A\) mediated synapse would generate an inhibitory post-synaptic potential (IPSP); thus we set the value to -80 mV [15]. The base parameter values of \( \tau_{syn} \), and \( E_{syn} \) of the post-synaptic membrane corresponding to \( syn \in \{AMPA, GABA_A\} \) neuro-receptor mediated synapses (defined in Equations (4) – (6)) are as in [9] and listed in Table II(B).

A literature survey indicates extensive dopaminergic modulation of synaptic transmission in all cell populations of the BG [39], [40], [41], [42], [43], [44], [45], [23], [32]. For simplicity, we have constrained the dopaminergic modulation in our BG model to only a few synaptic pathways as listed below. The mathematical implementation of dopamine modulation is applied to the peak membrane conductance corresponding to a synapse, and is informed by prior works [9], [46].

1) Weakening of the AMPA mediated synapses:
The D2 receptors primarily target the AMPA neuro-receptor mediated synapses, weakening their impact by around 20% [24]. The D1 receptors primarily facilitate the NMDA mediated synapses while the AMPA mediated synapses are left unaffected. The modulation of the AMPA mediated excitatory afferents from extrinsic sources (thalamus/cortex) to the Str-MSN-D2, Str-FSI and STN are implemented using Eq. (7):

\[
\tilde{g}_{ampa-d2}^P = \tilde{g}_{ampa} \cdot (1 - \epsilon_{ampa}^d \cdot \phi_{dop}),
\]

where \( \tilde{P} \in \{\text{Str-MSN, Str-FSI, STN}\} \) represents the afferent populations receiving extrinsic inputs; \( \epsilon_{ampa}^d \) is the modulation co-efficient and is set as 0.2 to emulate the 20% modulation of the AMPA based synapses [24]; 0 < \( \phi_{dop}^P < 5 \) is the level of dopamine affecting the afferent synapse to the population \( P \). Thus, for a maximum value of \( \phi_{dop}^P \), \( \tilde{g}_{ampa-d2} = 0 \); Conversely, for lack of dopaminergic modulation i.e. \( \phi_{dop} = 0 \), the maximal value of \( \tilde{g}_{ampa-d2}^P = \tilde{g}_{ampa} \).

2) Modulation of GABA\(_A\) mediated synapses:
The GABA-ergic inhibition of the GPc by the Str-MSN-D2 population is weakened by the D2 receptors [44], while the GABA-ergic inhibition of Snr by the Str-MSN-D1 population is facilitated by the D1 receptors [25]; it is however unclear whether such facilitation is via pre-synaptic or post-synaptic receptors. We have implemented both these modulatory pathways as in Equations (8) and (9).

\[
\tilde{g}_{gaba-d2}^P = g_{gaba} \cdot (1 - \epsilon_{gaba}^d \cdot \phi_{dop}),
\]

\[
\tilde{g}_{gaba-d2}^P = g_{gaba} \cdot (1 + \epsilon_{gaba}^d \cdot \phi_{dop})
\]

For simplicity in this work, we constrain the dopaminergic modulation variability of GABA-ergic synapses to the MSN population only, and assume dopaminergic modulation of inhibitory afferents of both the GPc and SNr from Str-MSN to be mediated by pre-synaptic D2 and D1 receptors respectively. Thus, in Eq. (8), \( \tilde{P} \in \{\text{Str-MSN - D2}\} \), while in Eq. (9), \( \tilde{P} \in \{\text{Str-MSN - D1}\} \). Also, the co-efficient of dopaminergic modulation of GABA\(_A\) in both pathways is set to a base value of 7.3% i.e. \( \epsilon_{gaba} = 0.073 \).

3) Modulation of AMPA afferents of the STN:
The STN sends out diffused excitatory projections to the GPc and SNr populations of the same channel as well as of neighbouring channels (see Sec II-D). Once again for simplicity, we have assumed pre-synaptic dopaminergic modulation of these AMPA mediated effectors, implemented using Eq. (7).
In addition to the above, the dopaminergic weakening of GABA_4 mediated synapses is modelled by reducing the membrane conductance to a fraction of $g_{\text{gaba}}$.

The base values of all the dopaminergic modulatory parameters are mentioned in Table II(B), while the values for the modulated conductance, $g_{\text{syn}}$, are mentioned in Table III. These values are based on [9], although the final values are set by a 'trial and error' approach during model simulation so as to obtain the target firing rate (see Sect. III-A). Readers may note that we did not vary the dopamine levels ($\Phi, \epsilon$) for this work, and all results are generated with the base parameter values as mentioned in Table II (B). Thus, the dopamine modulation parameters in this model serve to set its operating region.

D. Overview of SpiNNaker and its handling of a synapse

SpiNNaker (Spiking Neural Network Architecture) is a System-on-Chip (SoC) consisting of very-low-power ARM968 processors. The on-chip communication architecture and protocols are biologically inspired, allowing asynchronous (event-based), parallel processing of synaptic data during neural network simulations on the ARM processors (referred to as 'cores'). Each chip has 18 cores, of which around 15 – 16 are available for neural computation; the remaining cores are used for system management on the chip. Each core has 64 KB (Data) Tightly Coupled Memory (DTCM: analogous to a ‘cache’ on a conventional computer, i.e. for quick data access during neural computation) where the neuron and synapse data pertaining to that core is stored to be accessed during neural computation. In addition, each chip has a 128 MB Synchronous Dynamic Random Access Memory (SDRAM) that is shared by all the cores on the chip for storing simulation data. For details of current state-of-the-art in SpiNNaker development, we refer the reader to a recent topical review [47]. The model used in this work is implemented on a single 48-chip SpiNNaker board (please refer to Sect. II-D for details).

The SpiNNaker software chain, sPyNNer [13], provides an implementation of PyNN [48], which is used as a standard interface for all neural simulations on SpiNNaker. (PyNN is a python based library bespoke to building spiking neural network models, and runs on other simulators (e.g. NEST) besides SpiNNaker). Two essential criteria that form the backbone of the sPyNNer application interface are: the neuron model, which we have discussed in Sect. II-B; and the synaptic connections in the network, which we describe in the following text.

The synaptic connectivity in the model is implemented via
Fig. 3. The single-channel basal ganglia network, as exported from SpineCreator. Rectangular boxes are neural populations. The population name, number of elements and SpineML component are shown in the box. Grey circles represent Poisson spike train sources. Green arrows are projections with element to element connectivities that are parameterised as in the SpiNNaker based model and reported in Tables I – III. Projections with arrow heads are excitatory, those with circles for heads are inhibitory. The thinner, red lines which connect populations to arrowheads connect the membrane voltage variable \( v(t) \) (defined in Eq. (1)) in the effenter population to the synapse component in each projection (on a one to one basis), allowing the synaptic current to be computed.

sPyNNaker using the function ‘FixedProbabilityConnector’. Each connection between two populations consists of three attributes viz. (a) the probability of the synaptic connection \( p_{conn} \in (0, 1) \), which is a normalised representation of the total ‘fan-in’ from the pre-synaptic population to the post-synaptic population; (b) the delay of the synaptic connection \( d_{conn} \), representing the latency of a pre-synaptic cell spike in reaching the post-synaptic cell; (c) the synaptic (connectivity) weight that scales the synaptic decay exponential, and is the membrane conductance increment per spike of the post-synaptic neuron \( g_{syn} \) in the current work (defined in Eq. (6)).

All of the above-mentioned synaptic attributes corresponding to a certain projection are stored as a 32-bit ‘synaptic data word’; the first 16 bits consist of the synaptic weight, while the next 16 bits are distributed thus: 4 bits for synaptic delay; 1-bit for synapse nature (i.e. excitatory/inhibitory); 8 bits for neuron index (therefore capped to 256 neurons per core); and 3 bits are unused. This is shown in Appendix, Fig. 10(b).

E. Mapping the BG model to SpineML

A neuronal model that has been specified in SpineML consists of individual XML files, which define the behaviour of model ‘components’ viz. neuron bodies, post-synapses and weight-updates. In addition, there are separate XML files that define how the components are built into a network of neuronal populations that are connected with ‘projections’, where each projection consists of one weight-update and one post-synapse component. A set of ‘experiment files’ define how the model should be executed; each experiment contains a specification of the inputs for the network, the data that should be logged from the simulation and any experiment-specific network lesions or parameter modifications that should be made. SpineML is thus a declarative format for specifying a network model.

In order to execute a SpineML model, it is necessary to use a SpineML back-end, which parses the SpineML input files and generates executable code for the model. We used SpineML 2 BRAHMS [49] that generates code suitable for execution on a general purpose CPU and is the canonical back-end for SpineML. The single-channel BG model on SpiNNaker in Fig. 1 is mapped to SpineML and shown in Fig. 3. Inspection of the figures indicates that the network connectivity is the same, although there is an important difference in solving the synapse models. Fig. 3 shows a typical SpineML model in which ‘spike events’ are transmitted along projections via a weight-update component (a mechanism to implement fan-in from multiple pre-synaptic neurons) and then to the post-synapse component of the projection whose conductivity is incremented by the synaptic weight \( g_{syn} \). This \( g_{syn} \) is a parameter of the post-synapse component, which means that a spike afferent from population A and a spike afferent from a different population B increment the conductivity at the post synapse by the same amount. In contrast, in SpiNNaker, each spike is transmitted in a data packet which encodes \( \bar{g} \) (see the ring-buffer implementation in Fig. 10, Appendix), meaning that the spike from population A could increment the post-synaptic conductivity by a different amount than the spike from population B. Thus, there can be a difference between SpineML and SpiNNaker networks which have apparently been arranged in an identical manner. It is possible to create a SpineML network which faithfully reproduces the behaviour of the SpiNNaker network, but this leads to a more complex, unwieldy network (see the SpineML model bgbsb1_empt in the repository referenced below and compare with bgbsb1, which is shown in Fig. 3). The more complex, and more faithfully SpiNNaker-like SpineML model produced results that were not statistically different from those discussed in Sect. IV for the more natural SpineML models.

The SpineML model and associated results are available publicly at https://github.com/ABRG-Models/GPR-BSB/>

F. A scaled-up BG model with three channels

The basal ganglia has multiple parallel pathways that serve different parts of the cortex [50], as well as segregated voluntary and automatic behaviour pathways [51]. Furthermore, focussed inhibition and surround excitation are concepts that were proposed by [2]. We have scaled up our BG model to consist of three channels, where each channel is the single-channel model of Fig. 1. Thus, the total number of neurons in this three-channel BG model is 8043. The STN population of each channel sends out excitatory efferents to the SNr and GPe populations of the other two channels. These cross-channel connections produce the desired surround effect on the neighbouring channels by each channel \( \Psi \), thus indirectly emphasising the focal inhibition within \( \Psi \). The delay parameter of the cross-channel efferents of the STN is higher than that of the intra-channel pathway (see Table III (F)). The connectivity parameters are mentioned in Table III. The method of initiating competing sources and incorporating action-selection in the model is discussed in Sect. III.
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Fig. 4. The schematic diagram of a scaled up model of the Basal Ganglia where three channels are interconnected laterally. The STN population of each channel projects to GPe and SNr populations of every other channel. The connectivity parameters are mentioned in Table III (F), and referred to as ‘cross-channel’. The model demonstrates action-selection when presented with competing inputs (see Sect. III-B).

III. SIMULATION METHODS ON SpiNNAKER AND RESULTS

A. Base state dynamics of the single-channel model

Simulation methods: To mimic extrinsic input to the BG, a total of 25 Poisson distributed spike sources project to the Str-MSN and Str-FSI populations, while 2 Poisson sources feed the STN population. The Poisson number for all the spike sources is maintained at 3 Hz. The total duration of simulation is 10 s. The Poisson sources are applied stochastically sometime between 500 – 700 ms from the start of simulation, and for a total duration of 9.2 s. The individual neuron model equations on sPyNNaker are solved using a 2nd order Runge Kutta solver with a time-step 0.1 ms to achieve solution accuracy.

The average firing rate of each population over $r$ trials is derived using Eq. (10):

$$S_R = \frac{\sum_{r=1}^{R} S_m^r}{t_{sim} \cdot N_m \cdot R}$$

where $S_m^r$ is the total number of spikes fired for each trial $r$ by all $N_m$ neurons in the population $M \in \{SNr, GPe, STN\}$, and over the total simulation duration of $t_{sim} = 10$ s; $R = 10$ is the total number of trial simulation runs. The spike count histogram is derived by averaging the spike count for each 100 ms bin of $S_R$.

Results: The average firing rates for the STN, GPe and SNr populations are shown in Fig. 5(a), while the spike count histogram with 100 ms bins is shown in Fig. 5(b). The firing rate for: STN lies within the range 10 – 12 Hz; GPe lies within the range 30 – 32 Hz; SNr lies within the range 20 – 25 Hz. Thus, our results show a good similarity with those reported in [9] (see Fig. 2) and [15] (see Fig. 6). We now treat this model as the single ‘channel’ of decision-making and action-selection in the BG.

B. Simulating action-selection on SpiNNaker

Next, we aim to simulate action-selection using the three-channel BG model presented in Sect. II-F.

Simulation methods: Similar to the single channel model, Poisson input at 3 Hz is provided to all the three channels from around 500 – 700 ms and for a duration of 9.2 s. Total simulation time is 10 s at a resolution of 0.1 ms. At $t_1 = 3$ s from the start of simulation, the first channel receives a request for being ‘selected’, which is simulated by providing Poisson spike trains at 15 Hz, drawn from two separate pools consisting of 25 and 2 Poisson spike sources, and provided to the Striatum and STN populations respectively in the channel. At $t_2 = 6$ s, the second channel receives a request for being ‘selected’, which is simulated by 25 Hz Poisson spike trains provided to all the input pathway cells of the channel. Both 15 Hz and 25 Hz spike train inputs are present until $9.9$ s from start of simulation. Thus, all Poisson sources are withdrawn at $100$ ms before the end of simulation; after this time and until the end of simulation, all channels respond to $I_{dc}$ only. The outputs of the model are the SNr firing rates of all three channels. The results are averaged over 10 trials. The average firing rate time histogram with time bin widths of 1 s is shown in Fig. 6. All dopamine levels were kept at base values indicated in Table II (B).

Data collected from BG of awake resting rats suggest firing rates of: STN at around 10 – 12 Hz; GPe at $\approx 30$ Hz; SNr at somewhere in the range 25 – 30 Hz, and generally less than that of GPe [9]. Furthermore, the firing rate of the Str-MSN cells are $\approx 3$ Hz while that of the Str-FSI are $\lesssim 10$ Hz. We tuned the parameters in Tables II (B) and III to emulate these firing rates, which will define the base firing rates of the model populations.
Fig. 6. The average firing rate histogram, with a bin width of 1 s, of SNr populations in the (blue, dash-dot line) first, (pink, dash line) second, and (yellow, dot line) third channels of the BG model shown in Fig. 4. The model response demonstrates action-selection in the circuit. The third channel (yellow) does not receive any competing input and is therefore the neutral channel.

Results: Figure 6 demonstrates action selection in the three-channel model simulated on the SpiNNaker machine. When a channel receives a Poisson input with a higher frequency, the focal inhibition of SNr within the channel increases. At the same time, the STN cells of that channel provide increased excitatory projections to the SNr cells of the neighbouring channels, thus producing an ‘off-centre’ effect, and the firing rate of the SNr population in the competing channel drops relative to the other channels. We did not set a specific firing rate threshold to demonstrate selection of action, and rather let the inherent model dynamics take control. Thus, all parameters in each channel of the model are the same as those of the single-channel model.

IV. COMPARISON WITH SPINEML AND PERFORMANCE ANALYSIS

The model implemented on SpineML is intended to be topologically and parametrically identical to the one running on SpiNNaker. All parameters and network connectivities are configured the same in both systems. Figure 7(a) shows the firing rate histogram of the SNr, STN and GPe populations of the single-channel BG circuit simulated using the SpineML model. The figure demonstrates similar mean spiking rates for the STN and SNr as on the SpiNNaker-based model shown in Fig. 5; the firing rate lower bound of GPe cells are lower than that on SpiNNaker. The three-channel BG circuit on SpineML also demonstrates action selection behaviour when simulated with exactly the same parameter and input attributes. This is shown in Fig. 7(b). The results demonstrate qualitative and functional similarity between models simulated on the two independent platforms.

A. Statistical comparison of the single-channel models

Although we did not expect the numerical results of simulations to be identical, we tested whether the two implementations would generate statistically equivalent results. Table IV gives the results of statistical tests on the spike counts in each population of the single-channel model obtained from 30 repetitions of 10 s simulations run on both SpiNNaker and SpineML_BRAHMS. The standard error of the mean number of spikes was computed by the bootstrap method with 256 resamples. The difference of the means is also given, along with a bootstrapped estimate of the difference of the means from 256 resamples. Finally a bootstrapped test of the difference of the mean number of spikes was applied with 10000 resamples. The Achieved Significance Level is a measure of ‘the probability that the means are indistinguishable’ (see Appendix). The results indicate that whilst the spike counts are similar, none of the results can be said to be statistically equivalent according to this stringent test that the spike counts be indistinguishable in all populations.

B. Performance analysis

The PyNN script describing the BG model is mapped and executed on the SpiNNaker machine by the sPyNNaker software toolchain [13], which itself runs on a host machine, in three stages: pre-processing, execution and post-processing. Pre-processing involves translation of the PyNN-defined network into a form suitable for the SpiNNaker machine, and includes partitioning, routing, and generation and loading of data structures. In the context of performance testing, execution is defined as the time taken, once all data has been loaded, to run the simulation on the SpiNNaker machine. Post-processing refers to extraction of resultant data, generated by executing the model, from the SpiNNaker machine to the
host machine. Both uploading/extraction of data to/from the SpiNNaker machine is currently done via ethernet [4].

The single-channel BG model consists of $2.68 \times 10^4$ neurons and $\approx 0.68 \times 10^6$ synapses (estimated from projection probabilities). While each processor within a SpiNNaker chip is capable of simulating an upper limit of 256 neurons (discussed in Sect. II-D), memory requirements of the neuron model and synaptic connectivity for certain applications may cause this number to be reduced. In the current work, sPyNNaker maps the single-channel BG model on to 32 cores distributed across 2 SpiNNaker chips, residing on a single 48-chip SpiNNaker board. In case of the three-channel model, the total number of neurons and synapses are $8.043 \times 10^5$ and $\approx 2.05 \times 10^6$ respectively, and the model network is mapped by sPyNNaker on to 96 cores, distributed across 7 SpiNNaker chips.

Pre-processing is done on a 4-core 8 GB RAM desktop host machine, and takes 70.5 s for the single-channel BG model (three-channel: 191.0 s). The SpiNNaker hardware is designed to execute neuronal models in real time at a resolution $\geq 1$ ms. Both single- and three-channel BG networks are configured to simulate with a solver time-step of 0.1 ms in order to maintain solution accuracy. Due to this constraint, 1 s of model simulation time is executed in 10 s real (‘wall clock’) time. However, both the single- and three-channel models are guaranteed to execute within this 10 s. On execution completion, a further 119.1 s is required to extract output data for the single-channel model (three-channel: 574.7 s), giving an average total simulation time of 199.8 s for the single-channel model (three-channel: 776.0 s). The timing data recorded from the SpiNNaker execution of both the single-channel and three-channel models is shown in Fig. 8. Timing values are averaged across 10 repeated runs; the standard deviations across the 10 samples of each model were less than 1.3 s, 4 ms and 2.7 s for pre-processing, execution and post-processing respectively.

The above-mentioned data for SpiNNaker-based model simulation is now compared to that using SpineML and executing on a 4-core 8 GB RAM desktop host machine, extracting and saving data and ‘logs’ (post-processing). The results are also shown in Fig. 8, and indicate that the single-channel model simulated on SpineML performs pre-processing in 0.036 s, while 1 s of model simulation time is executed in 3.5 s real time. For the three-channel model, pre-processing time is 0.1 s, and the execution time equivalent of 1 s simulation time is increased significantly to 26.7 s real time. The post-processing time is insignificant in both cases and $\approx 0$. Clearly, the time of execution increases with scaling up of the model, and emphasises the advantage of SpiNNaker-based computation for larger models over conventional computers.

Power consumption on SpiNNaker: In a recent work, we used in-house Arduino-based power measurement equipment to measure power directly from a 48-node SpiNNaker board during model execution (the reader may refer to [20] for details). The main draw-back of this previous set-up was the coarse resolution (8.9 ms) of recording power from the SpiNNaker board. In this work, we have used an enhanced (Raspberry-pi-based) version of this equipment, allowing a resolution of up to 0.6 ms with cleaner recording, i.e. without noise/glitches. Thus, the sampling rate of recording the power is higher than 1 ms, the time-step of model simulation, and minimises the potential for data loss due to delays during communication with the SpiNNaker board via ethernet. Our study shows that the single-channel model execution uses $\approx 800$ mW, while the three-channel model execution consumes $\approx 1.8$ W shown in Fig. 9. The figure also confirms that the model execution time is not affected by scaling up to three channels, and is consistent at 100 s real time corresponding to a simulation time of 10 s. As power consumed during pre- and post-processing are negligible compared to that during model execution, we kept the post-processing time to a minimum; pre-processing times are handled by sPyNNaker and is not accessible to the user.

V. DISCUSSION AND CONCLUSION

We have presented a biologically-plausible and scalable model of the Basal Ganglia (BG) circuit, designed to run on the SpiNNaker machine — a biologically-inspired architecture built with low-power ARM processors, allowing inherent asynchronous, parallel computation, and in real time for time-steps

\begin{table}[h]
\centering
\caption{Table IV: Comparison of single-channel BG model simulated on SpineML and SpiNNaker.}
\begin{tabular}{|c|c|c|c|c|}
\hline
\textbf{BG population} & \textbf{Mean spike count} & \textbf{SpineML} & \textbf{SpiNNaker} & \textbf{ASL} \\
\hline
Sth-MSN-D1 & 3127 (4) & 2869 (82) & 258 (113) & 0.016 \\
Sth-MSN-D2 & 0.33 (0.13) & 0.5 (0.17) & -0.17 (0.2) & 0.22 \\
STN & 1.73 (0.38) & 1.47 (0.31) & 0.27 (0.49) & 0.28 \\
GPe & 1072 (6.2) & 1158 (1.7) & -86 (5.7) & <0.0001 \\
SNc & 8207 (6.9) & 8518 (4.3) & -311 (7.6) & <0.0001 \\
\hline
\end{tabular}
\end{table}
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Fig. 9. The power consumptions of the three-channel model using an in-house Raspberry-pi-based measurement system connected to the SpiNNaker board (see [20] for details). The duration of recording the power can be broken down into four regions: (i) booting the machine; (ii) pre-processing of data; (iii) model execution; (iv) post-processing (i.e. data extraction); the delay of around 4 s after booting the machine is inserted for clarity. The peak-to-peak power in region (iii) is 1800 mW. The measurement sampling interval is 0.6 ms in real time. This is shorter than the time-step of model simulation (1 ms real time) in order to ensure that we do not lose data due to circuit delays between the Raspberry pi and the SpiNNaker board.

> 1 ms. A single neuro-computational unit in our BG model is simulated with a conductance-based Izhikevich neuron model, facilitated by the underlying SpiNNaker software toolchain, sPyNNaker, which in turn is based on PyNN, a python-based neural network application interface. A columnar structure of the BG circuitry is first parameterised on SpiNNaker to set the base firing rates for all model cell populations, informed by existing literature. This forms the basic building block for a scalable framework, and is thought to be a single-channel for action-selection in the BG. To simulate action-selection by competing inputs, we scaled up the model to consist of three channels, and tested with two competing inputs in the presence of background noisy stimulus. Our results show that an input stimulus that is larger than the others is always the ‘winner’, indicated by a relative drop in the firing rate of the SNr population (representing the BG model output) in the competing channel. The reduced firing rate of the inhibitory SNr population implies a reduced inhibition of the thalamic/brainstem cells, which are known to be the recipients of the BG output. This in turn means that the ‘action’ that is solicited by a relatively larger (‘competing’) input is now ‘decided’ by the BG circuit to be ‘selected, and acted upon’, indicated by disinhibition of the target outputs. We have tested our model with a competing input of 15 Hz in the presence of a background noisy input of 3 Hz. This is further confirmed by ‘selection’ of a larger input of 25 Hz provided in the presence of both 15 Hz and 3 Hz inputs. On both occasions, the largest input wins. It is worth mentioning here that dopamine neurotransmitter-receptor levels are fundamental to facilitating decision-making and action-selection by the BG. Here, we tuned the base parameters simulating neutral dopamine levels; implementation of the three-channel model on SpineML, following exact same implementation procedures as on SpiNNaker, demonstrates action-selection by a larger input. Overall, the functional and qualitative behaviour of the models are in agreement. A difference of means test (see Appendix) indicates statistically significant numerical difference between the two platforms. We speculate that such difference is due to the stochastic nature of the model inputs, and simulates the numerical differences in recorded data from different brains, even when they are in the same state, or performing similar behavioural tasks. We believe that our comparative study will provide a basic framework for mapping SpiNNaker-based models to SpineML, as well as for performance benchmarking of SpiNNaker with conventional computers during neuronal simulation.

The main drawback of our model is the inability to implement parameters that are voltage dependent, and thus need updating during run-time. Thus, we were unable to implement the voltage dependent NMDA synapses, nor the gap-junction (resistive) connections in the Str-FSI populations. This is due to current computational constraints on SpiNNaker during runtime, and work is ongoing to provide such implementations in the future. Another drawback is the slow ethernet-based data transfer rates between the host-machine and SpiNNaker. This is indicated in the performance analysis where the post-processing (data extraction) times are observed to increase significantly with scaling up of the model. In comparison, the pre-processing (mapping high-level model description to simulator) and post-processing times for both single- and three-channel models implemented on SpineML are negligible. Model execution on SpiNNaker for this work is slowed down.

To verify our model results simulated on SpiNNaker, we mapped the model to SpineML, an XML-based platform representing model attributes as ‘components’, and executing the models with SpineML_2_BRAHMS, a bespoke simulator which converts the SpineML model into machine code and runs it on a conventional computer. We aimed for the BG model implementation on SpineML to have the exact same network topology and neuron attributes as the SpiNNaker version, and therefore retained all model connectivities and parameter values used in the latter. Model results on SpineML show qualitative similarity with those on SpiNNaker in terms of base firing rates of the single-channel BG model cell populations. Implementation of the three-channel model on SpineML, following exact same implementation procedures as on SpiNNaker, demonstrates action-selection by a larger input. Overall, the functional and qualitative behaviour of the models are in agreement. A difference of means test (see Appendix) indicates statistically significant numerical difference between the two platforms. We speculate that such difference is due to the stochastic nature of the model inputs, and simulates the numerical differences in recorded data from different brains, even when they are in the same state, or performing similar behavioural tasks. We believe that our comparative study will provide a basic framework for mapping SpiNNaker-based models to SpineML, as well as for performance benchmarking of SpiNNaker with conventional computers during neuronal simulation.
by a factor of 10 relative to real time. This is because, the underlying Izhikevich equations need to be computed with a time-step of 0.1 ms to achieve solution accuracy, while the inherent SpiNNaker design is for real time operation with time-steps $\geq 1$ ms. Thus, 10 s of model simulation time on SpiNNaker runs in 100 s real time. However, this execution time is guaranteed i.e. both single-channel and three-channel BG models execute in 100 s real time corresponding to 10 s simulation time — this consistency demonstrates the ability of SpiNNaker to scale network size without compromising on execution time. In contrast, although the single-channel model execution time on SpineML is lower than that on SpiNNaker ($\approx 3$ s real time for 1 s simulation time), that for the three-channel model scales up significantly, and by an order of 10 (approximately). Continuing research on the BG model implementation on SpiNNaker is looking into further scaling up of the model, which will serve to test and challenge the SpiNNaker machine on its real time computational capabilities.

Continuing development of an in-house equipment is looking into ways to power directly from a 48-node SpiNNaker board during model execution [20]. To measure power during execution of the BG model, we use a Raspberry-pi based system (enhancement from the Arduino-based system described in [20]), allowing the recording of power at 0.6 ms (real time) resolution. This is $\approx$ half the sampling resolution at which the model is set to execute (1 ms real time). The single-channel model uses 2 SpiNNaker chips (32 cores) and dissipates $\approx 0.8$ W; the three-channel model runs on 7 SpiNNaker chips (96 cores) and dissipates $\approx 1.8$ W; the corresponding energy costs are 80 Joules (J) and 180 J respectively. In comparison, the thermal design power is $\approx 3$ W; the three-channel model scales up significantly, and by an order of 10 (approximately). Continuing research on the BG model implementation on SpiNNaker is looking into further scaling up of the model, which will serve to test and challenge the SpiNNaker machine on its real time computational capabilities.

In conclusion, our study demonstrates the SpiNNaker platform as capable of simulating biologically-plausible decision-making and action-selection circuitry that executes in a parallel and asynchronous manner, and within guaranteed timescales. Furthermore, the platform demonstrates the potential for simulating large scale models without compromising on execution times. In addition, prior research has shown the low energy requirements of the SpiNNaker machine (e.g. compared to NEST [52]). Not surprisingly, therefore, use of SpiNNaker has been proposed in several robotic applications [53], [54]. Autonomous intelligent decision-making is a key desirable attribute in robotic applications, which can benefit wide-ranging societal requirements. We believe our work developing the BG model on SpiNNaker will strengthen endeavours to build intelligent decision-making machines.

APPENDIX

1) The Ring Buffer — implementing a synapse: Let us assume an example case where a neuron-X, residing in core-Y of one SpiNNaker chip, initiates a spike transfer that is to be delivered to the post-synaptic neuron-Y, residing in core-Y of the same chip, at a delay $d_{\text{conn}} \approx 3$ ms, and with $p_{\text{conn}} = 1$, thus guaranteeing a connection. After a series of activities initiated by this spike event (the details of which can be found elsewhere [1] and are outside the scope of this report), the first 16 bits of the synaptic data word representing $g_{\text{syn}}$ is now fetched from the chip’s SDRAM and placed in a ring buffer of core-Y to be used for the post-synaptic membrane current computation for neuron-Y.

The ring buffer is a right circular shift-register structure occupying 16 KB of DTCM of each core, and is the basic algorithm that defines the post-synaptic behaviour in an afferent neuron population. A depiction of the ring buffer is shown in Fig. 10(a). Each neuron in a core will have two rows in the ring buffer pre-booked and at its disposal — one corresponding to an excitatory projection, and another corresponding to an inhibitory projection. Furthermore, each

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row consists of sixteen ‘slots’, and each slot consists of 16 bits. To access the synaptic weight in the ring buffer at the appropriate delay slot, there is a right circular shift ‘pointer’, which forms the reference for the specific slot in which the 16-bit synaptic weight is placed.

In our above-mentioned example case \( d_{\text{conn}} = 3 \) ms, therefore, if the pointer is pointing currently at the slot 14, then our synaptic weight data will be placed in slot 1, i.e. circular shifted \( 3^{rd} \) position from slot 14. Furthermore, if we assume there are ‘n’ excitatory synapses arriving to our single afferent neuron-Y, and all synapses are to be activated after 3 ms, then the resultant synaptic weight that is placed in the ring buffer at the 3 ms delay slot is a linear summation of all the afferent weights (Eq. (6)). Note that for simplicity and demonstration purposes, we have assumed a solver time-step of 1 ms. (The solver time-step is 0.1 ms (afore-mentioned) in the SpiNNaker and SpineML models. The achieved Significance Level

\[
l(t^*) = \frac{\bar{z}^* - \bar{y}^*}{\sqrt{\sigma^2_x/n + \sigma^2_y/n}}
\]

If the original means of \( z \) and \( y \) were genuinely very close, then \( z^* \) and \( y^* \) won’t have been shifted very much and it is likely that \( l(t^*) \) will exceed \( l(t) \) with probability around 0.5. If they were not close, and the mean\((z) \gg \text{mean}(y)\), then very few \( l(t^*) \) will exceed \( l(t) \). We made 10000 \( x^* \) resamples; the proportion of those for which \( l(t^*) \geq l(t) \) is the Achieved Significance Level (ASL). The smaller ASL is, the less probable is \( H_0 \), and the more significant is the difference of the means.

The test makes no assumption about the shape of the distributions which generated the samples, but it does assume that \( \bar{z} > \bar{y} \).

**AUTHOR CONTRIBUTIONS**

BSB designed the model layout, implemented the model on SpiNNaker, and generated SpiNNaker-based model simulation results. SJ implemented the model on SpineML, generated all SpineML-based model simulation results, and carried out statistical test for model comparison. KG advised on BG model design and implementation. OR made the performance analysis for time requirements. IS built the power measurement equipment and performed power analysis of the model simulation on SpiNNaker. AR and ABS supported with sPyNaker, and implementation and validation of the SpiNNaker-based model. SBF provided advice, guidance and support for model implementation on SpiNNaker. BSB, SJ, OR and IS generated the figures. All authors contributed to writing the manuscript.

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