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Exploring Self-Repair in a Coupled Spiking Astrocyte Neural Network

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Abstract—It is now known that astrocytes modulate activity at tripartite synapses where indirect signalling via the retrograde messenger, endocannabinoids, leads to a localised self-repairing capability. In this paper, a self-repairing Spiking Astrocyte Neural Network (SANN) is proposed to demonstrate a distributed self-repairing capability at the network level. The SANN uses a novel learning rule which combines the spike timing dependent plasticity (STDP) and Bienenstock, Cooper, and Munro (BCM) learning rules: hereafter referred to as the BSTDP rule. In this learning rule, the synaptic weight potentiation is not only driven by the temporal difference between the pre and postsynaptic neuron firing times but also by the postsynaptic neuron activity. We will show in this paper that the BSTDP modulates the height of the plasticity window to establish an input-output mapping (in the learning phase) and also maintains this mapping (via self-repair) if synaptic pathways become dysfunctional. It is the functional dependency of postsynaptic neuron firing activity on the height of the plasticity window that underpins how the SANN self-repairs on the fly. The SANN also uses the coupling between the tripartite synapses and γ-GABAergic (GABA) interneurons. This interaction gives rise to a presynaptic neuron frequency filtering capability which serves to route information, represented as spike trains, to different neurons in subsequent layers of the SANN. The proposed SANN follows a feed-forward architecture with multiple interneuron pathways and astrocytes modulate synaptic activity at the hidden and output neuronal layers. The self-repairing capability will be demonstrated in a robotic obstacle avoidance application and simulation results will show that the SANN can maintain learned manoeuvres at synaptic fault densities of up to 80% regardless of the fault locations.

Index Terms—Spiking neural networks, astrocyte, self-repair, fault tolerance, obstacle avoidance.

I. INTRODUCTION

W

ILE state-of-the-art hardware devices and neuromorphic chips [1]–[6] replicate to an extent a brain information processing paradigm, they are not fault-tolerant and can develop faults due to incorrect operations in post-manufacturing [7], wear-out failures [8], or radiation effects [9]. Therefore, fault tolerance is still a key challenge for modern hardware systems. Traditional approaches to address this challenge generally include several stages, e.g. fault detection [7], fault diagnosis [10], and correction/reconfiguration [11], [12]. Additionally, other techniques like redundancy [13], error correction method [14] can also be used to enhance the system fault tolerance. However, most of these approaches are not distributed and require a central control unit which can also be compromised due to faults. It has recently been reported [4], [15], [16] that the coupling of neurons and astrocytes in the brain may provide an elegant solution to this problem. Specifically, a network of excitatory and inhibitory synapses with spiking neurons was employed to develop a self-adaptive strategy for a robotic controller [17]. The authors have proposed a plastic spiking neural network model which was used to develop a fault-resilient robotic controller [18]. The approach of [19] showed that astrocytes have the potential to provide a distributed self-repairing function (e.g. when the spinal cord has injuries).

In this paper, a self-repairing spiking astrocyte neural network (SANN) is proposed where a novel learning rule (BSTDP) combining the spike-timing-dependent plasticity (STDP) and Bienenstock, Cooper, and Munro (BCM) learning rules is used which serves to initiate/suppress the learning process depending on the firing activity of pre and postsynaptic neurons. Specifically, it will be shown that the BSTDP rule is capable of (1): creating an input output mapping during a conventional learning phase and (2): maintaining this mapping in the presence of dysfunctional synaptic connections: this we define as self-repair. The BSTDP rule couples an γ-GABAergic (GABA) interneuron with tripartite synapse [20] which serves to modulate the height of plasticity windows as a function of presynaptic neuron frequency. This capability endows the SANN with a computationally useful frequency filtering capability where spike trains are routed through different synaptic pathways depending on presynaptic frequency. The proposed SANN is applied to a robotic obstacle avoidance task which is a well known benchmark in many robotic laboratories. Results will show, for the first time, that a computational network that captures known interactions between neurons and astrocytes can not only provide a local learning capability, whereby learning commences when either or both pre and postsynaptic neuron become relatively inactive, but also yield a self-repairing capability post learning to maintain learned mappings. Furthermore, results will also demonstrate that the self-repairing capability of SANN can maintain input-output mappings for a distributed fault density of up to 80%.
The main contributions of this paper are as follows:

1) A biologically inspired astrocyte-neural network (SANN) and learning rule is proposed. The BSTDP learning rule combines the STDP and BCM learning rules to initiate a conventional learning cycle and maintain learned input-output mappings in the presence of dysfunctional synapses, e.g. failures.

2) The BSTDP rule captures the biological interplay between GABA interneuron, postsynaptic neurons and the plasticity of synapses. This novel interaction gives the SANN the capability to continuously route data, represented in spike train frequencies, to different areas of the SANN and initiate a repair process in the presence of faults.

3) The fault tolerance of the proposed SANN architecture is evaluated through simulations using a robotic application. Results demonstrate the resilience of the SANN-based robotic controller in an obstacle avoidance application under various fault densities.

The rest of paper is organized as follows. Section II provides the background and motivation. Section III presents SANN models, the BSTDP learning rule and network structure. Section IV provides simulation results that demonstrate the performance of the SANN-based robotic controller under various fault densities and conditions. Section V concludes and gives insight into the potential directions for the future work.

II. BACKGROUND AND MOTIVATION

Astrocytes in the central nervous systems can encapsulate about ten thousand synapses and multiple neurons (e.g. ~6 in the cortex) [21]. These cells interact with synapses and neurons to modulate synaptic activity [22], namely the tripartite synapse [23]. It has been shown that at the tripartite synapse, the astrocyte cells perform a distributed and fine-grained repair [15], see Fig. 1, whereby when synapses are damaged, the Probability of neurotransmitter Release (PR) at synapses drops which leads to fall off in the activity at postsynaptic neurons. However, it is now commonly known that a negative feedback signal, often referred to as the retrograde messengers (endocannabinoids) pathway, exists where the messengers are represented in spike train frequencies, to different areas of the SANN and initiate a repair process in the presence of faults.

Recent research shows that the GABA interneuron participates in the activities of neurons and synapses [26] where for low presynaptic frequencies the GABA interneuron has an inhibitory affect on the presynaptic neuron. The signalling pathways between the GABA interneuron and tripartite synapse are shown in Fig. 2. The conventional tripartite synapse has three network is seriously damaged by a fault density of 80% [4]. This hardware architecture was optimized for the low area overhead [24] and achieved less hardware resource utilization than the original work of [4], therefore provides an efficient design solution for large scale networks. A small SANN has been applied to a mobile robotic car application [25] with a fixed input spike train. However, the work in this paper goes much further and proposes a SANN for a more complex robotic task in that the SANN takes on the role of a controller where input-output mappings are initially learned, allowing the robot to navigate around a series of obstacles. Furthermore, the SANN controller maintains the obstacle avoidance capability when faults occur. To the best of the authors’ knowledge, this is the first time that a network combining astrocytes with neurons has been shown to implement a learning and fault-tolerant capability which can be harnessed in a challenging application.

III. SELF-REPAIRING ASTROCYTE NEURAL NETWORK MODEL

In this section, the SANN architecture is presented together with the BSTDP learning rule and neuron/synaptic models. Specifically, we will start with communication between excitatory neurons, GABA neurons and astrocytes at tripartite synapses. From this we will distil out biological realistic models that capture the essence on the biological interplay between these cells while at the same time minimising the computational overhead.

A. Activity-dependent Mechanism in the Tripartite Synapses

Recent research shows that the GABA interneuron participates in the activities of neurons and synapses [26] where for low presynaptic frequencies the GABA interneuron has an inhibitory affect on the presynaptic neuron. The signalling pathways between the GABA interneuron and tripartite synapse are shown in Fig. 2. The conventional tripartite synapse has three
terminals shown by Fig. 1, which are the presynaptic axon, the postsynaptic dendrite and the astrocyte cell. However, the GABA interneuron introduces an additional terminal and we now consider this. When a spike train of frequency \( f_{\text{pre}} \) arrives at the presynaptic axon, the neurotransmitter (in this case glutamate) is released from cleft. It binds to the receptors at the postsynaptic dendrite, which depolarizes the postsynaptic neuron. The spikes presenting at the presynaptic axon also arrive at the GABA interneuron [26] causing GABA to be released which subsequently binds to GABA-B receptors at the astrocyte cell. This triggers the release of inositol 1, 4, 5-trisphosphate (\( IP_3 \)). For a low \( f_{\text{pre}} \) shown in Fig. 2A, the amount of released \( IP_3 \) is insufficient to cause the release of calcium \( Ca^{2+} \) from the Endoplasmic Reticulum (ER) due to degradation of the \( IP_3 \). Thus no \( Ca^{2+} \)-induced glutamate is released from the astrocyte. However, GABA also binds to the GABA-A receptors at the presynaptic terminal which has an inhibitory effect causing a low \( f_{\text{pre}} \). As \( f_{\text{pre}} \) is increased, as shown in Fig. 2B, more GABA binds to the GABA-B receptors and at some “trigger” frequency a sufficient amount of \( IP_3 \) is released to overcome \( IP_3 \) degradation and a low \( IP_3 \) threshold \( Th_{IP_3} \) is reached, causing \( Ca^{2+} \) released from the ER and a subsequent \( Ca^{2+} \) transient. The release of \( Ca^{2+} \) results in \( Ca^{2+} \)-induced glutamate release which subsequently binds to group I metabotropic Glutamate Receptors (mGluR) at the presynaptic terminal. It overcomes the inhibitory effect of GABA-A leading to an increase in PR of the presynaptic terminal, see Fig. 2B (green line). If \( f_{\text{pre}} \) continues to increase, the \( IP_3 \) within the astrocyte also increases and eventually reaches the upper threshold \( Th_{IP_3} \), as shown in Fig. 2C, and the oscillatory \( Ca^{2+} \) transient ceases. In this case glutamate is no longer released causing the inhibitory effect of GABA-A to be dominant again and the PR decreases at presynaptic terminal, see Fig. 3 (yellow line). Therefore, the interactions between the inhibitory GABA interneuron and the astrocyte can give rise to a frequency selective PR behaviour at tripartite synapses. This behaviour points towards a “band-pass” filtering characteristic for presynaptic spike trains where the passband will be closely related to cellular morphology, receptor distribution and many other aspects. Therefore, we assume that, as morphology and receptor density differs across synaptic sites, there will be a spread in the centre frequency of bandpass filters across the SANN, which will give rise to an advantageous SANN signal routing capability (This is discussed later in the paper). Effectively what is being filtered is the release of glutamate where with PR approaching unity, almost every spike arriving at a presynaptic terminal will cause the release of glutamate. Consequently, with the rapid arrival of glutamate in the cleft a strong STDP based learning phase occurs. However, as PR drops then fewer spikes release glutamate and the plasticity of synapses falls. Therefore, we propose that PR is effectively a local switch which can turn on/off learning at synaptic sites by modulating the height of the plasticity window: as \( PR \to 1 \) the plasticity window fully opens and with \( PR \to 0 \) the window closes. Section III-D of this paper proposes that activity at postsynaptic neurons also controls the plasticity of synapses.

To avoid the computational complexity discussed in Fig. 2, the interactions between GABA and the tripartite synapse can be approximated by a Gaussian function, which is shown by Fig. 3. It can be seen that the synaptic PR is a function of \( f_{\text{pre}} \), where the three stages of (a)-(c) correspond to the descriptions outlined in Fig. 2.

![Fig. 2. Signalling interactions between a tripartite synapse and a GABA interneuron. (A). A low \( f_{\text{pre}} \). (B). A high \( f_{\text{pre}} \). (C). \( IP_3 \) versus \( f_{\text{pre}} \).](image)

![Fig. 3. Spike frequency \( f_{\text{pre}} \) vs synaptic PR for a tripartite synapse and GABA interneuron. In the stage (a), the low \( f_{\text{pre}} \) is insufficient to cause the release of \( Ca^{2+} \) in the astrocyte, and no gliotransmitter is released. Thus the PR is low due to the inhibitory effect from the GABA interneuron. For the stage (b), if the \( f_{\text{pre}} \) is high, the \( IP_3 \) is sufficient (reaching the the low threshold \( Th_{IP_3} \)) to trigger the release of \( Ca^{2+} \). The gliotransmitter is released which overcomes the inhibitory effect of the GABA, thus the PR is increasing. In the stage (c), the \( f_{\text{pre}} \) is too high and causes the \( IP_3 \) reaching the upper threshold \( Th_{IP_3} \). It stops the release of \( Ca^{2+} \) and no gliotransmitter binds to the mGluR at the presynaptic terminal, which causes the inhibitory effect of GABA to be dominant again and the PR decreases.](image)
The Gaussian function in Fig. 3 is described by
\[
PR = e^{-\frac{(f_{pre} - f_s)^2}{2\sigma^2}}, \quad (1)
\]
where \( f_s \) is the sensitive (centre) frequency of the presynaptic terminal, \( \sigma \) is the width of Gaussian passband; both of these parameters are set by encoding scheme.

**B. Neuron Model**

Previous research have proposed several spiking neuron models, such as Hodgkin-Huxley model [27] and FitzHugh-Nagumo model [28], [29]. However, these models are computing-intensive as they used complex mathematics to describe the biological behaviours of the neuron [30]. Thus simplified models, including the Izhikevich [31] and Leaky Integrate-and-Fire (LIF) models [32], were proposed and are now widely used in many approaches [33]–[35]. In this paper, the simplified neuron model of LIF [32] is used, which is given by
\[
\tau_m \frac{dv}{dt} = -v(t) + R_m \sum_{i=1}^{n} I_{syn}^i(t), \quad (2)
\]
where \( v \) is the membrane potential of the neuron, \( I_{syn}^i \) is the injected current from \( i^{th} \) synapse and the total number of synapse is \( n \). The parameter descriptions of \( \tau_m, R_m \) and \( v_{th} \) and their values in this work can be found in Table I. The neuron goes to the refractory period when \( v > v_{th} \) and the refractory period is \( \sim 2 \) ms in this approach.

**C. Synapse Model**

In this work, a probabilistic-based model is used for the synapses, which is based on the synaptic neurotransmitter releasing mechanism similar to the approaches of [15], [36]. In this model when a pre-synaptic spike presents at the synapse, a random number generator generates a value \( \text{rand} \) (between 0 and 1) which is used to compare with PR of the synapse. If \( \text{rand} \leq PR \), the synaptic current is generated, which is described by
\[
I_{syn}^i(t) = \begin{cases} 
    r_I \ast w_i^{syn}(t)/n, & \text{rand} \leq PR \\text{rand} > PR 
\end{cases}, \quad (3)
\]
where \( \text{rand} \) is the generated random number, \( r_I \) is the current production rate, \( w_i^{syn} \) is the weight of the \( i^{th} \) synapse associated with neuron. All the parameters for the neuron and synapse models can be found in Table I.

**TABLE I**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Parameter description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \tau_m )</td>
<td>Neuron membrane time constant</td>
<td>240 ms</td>
</tr>
<tr>
<td>( R_m )</td>
<td>Neuron membrane resistance</td>
<td>1.2 ( \Omega )</td>
</tr>
<tr>
<td>( v_{th} )</td>
<td>Firing threshold value</td>
<td>9 mv</td>
</tr>
<tr>
<td>( r_I )</td>
<td>Synaptic current production rate</td>
<td>8</td>
</tr>
</tbody>
</table>

**D. BSTDP Learning Rule**

In this paper, a new learning algorithm with the capability of not only mapping new input data patterns to a desired output pattern, but also sensing when synaptic pathway(s) becomes dysfunctional and initiating re-learning or repair, was developed. If synaptic connections become broken or damaged, the corresponding currents injected into the neuron diminishes, which will reduce or even cause the firing activity of the postsynaptic neuron to cease. The BSTDP continually monitors the activity of postsynaptic neurons and when the level of activity of these neurons drops below a predetermined threshold a feedback signal initiates a localised learning phase. As learning process progresses, the activity of postsynaptic neuron increases which strengthens this indirect retrograde signalling pathway resulting in an increase in the astrocytic \( IP_3 \). With sufficiently high postsynaptic activity astrocytic \( IP_3 \) concentration will cross the upper \( IP_3 \) threshold and the \( Ca^{2+} \) transient ceases. Thereafter, no glutamate is released and PR falls off to a low background level resulting is shut down of learning: the plasticity window in (5) has collapsed. Hence as astrocytic \( IP_3 \) can be elevated from either direct signals by GABA interneurons or the indirect retrograde pathway originating from the postsynaptic neuron then both the pre and postsynaptic activity modulate the plasticity at synaptic sites. Consequently even if the frequency of the presynaptic neuron \( f_{pre} \) is fixed and falls within the bandpass range, as outlined in Fig.2, the plasticity window will only open if the postsynaptic neuron activity is below a predefined maximum. Effectively the height of PR in Fig.2 is modulated by the postsynaptic neuron frequency and this leads the authors to propose that the link between postsynaptic neuron activity, PR and synaptic plasticity may explain the biophysical processes underpinning the BCM learning function. Furthermore, there is strong experimental evidence for this feedback pathway where it has been shown [15], [26] that the retrograde messenger (endo-cannabinoids), which are synthesised in active postsynaptic neurons, act as an indirect signalling messenger, via astrocytes and the \( IP_3 \) pathway, that modulates PR, and hence plasticity, at local and distal synapses.

However, due to the computational overhead that would be incurred in modelling this feedback pathway and also its causal effect is adequately captured in the BCM function, we adopt the BCM model to map postsynaptic neuron activity directly to the plasticity of synapses. Hence, we now formulate a model for the BSTDP rule that merges conventional STDP learning with the proposed indirect pathway (BCM). The BSTDP rule updates the synaptic weights according to the timing difference between the pre and postsynaptic spikes, and is described by
\[
\delta w(\Delta t) = \begin{cases} 
    A_0 \exp\left(\frac{\Delta t}{\tau_+}\right), & \Delta t \leq 0 \\
    -A_0 \exp\left(-\frac{\Delta t}{\tau_-}\right), & \Delta t > 0 
\end{cases}, \quad (4)
\]
where \( \delta w(\Delta t) \) is used to update the synaptic weight, \( \Delta t \) is the time difference between presynaptic and postsynaptic spikes, \( A_0 \) is the plasticity window height, \( \tau_+\), \( \tau_- \) control the width of the plasticity window, and they are \( 40 \text{ms} \) in this approach.
Since the indirect feedback pathway appears to underpin the BCM rule we can use the established BCM function to model the relationship between postsynaptic activity and plasticity at synapses, as follows. The height of the plasticity window $A_0$ is modulated by a BCM type behaviour using the actual and target postsynaptic firing rates according to

$$A_0 = \frac{A}{1 + e^{a(f-f_0)}} - A_-, \quad \text{(5)}$$

where $A$ is the maximum plasticity window height, $A_-$ is the maximum plasticity window height for depression, $f$ and $f_0$ are the actual firing rate (i.e. a running average over forty seconds) and target firing rate of the postsynaptic neuron, respectively. The parameter $a$ controls the plasticity window opening and closing rates and was set to 0.2 in this work. All the parameters for the BSTDP learning rule can be found in Table II.

When $f \gg f_0$, then $A_0 \approx A_-$, which leads to long-term depression (LTD) and for $f \ll f_0$, $A_0 \approx A - A_-$, which gives rise to long-term potentiation (LTP). It is clear that if the postsynaptic neuron is in the early learning phase or the firing rate drops off due to the faulty synaptic connections, then the plasticity window height $A_0$ is positive (the learning window is open). This initiates a learning phase where the synaptic weights of the remaining non-dysfunctional synapses potentiate, eventually stabilising when the neural firing rate $f$ approaches $f_0$. Thus when faults occur and $f$ diminishes, this process will recover the postsynaptic neuronal firing rate to the pre-fault level. This is the self-repairing process which will be shown in the next section to maintain the functionality of the network when the synapses are damaged or broken. The results under various faulty conditions will also be provided to demonstrate this capability.

E. Self-repairing SANN architecture

The proposed SANN follows the architecture of feed forward neural networks. Fig. 4(a) is an example of the SANN architecture which includes an input, a hidden, and an output layers. Note that we have now modelled the interaction between the GABA neuron and the tripartite synapse as a Gaussian function: hereafter referred to as “Gaussian” synapses. The SANN uses multiple pathway connections, between neurons in different layers, with different delays for each pre and post neuron pairing, as shown by Fig. 4(b). Multiple pathways with delays are required to facilitate spatial data for the formation of a postsynaptic potential. Note that Fig. 4(a) is an example with only one hidden layer, but the SANN could also include multiple hidden layers.

In the proposed SANN information is passed from layer to layer in a forward direction where the input sensor data is encoded into spike trains (see later). Neurons in the middle and output layers have an associated Gaussian passband, implemented using (1) and (5), and therefore each neuron exhibits a unique “receptive field”. For example, consider the case for the SANN in Fig. 4(a) where there are $n$ unique input patterns. Each pattern is assigned to a single neuron in the hidden layer and therefore this layer will contain $n$ neurons where each of these neurons has a receptive field associated with one of the $n$ input patterns. This association is achieved using the centre frequency $f = f_s$ of the Gaussian function whereby a different $f_s$ value is used for each Gaussian synapse and this spread in $f_s$ will be sensitive to a unique input pattern: because of this spread in $f_s$ each of hidden layer neurons will be associated with one only possible input pattern. Hence in Fig. 4(a) the three input spike patterns can be accommodated with a corresponding mapping to the output layer neurons.

In real networks many processing neurons exist where groups of these neurons will be sensitive to different input patterns. Therefore, in these networks there is likely to exist neuron grouping that is receptive to a particular pattern where some will fire maximally to the pattern while others less so. It may even be possible to tune a neuron receptive fields as the shape of a receptive field is defined by the synaptic inputs to that neuron. These local or lateral connections are often inhibitory and by adjusting the associated weights it is conceivable that part of the transmitted information could be removed. However, this is beyond the scope of this paper and because of the computational overhead we restricted the number of hidden layer neurons by setting their receptive fields to match known input patterns.

IV. Simulation Results

The test bench setup is discussed followed by simulation results which show the performance of the SANN in an obstacle avoidance task. This task is employed as an example to demonstrate how the SANN can self-adapt to different fault conditions. The simulation results are provided from software simulations of the biological SANNs. Then the hardware acceleration is discussed in the final subsection which demonstrates the possibility of deploying the SANN to real-world robotic hardware platforms.

A. Example Application Setup

The obstacle avoidance data was collected using the Psi swarm robot shown in Fig. 5 which was developed by the York Robotics Laboratory, University of York, UK [37]. In this task, four infrared sensors were used with one at the front, one at each side (right and left) and one at the back where each sensor provided the sensory input data for the SANN controller, and the SANN output neurons gave the commands to drive the motors. The Psi swarm robot was deployed and the sensory data and corresponding controller decisions were recorded to local memory on the robot, and then downloaded.
after the experiments. Moving forward is the default action for the robot and it has the highest priority whereas moving backward has the lowest priority. Thus the action priorities are $F > R > L > B$, where $F/R/L/B$ represent moving forward, turning right, left and moving backward, respectively.

In this work sensors have a threshold level at which an object is detected and for sensory levels above this threshold the sensor value is mapped to a linear spike train frequency of 35 Hz, otherwise the mapping is to a frequency of 25 Hz. The output layer has four neurons corresponding to the four possible actions that can be taken by the Psi swarm robot to avoid obstacles. The target firing rates for the neurons in the hidden and output layers were arbitrarily set to 25 Hz and 10 Hz, respectively. Each synaptic connection has multiple pathways which are 8 and 16 for the connections in the hidden and output layers, respectively. Each synaptic pathway is delayed by 1 ms from the neighbouring pathway. Thus the minimum delay of a pathway is zero and the maximum is 7 ms for hidden and output layers, respectively; these delay times were arbitrarily chosen. The SANN architecture is given by Fig. 6 and each sensory neuron in the hidden layer generates an input spike train and in this simulation, the maximum number of possible input spike train in a pattern is 15: no action is needed when all sensors are activated as the Psi swarm robot is trapped.

### B. No Fault Condition

Consider the scenario where there is an obstacle directly in front of the robot (denoted by pattern A in Fig. 6) which would give rise to an input spike train of 35, 25, 25, and 25 Hz across all four sensory neurons. As this pattern matches the spread in $f_s$ across synapses associated with neuron #9 in the hidden layer then the plasticity window for these synapses will open and learning will be initiated and continue until neuron #9 reached a predefined firing frequency of 25 Hz. Fig. 7(a) shows that this is indeed the case where its firing rate increases gradually during the learning phase and eventually at around 170 s the firing rate of neuron #9 reaches 25 Hz and thereafter stabilises. The corresponding weights associated with neuron #9 potentiated during this learning phase, as shown in Fig. 7(b), and also stabilise at $\sim 170$ s because at 25 Hz the plasticity window closes and learning ceases. Activity at neuron #9 must be mapped to an action where the next highest priority is turning right, and we associated this action with neuron #2 in the output layer. Fig. 7(c) shows that the neuron #2 begins firing at $\sim 150$ s and this is because a period of learning is required before the weights associated with neuron #2 are potentiated to a level whereby the postsynaptic potential is sufficient to cause firing of neuron #2: note that STDP learning takes place for all synapses associated with neuron #2 even for subthreshold postsynaptic potentials due to Slow Inward Currents (SICs) as a result of $Ca^{2+}$ activity in the nearby astrocyte. The synaptic weights associated with neuron #2 are shown in Fig. 7(d) and again it can be seen that these weights are potentiated and stabilise after a period of learning; as neuron #2 firing rate approaches the target frequency of 10 Hz (Fig. 7(c)) the learning window closes and finally the associated synaptic weights stabilize. Additional results (not provided due to space) demonstrated that none of the remaining output neurons were active for this input firing pattern.

Fig. 7 shows the SANN learning process under the input pattern A, i.e. an obstacle is directly in front of the robot. When the input pattern changes to pattern B, obstacles are placed at both the front and right side of the robot. This pattern is uniquely associated, through the spread in the centre frequencies of the Gaussian synapses, with neuron #13 in the hidden layer and neuron #3 in the output layer. In the same way as for pattern A, synapses associated with both these neurons begin to potentiate until both neurons reach their target frequency and a similar learning process takes place for all other patterns. Therefore, after a period of training across all possible input patterns, the SANN will have implemented a mapping between input patterns generated by the sensory neurons and the output neurons. For the purposes of demonstrating the SANN under fault conditions we now
fix all the weights across the SANN at their post trained values and in the next section we present simulation results that demonstrate how the SANN maintains this mapping under fault conditions.

C. Fault Condition

Consider the case where pattern A is presented to the SANN, and at $t = 600$ seconds, seven of the eight pathways linking neuron #2 in the input layer with neuron #9 in the hidden layer developed faults: in this work we implement a fault on a pathway by permanently setting $PR$ to zero. At the same time eight of the sixteen pathways linking neuron #9 in the hidden layer with neuron #2 in the output layer also developed faults. This reduces the activity at both neuron #9 in the hidden layer and neuron #2 in the output layer. Fig. 8(a) shows the firing rate of the neuron #9 of hidden layer and it can be seen that this rate drops off to $\sim 20$ Hz from its target frequency of 25 Hz due to the introduction of the above faults. This frequency difference causes the plasticity window to reopen (modelled by (5)) and re-learning or repair commences (modelled by (4)). Fig. 8(b) shows that during this repair process the synaptic weights associated with neuron #9 potentiate over a “repair” period of 600 to $\sim 700$ seconds and thereafter remain stable as the firing rate of this neuron reaches its target value of 25 Hz. The drop in the firing rate of neuron #2 in the output layer (see Fig. 8(c)) is much more significant as half of the associated synaptic pathways are broken and, in addition, the output spike frequency of neuron #9 in the hidden layer has reduced due to damage in the hidden layer. Fig. 8(c) shows the recovery in the activity of neuron #2 to its pre-fault level (10 Hz) and again this occurs due to re-training which causes the remaining healthy (undamaged) synaptic pathways to potentiate, as shown by Fig. 8(d).

Now consider the case where faults are scattered randomly across the SANN with a fault density of 40%. The aim of this simulation is to evaluate how well the SANN maintains input-output mappings with a network-wide distribution of faults. Four different input patterns were presented to the SANN where each pattern activates a different output neuron. Fig. 9 shows the firing rates of all four output neurons where it can be seen that after faults occur, the firing rate of all four neurons drop significantly, but after a period of re-learning or repair these neurons become active again and eventually reach their target firing rate. Note that the recovery time of each output neuron is different and this correlates with the fault distribution: the causal effect of the drop in activity of a hidden layer neuron will be different across all fours output neurons. This results clearly demonstrate the ability of the
SANN to recover from a scattering of faults.

D. Fault densities variations

The fault tolerance capability of the SANN was also evaluated for different fault densities. The fault density is defined as the percentage of the dysfunctional (faulty) synapses against the total number of synapses. In this simulation, only pattern A is used as the input and faults were localised to pathways between the input sensory neuron #2 and neuron #9 in the hidden layer. Faults were set to occur at 300 seconds and two metrics were used to evaluate the self-repairing performance of the SANN: the lowest firing rate \( f_L \) of a neuron after a fault occurs, which reflect the severity of the faults on neuronal activity, and the fault recovery period \( T_R \) which reflects the recovery period or speed of recovery. The SANN was evaluated under fault densities of 20%, 40% and 80%. Fig. 10 shows the fault recover characteristics for neuron #9 in the hidden layer. It can be seen clearly from this plot both the neuronal activity and the repair duration correlate with the fault density.

Table III gives the \( f_L \) and \( T_R \) for neuron #9. As more synaptic connections become dysfunctional the longer the recovery time, as expected. However, despite the correlation between recovery time and faults density, the SANN does recover its functionality even under a high-density fault condition.

E. Hardware acceleration of the SANN

Note that the recovery time of 20-65 seconds in Section IV-D is in terms of biological time but it can be accelerated with hardware implementations. Based on the authors’ previous works of [4], [24], the FPGA implementation of an SANN can execute at clock speeds of 10-50 MHz, where one clock period is matched to one millisecond of the biological time step. Therefore, the recovery time of 65 seconds in biology would only require 6.5 ms when implemented on FPGA hardware, so this is within real-time constraints for many visual processing applications. This achieves an acceleration rate of \( 10^4 \) thus making this approach viable for real-time implementations of fault tolerance hardware. This is one direction of future work currently under investigation.

In summary, compared to conventional fault tolerance approaches, the proposed method has several advantages including: a) A distributed fault-resilient mechanism. The proposed SANN is a distributed neural network which does not require a
centre controlling unit and therefore the SANN demonstrates a higher degree of fault tolerance; b). Fine-grained fault-tolerant mechanism. The SANN supports repair at the level of synapses which, when compared to other more conventional approaches, is at a basic component or multi-transistor (logic gate) level. We view this as fine grained when compared to, for example, the approach in [38], where the entire column inside the FPGA device is reconfigured during partial reconfiguration (i.e., this is very much coarse grain repair); and finally c). Low cost/overhead where unlike the redundancy [13] (e.g., triple modular redundancy), error detection/correction [7], [11] methods etc., the SANN does not require additional components to detect and repair faults. Therefore, the proposed SANN architecture can provide a low cost, distributed, fine-grained fault-tolerant route to autonomous hardware.

V. CONCLUSION
A distributed self-repairing spiking astrocyte neural network with an information routing capability, underpinned by known interaction between an inhibitory GABA interneuron coupled with a tripartite synapse, has been presented in this paper. To minimise the computational overhead where interaction this interaction was realised in a Gaussian synaptic function which serves to route spike trains of different frequencies to different parts of the network. Furthermore, a biologically plausible link between postsynaptic neuron frequency and the probability of neurotransmitter release at synaptic sites was proposed and captured in the BSTDP learning rule, which combines the conventional STDP with the BCM rule. The paper has demonstrated that the rule provides a novel self-repairing capability which can not only establish an initial mapping between input data patterns and desired output response but also maintains this mapping in the presence of both localised or globally distributed faults. The fault-tolerant capability of the SANN was demonstrated in an obstacle avoidance task with results demonstrating that pre-learned input/output mappings can be maintained for fault densities up to 80% regardless of whether the faults are localised or globally distributed. Future work will apply the SANN to the domain of robotics for mission critical tasks and swarm applications. In these applications the robots’ functions can be compromised due to faults (such as caused by radiations, component ageing etc.), therefore fault tolerance is an extremely important capability in maintaining the key functionality of robotic and other hardware systems.

REFERENCES


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<th>Fault density</th>
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<td>20%</td>
<td>24.6</td>
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<tr>
<td>40%</td>
<td>20.5</td>
<td>35</td>
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<tr>
<td>80%</td>
<td>19.8</td>
<td>65</td>
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