Coping With Multiple Visual Motion Cues Under Extremely Constrained Computation Power of Micro Autonomous Robots

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This work was supported in part by the EU HORIZON 2020 Project Spatial-temporal information processing for collision detection in dynamic environments (STEP2DYNA) under Grant 691154; in part by the Ultra-layered perception with brain inspired information processing for vehicle collision avoidance (ULTRACEPT) under Grant 778062; in part by the China Postdoctoral Science Foundation under Grant 2019M662837; and in part by the National Natural Science Foundation of China under Grant 91648203, Grant 1913601, and Grant U1913205.

\textbf{ABSTRACT} The perception of different visual motion cues is crucial for autonomous mobile robots to react to or interact with the dynamic visual world. It is still a great challenge for a micro mobile robot to cope with dynamic environments due to the restricted computational resources and the limited functionalities of its visual systems. In this study, we propose a compound visual neural system to automatically extract and fuse different visual motion cues in real-time using the extremely constrained computation power of micro mobile robots. The proposed visual system contains multiple bio-inspired visual motion perceptive neurons each with a unique role, for example to extract collision visual cues, darker collision cue and directional motion cues. In the embedded system, these multiple visual neurons share a similar presynaptic network to minimise the consumption of computation resources. In the postsynaptic part of the system, visual cues pass results to corresponding action neurons using lateral inhibition mechanism. The translational motion cues, which are identified by comparing pairs of directional cues, are given the highest priority, followed by the darker colliding cues and approaching cues. Systematic experiments with both virtual visual stimuli and real-world scenarios have been carried out to validate the system’s functionality and reliability. The proposed methods have demonstrated that (1) with extremely limited computation power, it is still possible for a micro mobile robot to extract multiple visual motion cues robustly in a complex dynamic environment; (2) the cues extracted can be fused with a lateral inhibited postsynaptic network, thus enabling the micro robots to respond effectively with different actions, accordingly to different states, in real-time. The proposed embedded visual system has been modularised and can be easily implemented in other autonomous mobile platforms for real-time applications. The system could also be used by neurophysiologists to test new hypotheses pertaining to biological visual neural systems.

\textbf{INDEX TERMS} Bio-inspired, neural network, visual motion perception, micro robot, multiple cues.

\section{I. INTRODUCTION}

Computer vision has underpinned the rapid development of autonomous mobile robots in various applications, such as surveillance, transportation and manipulation [1]–[3]. A distinctive feature of computer vision is that its performance is strictly determined by the scale of available computational resources [4], [5]. In many remote robotics applications, such as ruin investigation [6], moon rovers [7] and underwater surveillance [8], micro mobile robots often play a unique role because of their size. However, conventional computer vision systems demand a highly capacious computational resource, and may not be readily incorporated into micro mobile robots for autonomous navigation. Novel approaches that function with less computational power are desperately needed for those applications.
Like robots, many animal species must address quite similar visual challenges in order to navigate (and survive) in a dynamic environment. These challenges are visual motion perceptions which, in animals, are perceived and computed by their extremely efficient visual neural systems. For example, in flying insects, their daily behaviour such as flight stabilisation [9], visual tracking [10], predator detection and avoidance [11], navigation [12] or landing control [13] are controlled mostly by visual cues. Neurophysiological and anatomical studies have shown that these behaviours rely on specific sensory neural pathways. Each of them is highly efficient in extracting specific visual motion patterns in dynamic scenarios with minimum energy costs or structural complexity.

Benefiting from the unique characteristics [14], bio-inspired computational visual perceptive neural systems have been extensively developed and gradually applied in the field of robotics to undertake rapid visual motion recognition tasks. For example, the lateral inhibitory neuron Lobular Giant Movement Detector (LGMD1 and LGMD2) [11], the Small Target Motion Detector (STMD) [15], [16] and the typical lobular plate tangential cells (LPTC) Elementary Motion Detector (EMD) [17], [18] are all highly effective visual neural systems that have been modelled and successfully emulated in small robot platforms.

Although individual neurons can exhibit a very high degree of selectivity for a stimulus produced in response to a specific feature, a single neuron is of limited utility for stimulus encoding [19]. One reason for this is that an individual neuron could exhibit sensitivity to multiple stimuli so that the output response could be mixed and lack precision; another reason is that single neuron’s performance fluctuates significantly according to different parameters and scenarios. To eliminate these influences in the vision system of some animals, multiple visual perceptive neurons with common structures and similar neural mechanisms coexist [9], [20], [21]. This mechanism is considered to be a critical prerequisite which enables these species to interact robustly with the dynamic complex real-world. For example, the LGMD1 and LGMD2 are a pair of wide-field detectors in the optic lobe of the locust’s which detect looming objects [22]–[25]. The latest research has shown that their neurophysiological behaviours vary significantly despite the fact that their morphological structures have only minor differences. For instance, the LGMD2 selectively responds to darker objects moving on a colliding trajectory against brighter backgrounds whilst the LGMD1 responds to all looming objects [11]. Nonetheless, it is clear that multiple visual cues extracted by different neurons could be the key to the agile reactions and decisions made by animals in relation to real-world dynamic events.

For micro mobile robots, it is critically important that they react to the dynamic environment with rich visual cues, with the highly restricted computing resources. To overcome these challenges, we propose in this paper a “Visual Motion cues Discrimination Neural Network” (ViMDNN) which functions effectively under extremely constrained computation power, and which enables a micro robot to detect and react autonomously to varied visual motion scenarios in real time. The proposed ViMDNN is constructed in two parts: a presynaptic visual cue detective array and a postsynaptic visual cues fusion neural network, followed by action triggering neuron groups to trigger reaction motion commands. The presynaptic neural array comprises four individual Visual Motion Perception Neurons (VMPN) that detect specific visual motion cues. Each VMPN contains a unified neural structure known as the Extended-LGMD (E-LGMD), which is an optimised neural prototype inspired by the insect LGMD neurons. In the postsynaptic part of the system, visual cues pass results to corresponding action neurons through the feature of lateral inhibition. A Translational Motion Cue Identification System(TMCIS) compares a pair of directional cues to generate translational cues, which are given the highest priority for reactive control, followed by the dark collision cues and approach cues.

The rest of this paper is organised as follows. An overview of related works is given in section II. The proposed models, including the E-LGMD and their postsynaptic connections, are described in section III. Section IV illustrates our experiments on both simulated platforms and real robots. In section V, we further discuss the proposed system and future research.

II. RELATED WORK

A. CONVENTIONAL METHODS OF VISUAL MOTION DISCRIMINATION

The majority of conventional methods for visual motion discrimination focus on distinguishing between the physical difference in the foregrounds and backgrounds, including their size, shape and texture details [26]–[28]. So far, preliminary visual motion perception techniques utilise three typical methods: geometric feature detection [29]–[31], background subtraction [32], [33] and optical flow [34], [35].

These methods are powerful in identifying specific objects in complex backgrounds. However, they are highly dependent on prior knowledge of the objects of interest. Even though some learning methods exist to increase the adaptability in unfamiliar and dynamic environments [36]–[38], the growing demand for computational resource is considered as an increasing challenge for real-time processing on a mobile platform [39].

B. BIO-INSPIRED VISUAL MOTION DETECTORS

Nature provides abundant methods to detect moving objects, which can be classified into looming detectors and lobular plate tangential cells (LPTCs) and Small target motion detectors (STMDs).

Nature provides abundant inspiration for detecting moving objects. The lobular plate tangential cells (LPTCs), LGMD and Directional Selective Neurons (DSN) [40] are generally regarded as typical neurons that respond to wide-field
motion and have been modelled into computational algorithms widely.

The lateral inhibition-based neural models have been applied to robots for many years [41]–[44]. The early models of LGMD1 typically included four layers [41], to which a fifth layer (grouping layer) was eventually added to increase robustness against cluttered backgrounds [43]. Several approaches have been proposed to simulate the selectivity between collision and receding motions [45], [46]. However, the biological basis has not yet been elucidated [11]. The study of LGMD2 by computational modelling commenced only recently and little work has been reported [44], [47]. Fu et al. proposed the first LGMD2 model using ON/OFF pathways and Spiking Frequency Adaptation (SFA) to achieve motion direction selectivity [44]. Unmanned Air Vehicles (UAV) are ideal lightweight platforms for LGMD1 and LGMD2 models [48], [49]. In [49], an event camera serves as input sensor to maximise computation efficiency.

There are also other types of bio-inspired neural models which, being simple but sensitive, have been deployed on light-weight platforms. For example, in fruit flies, the well-known LPTC model elementary motion detector (EMD) has been shown to be responsible for detecting translation motion [50]–[52] and has been applied on robots to enable course stabilisation and navigation [17], [53]. The Small Target Motion Detector (STMD) found in dragonflies and hoverflies is specifically sensitive to movements caused by dark objects with very small or limited size [15], and has shown potential in the development of small target automatic detection and tracking systems [16], [54]. Honey bees are widely studied for their flight control behaviour evoked by specific visuo-motor neurons [12], [55], [56]. There is also evidence showing that the praying mantis utilises individual neuro-mechanical visual-motor pathways to control prey-orienting movements [57]–[59]. Most of the above mentioned neural models exhibit single functionalities, that can barely distinguish between, or recognise multiple visual motion cues.

C. COORDINATION OF VISUAL MOTION DETECTING NEURONS

Multiple neural models can be integrated to achieve complex tasks. In [53], [60], [61], hybrid models combining both EMD and LGMD systems are proposed. A high-reliable flight control system for UAVs is proposed in [60]. However, these models serve two independent tasks: course stabilisation and collision avoidance. Thus, the system lacks exhaustive recognition of the current visual situation.

Several attempts have been made to combine bio-plausible lateral inhibition models to carry out tasks that require higher level recognition [40], [61]–[63]. As one variant model of LGMD, the DSN is sensitive to translational motion towards a specific direction. DSNs are commonly deployed in an array to identify whole-field translational motion using a competitive mechanism [40], which creates a translational selective neural network (TSNN). Further studies have investigated the effectiveness of TSNN combined with LGMD using a computational switching gene [61]. Fu et al. proposed a compound visual model that employs both LGMD1 and LGMD2 in competition within one robot to compare their different selectivities [64].

Hu et al. proposed a neural structure employing more than ten DSN-like neurons with their corresponding directions arranged in a circle to detect rotation motion detection [62] or even spiral motion [65].

Compared to visual models with single functionalities, these studies demonstrate the potential for recognising dynamic and complex visual motion scenes by integrating multiple bio-inspired visual motion detectors. However, the redundancy in structure could be a significant problem for constrained computing platforms such as micro robot platforms or UAVs.

III. MODELS AND METHODS

In this section, we present the proposed neural models in detail. The proposed visual system compromises of four subsystems, including 1) the image capturing and preprocessing; 2) visual motion perceptive neural array; 3) visual motion cues fusion neurons and 4) action neurons, as illustrated in figure 1. Image data is captured and preprocessed first, then transmitted into four individual VMPNs. In the following process, the visual motion cues fusion neurons arbitrate the right visual motion event and select the corresponding pathway to the action neurons. Finally, the results of visual motion perception are indicated by different reactive motor commands autonomously triggered by the micro robot in real-time.

![FIGURE 1. A schematic of the proposed ViMDNN model. Four VMPNs based on lateral inhibition model (LGMD1, LGMD2, DSNL and DSNR) that serve as visual motion perception neurons are deployed in this system. Their recognised individual visual motion cues are fused and integrated subsequently to generate responding motion commands by the action neurons. The visual cues fusion neurons arbitrate the corresponding output by the defined priority. The dashed arrows represent the inhibition.](image-url)

As revealed in biological studies, the perception and discrimination of multiple visual motion cues compromises of two stages: 1) the perception of individual visual motion patterns and 2) the fuse and selection of corresponding visual motion cue. In ViMDNN, excitations from all the four VMPNs are gathered and fused. The post-synaptic structure contains two parts: one TMCIS that compares the directional visual motion cues and one array of visual cues fusion
neurons. This bio-plausible structure may reflect an insect’s neural connections that use parallel pathways to process visual information and their mechanisms of natural behaviour generation mechanism [66].

**A. THE VISUAL MOTION PERCEPTIVE NEURONS**

As shown in figure 1, each VMPN model is responsible for a specific visual motion pattern, including the LGMD1/LGMD2 pair that detect approaching motion and DSNs that are sensitive to translational motion. In this work that ground mobile robot is utilised, the left/right translational motion is dominant (caused by ego-motion or objects passing by). In contrast, the up/down translational motion seldom happens. Thus only two DSNs are required. Along with the pair of LGMD to discriminate approaching objects, the final system is composed of four VMPNs, i.e. the LGMD1, LGMD2, left-sensitive DSN (DSNL) and right-sensitive DSN (DSNR).

1) **THE LAYER ARRANGEMENT OF E-LGMD**

The E-LGMD is a layered neural model formed by six layers and several individual neural processing cells and an auxiliary FFI pathway, as illustrated in figure 2. The six layers are the 1) Photoreceptor layer (P layer), 2) Excitation layer (E layer), 3) Inhibition layer (I layer), 4) Pre-Summing layer (pre-S layer) 5) Summing layer (S layer) and 6) Grouping layer (G layer). The luminance change of local individual single eye (pixel) is gathered by each cell in the P layer.

The excitation pass into the excitation layer without any delay; however, the inhibition layer accepts delayed signals according to the desired feature. Before the excitations and inhibition meet in the summing layer, an ON/OFF separator is inserted between the excitation/inhibition layers and the summing layer, thus forming a pre-S layer. Followed by the summing, a group layer is introduced for signal enhancement. The K cell gathers excitations in the G layers for generating E-LGMD spikes. The spikes of E-LGMD neuron are generated afterwards.

2) **THE PHOTORECEPTOR LAYER**

In locusts, the first layer of LGMD consists of photoreceptors that represent the excitations from the lamina, which are the luminance change. Similarly, in E-LGMD, the P layer is formed by grid-shaped P-cells that convey the difference between adjacent frames captured by the camera, with a residue part serving as the visual persistence effect. Respecting to the biological facts, all the VMPNs accept the same visual inputs from a single photoreceptor layer that a single P layer acts as their common input. The P layer at frame f can be defined as a matrix P(f):

$$P(x, y, f) = L_0(x, y, f) - L_0(x, y, f - 1) + \sum_{i=1}^{n_p} e^{iui} \cdot P(x, y, f - i)$$  \hspace{1cm} (1)

where f donates the frame index. $L_0$ is the grey scale image inputs from the camera, which represents the luminance value. $L_0(f)$ and $L_0(f - 1)$ are the current and last images. In the visual persistence part, $n_p$ is the confined steps of persistence. The visual persistence coefficient $\mu < 0$ determines the decaying speed. The index i indicates the last ith frames. To balance the performance and complexity, the depth of visual persistence $n_p = 1$, and the decaying coefficient $\mu = −2$. These parameters are selected empirically with consideration of a balance between the model’s functionalities and the compatibility for embedded environments.

3) **THE EXCITATION AND INHIBITION LAYERS**

As the core mechanism of E-LGMD modelling, the lateral inhibition is accomplished by utilising two types of layers that exhibit reversed features. The excitation layer (the E layer) holds all the current visual motion cues, which are directly retrieved from excitations in the P layer.

$$E(x, y, f) = P(x, y, f)$$  \hspace{1cm} (2)

The inhibition layer spread inhibitory features with short latency:

$$I(x, y, f) = \omega_l \cdot \sum_{i=-r}^{r} \sum_{j=-r}^{r} P(x, y, f - 1) \cdot W_l(i + r, j + r)$$  \hspace{1cm} (3)

where $\omega_l$ stands for the inhibition coefficient, $W_l$ is the inhibition kernel pattern. The variation in $W_l$ represents whether the selectivity is omnidirectional or directional,
TABLE 1. Configurations set for the applied VMPNs.

| Instance | ncell  | $\omega_f$ | $W_f$ | $g_{on}$ | $g_{off}$ | grouping | $C_\omega$ | $t_{de}$ | $C_\alpha$ | $C_\beta$ | $t_{spikes}$ | $t_{pre}$ |
|----------|--------|------------|-------|----------|-----------|----------|-----------|--------|-----------|-----------|-------------|-----------|
| LGMD1    | 7200 (100x72) | -0.4 | $\begin{bmatrix} 1 & 1 & 1 \\ 1 & 0 & 1 \\ 1 & 1 & 0 \end{bmatrix}$ | 1 | 1 | yes | 4 | 30 | 10 | 11 | 100 | 80 |
| LGMD2    | 7200 (100x72) | -0.4 | $\begin{bmatrix} 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 \\ 1 & 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 \end{bmatrix}$ | 0 | 1 | yes | 4 | 30 | 10 | 11 | 100 | - |
| DSNL     | 5600 (80x70)  | -0.53 | $\begin{bmatrix} 1 & 1 & 1 & 1 & 1 \\ 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \end{bmatrix}$ | 1 | 1 | no | - | 20 | 17 | 48 | - | - |
| DSNR     | 5600 (80x70)  | -0.53 | $\begin{bmatrix} 1 & 1 & 1 & 1 & 1 \\ 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \end{bmatrix}$ | 1 | 1 | no | - | 20 | 17 | 48 | - | - |

which determines the feature of formed VMPN. The $r$ indicates the size of applied inhibition pattern $W_f$. When pattern size is $3 \times 3$, $r = 1$, and for size of $5 \times 5$, $r = 2$.

The applied inhibition kernels meet the list in Table 1. Notice that for LGMD1/LGMD2 pair, the kernels are symmetrical, while for DSNs, horizontal biases are applied.

4) THE PRE-SUMMING LAYERS

In the earlier modelling work of LGMD, the lateral inhibition is performed in the $S$ layer, where the excitation from $E$ cells and inhibition from $I$ cells meet. The $S$ layer in the earlier model is calculated by:

$$S_{RAW}(x, y, f) = E(x, y, f) - I(x, y, f)$$  \hspace{1cm} (4)

in which the $S_{RAW}(x, y, f)$ donates the raw $S$ cells in earlier simplified models [43], [64]. However, this method should be improved since it cannot separate the signal onset and off-set, which could be the major difference between LGMD1 and LGMD2. In the pre-$S$ layers of the proposed model, the ON/OFF excitations are processed individually. Moreover, concerns about the overflow issue, an anti-overflow limiter is applied to each of the layers:

$$S_{ON}(x, y, f) = [E(x, y, f), E(x, y, f) - I(x, y, f)]^\oplus$$  \hspace{1cm} (5)

$$S_{OFF}(x, y, f) = [E(x, y, f), E(x, y, f) - I(x, y, f)]^\ominus$$  \hspace{1cm} (6)

in which the $S_{ON}(x, y, f)$ and $S_{OFF}(x, y, f)$ stands for the on-channel and off-channel respectively. The anti-overflow limiter ensures that the output amplitude of pre-$S$ cells does not exceed the input of $E$ cells. A simple illustration of how excitation and inhibition affect the pre-$S$ layers are shown in figure 3

5) THE SUMMING LAYER

After joining the excitation and inhibition together into the pair of pre-$S$ layers, the final $S$ layer is regulated by a switching mechanism.

$$S(x, y, f) = S_{ON}(x, y, f) \cdot g_{on} + S_{OFF}(x, y, f) \cdot g_{off}$$  \hspace{1cm} (7)

in which the $g_{on}$ and $g_{off}$ are switching parameters that are either 0 or 1. The values for each VMPN are listed in table 1. Notice that the LGMD2 has turned off the ON channel to reject on-sets.

6) THE GROUPING LAYER

The $G$ layer of the VPMN enhances spatial contrast. The process contains two steps.

In the spatial enhancement process, a passing coefficient, which is determined by local excitation strength, is set for each local cell. The array of passing coefficient $Ce$ is computed by a 2-D filter:

$$Ce(f) = \sum_{i=r}^{r} \sum_{j=r}^{r} S(x, y, f) \cdot W_G(i + r, j + r)$$  \hspace{1cm} (8)

in which $W_G$ is the influence from neighbouring cells, which can be described as:

$$W_G = \frac{1}{9} \begin{bmatrix} 1 & 1 & 1 \\ 1 & 1 & 1 \\ 1 & 1 & 1 \end{bmatrix}$$  \hspace{1cm} (9)

The spatial enhanced $G$ layer is then obtained by:

$$G(x, y, f) = S(x, y, f) \cdot Ce(x, y, f) \cdot \omega(f)^{-1}$$  \hspace{1cm} (10)
where $\omega$ is a scale that measures the passing coefficient intensity of the whole frame:

$$\omega(f) = 0.01 + \max \left| C_\omega \cdot C^{-1}_{\omega} \right|$$  \hspace{1cm} (11)

where $C_\omega$ is a certain value.

In the second step, small signals are blocked by a certain threshold:

$$\tilde{G}(x, y, f) = \begin{cases} G(x, y, f) & \text{if } G(x, y, f) \geq t_{de} \\ 0 & \text{otherwise} \end{cases}$$  \hspace{1cm} (12)

where the $t_{de}$ is the constant threshold that filters out small signals. For modelling DSN, only the second process (eq.12) is involved since grouping will increase undesired non-linearity to later process in TMCIS.

7) THE SPATIAL INTEGRATION NEURON

The spatial integration neuron (K cell) in this VMPN model represents the average excitation levels in the previous layer. It first gathers neural excitations from all cells in the previous layer:

$$k(f) = \sum_x \sum_y \left| \tilde{G}(x, y, f) \right|$$  \hspace{1cm} (13)

Following by that, the value $k_f$ is transformed by a normaliser:

$$\kappa(f) = \frac{1}{2} \cdot \tanh \left( \frac{\sqrt{k(f)} - n_{cell} \cdot c_a}{n_{cell} \cdot c_\beta} \right) + \frac{1}{2}$$  \hspace{1cm} (14)

where the $c_a$ and $c_\beta$ are constants that can shape the normalising function with varied gain settings for small and big signals. The $n_{cell}$ is the number of element cells in the $G$ layer.

The structures followed by this K cell is dependent on the actual model. In this work, the output of four utilised VMPNs are represented by $k_1(f), k_2(f), k_L(f)$ and $k_R(f)$ respectively. For the LGMD1/LGMD2 model, $k_1(f)$ and $k_2(f)$ cells are followed by the spiking mechanism that represent a prominent approaching event. For DSN modelling, the comparison of $k_L(f)$ and $k_R(f)$ will be described later for the TMCIS modelling.

8) THE SPIKING MECHANISM FOR LGMD1/LGMD2

In E-LGMD process, if the spatial integration neuron exceeds a determined threshold, a spike is produced:

$$s_{\text{spike}}(f) = \begin{cases} 1 & \text{if } \kappa(f) \geq t_{\text{spike}} \\ 0 & \text{otherwise} \end{cases}$$  \hspace{1cm} (15)

An impending collision is confirmed after several successive spikes. Longer decision time lead to solid and reliable outputs, but it is harder to respond to sudden and fast collision events. In our tests, the decision time is set to 3-6 frames (100-200 ms) for best performance.

9) THE FFI PATHWAY OF LGMD1

In LGMD1 circuitry, an auxiliary feed-forward inhibition pathway is responsible for detecting large and sudden whole-field visual motion, which could be introduced by self-rotation or other unknown significant motion. The FFI neuron and lateral inhibition work together to prevent false spiking alarms in these situations. When the VMPN is configured into other neuron types, the FFI mechanism is bypassed.

The FFI cell is proportional to the average excitation level of $E$ layer with one frame delay:

$$k_{\text{FFI}}(f) = \sum_x \sum_y \left| E(x, y, f - 1) \right| n_{cell}^{-1}$$  \hspace{1cm} (16)

A constant threshold is set to enable the FFI mechanism. This threshold should be high enough that it will not interfere with normal reactive control.

$$s_{\text{spikeFFI}}(f) = \begin{cases} 1 & \text{if } k_{\text{FFI}}(f) \geq t_{\text{FFI}} \\ 0 & \text{otherwise} \end{cases}$$  \hspace{1cm} (17)

10) THE PARAMETERS SETTING

By setting the unified E-LGMD models with different parameters, they can be initialised into varied VMPNs with unique functionalities. This facilitates the implementation of multiple models with little extra ROM occupation. The parameters’ configuration of applied VMPNs are described in table 1.

Currently, there is no learning or adaptive mechanism applied in E-LGMD thus most of the parameters are empirically fixed considering the functionalities of designed models. However, there are several methods developed to optimise the parameters [67]. As mention before, some parameters are selected for better compatibility with embedded environments. Notice that the $W_j$ are fractions with denominators of 4 or 8. Meanwhile, some parameters are determined by the physical properties of the model such as the value of $n_{cell}$ and the gating parameters.

B. THE POSTSYNAPTIC NEURONS

As a higher-level architecture, the ViMDNN coordinates multiple VMPNs according to their functions and generates the right behavioural response to each type of visual challenge. Therefore the robot with the ViMDNN can cope with different visual stimuli reliably. If only with single function model, for example, only with LGMD, the robot will not be able to respond to translating objects correctly; only with DSNs, it can hardly cope with collision events. The ViMDNN provides the right structure for a robot to initiate the right response to moving visual cues in the real world.

However, different VMPNs cannot simply be integrated together. The LGMD produce intensive spikes when an object approaches. This spiking mechanism contributes to rapid and solid results but with strong non-linearity. On the contrary, the DSNs are tuned to be sensitive to certain motion directions, although they also exhibit minor excitations to approaching objects or even translating objects towards other directions. Therefore, the DSNs must be merged and...
transformed into spiking outputs before they meet the LGMD neurons. The TMCIS is deployed to achieve this goal.

1) THE TRANSLATIONAL MOTION CUE IDENTIFICATION SYSTEM

For flying insects such as fruitflies, two steering strategies are applied, which are course stabilisation and obstacle avoidance. Required visual information could be obtained by comparing between left and right visual motion trends through the visual motion detector EMD [68]. It is demonstrated that when triggering saccade reactions, the direction of saccades (left or right) is opposite to the side who experience larger visual motion. This approach has also inspired several controlling models in robotics [69], [70] with EMD neurons.

Similarly, a TMCIS based on E-LGMD neurons can provide directional information for steering control of robots, compromising results from the divergence of a pair of DSNs. The output of TMCIS represents the dominating motion direction. The structure of TMCIS and the comparison to fly’s Spatial-temporal Integration if Motion (STIM) model is illustrated in figure 4.

![Figure 4](image_url)

**FIGURE 4.** (a) The spatio-temporal integration of motion (STIM) model inspired by fruitflies (figure adapted from [71]). (b) A schematic of the proposed TMCIS model. Notice the difference of inhibition masks in DSNL and DSNR.

The basic idea of obtaining the directional motion is to compute the divergence of the DSN cells directly:

\[ d_{RAW}(f) = \kappa_L(f) - \kappa_R(f) \]  

(18)

in which the output value \(d_{RAW}(f)\) contain the information of both the direction and strength of translational motion. However, due to the noise introduced by calculating DSN, this approach is not practical and should be improved by further noise reduction and gating techniques.

Here in TMCIS, it evaluates the divergence between the pair of DSNs, namely \(d_{TMCIS}(f)\). First, the output of both DSNs are filtered by a single-pole recursive filter (also can be regarded as a leaky integrator) to reduce noise:

\[ \tilde{\kappa}_L(f) = \kappa_L(f - 1) \cdot \eta_L + \kappa_L(f) \cdot (1 - \eta_L) \]  

(19)

\[ \tilde{\kappa}_R(f) = \kappa_R(f - 1) \cdot \eta_R + \kappa_R(f) \cdot (1 - \eta_R) \]  

(20)

where \(\eta_L\) and \(\eta_R\) are the decaying factors within range \((0,1)\).

A smaller value contributes to a lower filtering strength.

A greater value contributes to smoother output but with longer latency. This is a practical approach of low-pass filters for micro-controllers. In TMCIS, both decaying factors are empirically set to 1/7.

Then the divergence of two DSNs \(d_{TMCIS}(f)\) is calculated by:

\[ d_{TMCIS}(f) = \frac{(\tilde{\kappa}_L(f) - \tilde{\kappa}_R(f))^2}{\sqrt{\tilde{\kappa}_L(f)^2 + \tilde{\kappa}_R(f)^2}} \cdot \text{sign}(\tilde{\kappa}_L(f) - \tilde{\kappa}_R(f)) \]  

(21)

where the strength of the divergence \(|d_{TMCIS}(f)|\) shows the strength of dominate direction of motion. This operation slightly compresses small signals to increase the robustness of direction detection, because a small signal means the \(\kappa_L\) and \(\kappa_R\) grows simultaneously, which is not the case of translational motion. A simple illustration of this mechanism is shown in figure 5.

![Figure 5](image_url)

**FIGURE 5.** The illustration of obtaining \(d_{TMCIS}\) from the outputs of DSNs. Notice that the output is suppressed (flattened) along the line: \(\kappa_L - \kappa_R = 0\).

\(a: THE COMPARING CELL\)

A solid decision of translational motion is made when \(d_{TMCIS}(f)\) has steady output. A tri-state hysteresis neuron is applied here to reduce sudden and frequent fluctuations. The tri-state hysteresis neuron has two constant on-set thresholds: one on each side and an off-set threshold zero, which contribute to three outputs: left, right and centre. They are represented by \(-1, 1\) and 0 respectively. The decision of a certain direction is selected when the input strength (absolute value) is greater than the on-set threshold, however, it will return to the centre state once the input strength drops below the off-set threshold and cross to the other side. When the value is between two thresholds, the decision remains unchanged:

\[ T_{Dir}(f) = \begin{cases} 
1(\text{right}) & \text{if } d_{TMCIS}(f) > t_{TR} \\
-1(\text{left}) & \text{else if } d_{TMCIS}(f) < t_{TL} \\
0(\text{centre}) & \text{else if } d_{TMCIS}(f) \cdot T_{Dir}(f - 1) \leq 0 \\
T_{Dir}(f - 1) & \text{otherwise} 
\end{cases} \]  

(22)

in which \(T_{Dir}(f)\) is the decision at frame \(f\). The \(t_{TR}\) and \(t_{TL}\) are two thresholds for the left side and right on-set (\(t_{TR} > 0, t_{TL} < 0\)). In this work, the values are set to \(t_{TR} = 4\) and \(t_{TL} = -4\).
b: THE CROSS PARTIAL IMAGE CONFIGURATION

Consider the situation that a right motion happens in the rightmost part of the view. It is usually not hazardous since the object has almost moved outside the field of view. Similarly, the leftmost part of the view can be skipped when judging a hazardous left-moving object. Thus, the input of DSNs can be tailored to increase selectivity and reduce processing time. The DSNL takes only the right part of the input image into the process, and the DSNR calculates only left part. In TMCIS model, the overlap of 60% provides good separation, as shown in figure 4. A simple test of TMCIS with video stimuli is shown in figure 6.

2) THE VISUAL CUES FUSION NEURON

As illustrated in figure 1, the proposed visual cues fusion neurons collect visual motion cues from VMPNs and the TMCIS, then generate a single output directly to the action neurons. The extracted visual motion cues could be either of: 1) safe, 2) approaching motion, 3) approaching motion caused by dark objects, 4) translational motion to the left and 5) translational motion to the right.

The visual cues fusion neurons use a prioritised preemption mechanism to organise the orders of input neurons. In this work, five cells representing possible identified visual motion are assigned with defined priority to connecting presynaptic neurons, including a default cell. A neuron can inhibit other neural excitations whose priority is lower to presynaptic neurons, including a default cell. A neuron can inhibit other neural excitations whose priority is lower to presynaptic neurons, including a default cell. A neuron can inhibit other neural excitations whose priority is lower to presynaptic neurons, including a default cell. A neuron can inhibit other neural excitations whose priority is lower to presynaptic neurons, including a default cell.

The time used for each process also depends on the scale of active neurons. For each E-LGMD, even though five layers such as excitation transferring and neighbouring operations, which make it a desired model for embedded platforms with constrained computational resources. However, it is still not an easy task for adapting this neural structure successfully.

1) THE ROBOT PLATFORM

A micro ground robot Colias-IV is selected, as shown in figure 7, to be the application platform to demonstrate the high computing efficiency and low-hardware demanding. The Colias-IV which has constrained computation resources is a low-cost vision-based micro ground robot developed for swarm robotic applications and bio-robotic research [72]. The Colias-IV employs a circular body with 4cm in diameter that has two layers of circuit boards. The bottom part serves as an only primary robotic platform such as power supply, motion control and primary sensing. Driven by a pair of differentially driven wheels and a third pin-stand in front, the Colias-IV can run up to 35cm/s at full speed. The upper part, which called the Colias-Sensing-Unit (CSU), is designed for onboard high-level sensing such as vision. The CSU contains a tiny CMOS camera as a visual sensor that captures RGB image sequences at 30 frames per second (fps). The field of view angle (FOV) is about 70 degrees horizontally. The CSU utilises an STM32F427 microcontroller as the main processor, which is running at 180MHz. It contains 256KBytes internal RAM space and 2MBytes ROM space.

2) THE COMPUTATIONAL COMPLEXITY ANALYSIS

For embedded programming, both spatial complexity and temporal complexity are critical issues and should be optimised. In addition to the basic principle of optimising ELGMD-based model described in [42], the $P$ layer is regarded as a common part for all the VMPNs for further memory saving. Thus the $P$ layer can be stored separately and calculated only once for each frame. Regarding the calculation procedure, two consecutive $P$ layers should be accessible for implementing delayed inhibition, and for differential image calculation and temporary storage, three input buffers are allocated. Due to the maximum available RAM space, the utilised scale of the input image is 100x72. The breakdown of total occupied memory space is shown in figure 8.

The time used for each process also depends on the scale of active neurons. For each E-LGMD, even though five layers...
exist, calculations for some layers can be omitted or merged in a single loop to save time. As briefly illustrated in algorithm 25 and figure 9, the calculations of I layer and S layers are combined into one loop, the calculation of Ce map and the G layer are also combined. Meanwhile, the boundary of convolution is omitted that the time cost for convolution operation can be further reduced by skipping the edge cells. Thus, the computation process of the full process mainly consists of three loop phases: 1) the P layer, 2) I layer and S layer of each VMPN, and 3) the loop for G layer and spatial integration for each VMPN. When the input image has the size of $H_{in} \times W_{in}$ pixels and convolution kernel size of $k$, the computational complexity of each phase can be estimated as $N_P = O(H_{in} \cdot W_{in})$, $N_{IS} = O(k^2 \cdot (H_{in} - 2) \cdot (W_{in} - 2) + H_{in} \cdot W_{in})$ and $N_G = O(k^2 \cdot (H_{in} - 4)(W_{in} - 4))$, where $N_P$, $N_{IS}$ and $N_G$ represent time for the three phases.

$$N_{VMPNs} = N_P + 2N_{LGMD} + 2N_{DSN}$$

(23)

where $i$ represents the index of utilised VMPNs. A test of time consumption in different image scales is shown in figure 10a. The timing profile of utilised model that has a scale of $100 \times 72$ is shown in figure 10b.

As a result, these further optimisations, the total time used for processing all the modules including four E-LGMD structures has reached $23 \pm 2$ ms, which is within the duration of one single frame (typically 33ms). This indicates that the proposed compound visual motion detection system can be performed in real-time on the Colias-IV platform.

The power consumption of the model is mainly dependent on the MCU’s clock frequency due to the architecture of the embedded processor. In its typical condition (180MHz), the power consumption of MCU is around 76mA with the capability of processing 225 Dhrystone Million Instructions Per Second (DMIPS). The power consumption breakdown is briefly listed in table 2.

### IV. EXPERIMENTS AND RESULTS

The performance of proposed ViMDNN is evaluated in systematic experiments including basic functional tests with virtual visual stimuli, real-world challenges and simple autonomous control scenarios.
FIGURE 10. Timing profile of real-time processing on the embedded processor. (a) The measured time cost in different phases per cell when the size of input image changes, showing the measured results of LGMD1 and DSNL as examples. The x-axis is in log scale. Notice that in the most efficient situation, the input size should be 400-600 pixels. (b) The timing profile observed and logged by a multi-channel oscilloscope (Sampling Rate 1GHz) using 100×72 cells. The time measuring labels are added manually. In the duration of a full frame (33 ms), four VMPNs are processed successively after the P layer calculation, and the whole execution is completed within 24ms. The change of calculation phases is indicated by the signal flips.

TABLE 2. The power consumptions of major components.

| Module Name & Description | typical | max | unit |
|--------------------------|---------|-----|------|
| MCU standby *            | 2.5     | 6   |      |
| MCU core (150MHz) **     | 46      | 52  |      |
| MCU peripherals (150MHz) | 13      | 15  |      |
| MCU core (180MHz)       | 57      | 65  |      |
| MCU peripherals (180MHz) | 19      | 25  |      |
| Camera standby           | 0.8     |     | mA   |
| Camera active            | 12      | 18  |      |
| CBU (without motor)      | 20      | 75  |      |
| DC Motor x2              | 60 ***  | 300 |      |

1) SIMULATED APPROACHING OBJECTS
Firstly, we would like to compare the selectivity between LGMD1 and LGMD2 when an object is on the approaching trajectory, as shown in figure 11a. When challenged by a dark approaching object (figure 11e up-left), both LGMD1 and LGMD2 generate increasing membrane potential rapidly as the object expands in the retina. When the object is bright while the background is dark (figure 11e up-right), the LGMD2 neuron is inhibited during the whole approaching process as expected. The reactions of DSN towards virtual approaching visual stimuli are also tested, as shown in figure 11b. Since the object on an approaching trajectory donates brightness change on both sides evenly, it is clear that both DSNs generate spikes evenly and show no direction preference on any object/background contrast group.

2) SIMULATED TRANSLATING OBJECTS
When the simulated visual stimulus is moving at translating trajectory (figure 11e down), the LGMD1 and LGMD2 neurons show only constant and weak outputs during the whole process for either dark or bright stimulus (figure 11c). In this situation, the peak values for both motion directions (left and right) is almost the same, indicating that the pair of LGMD neurons is not sensitive to motion directions.

Meanwhile, the outputs of both DSNs differ along with the motion direction. For the motion from left to right regardless of the object/background contrast, the DSNL is weaker than DSNR, and vice versa. The divergence between these two DSNs, which is close to the TMCIS output described in the above section, have almost the same strength for motion in two opposite directions. The results demonstrated the functions of biased inhibition mechanism set in the I layer.

The results of tests on simulated visual stimuli with different object/background contrast set are compared and concluded in figure 12. The results shown in figure 12a show that LGMD2 only detects an approaching object when it is darker than the background. Moreover, the sensitivity of proposed LGMD1 and LGMD2 structures decrease when the object becomes more obscure against the background, resulting in a shorter alarm time for responding reactions.

The peak output of DSNs in the object translating scene has shown stable and distinctive selectivity within a single motion scene, as illustrated in figure 12b. It shows that the sensitivity of each DSN is related to the object/background contrast. The clearer the object stands out from the background, the higher values generates. Results of additional tests on translating objects from the opposite side have shown little difference in the peak value, except the swapping of places. This result reveals that the peak value of DSN is related to the contrast with the given direction, but the polarity of motion does not affect the performance of DSN, which is different from the LGMD2 model.

Figure 12c has drawn a relationship between the time to collision (TTC) and the motion speed. When the motion gets faster, the TTC decreases, which result in a shorter

A. RESPONSES OF VMPNs TO VIRTUAL VISUAL STIMULI
The proposed VMPN models are firstly tested by virtual visual stimuli to demonstrate their fundamental features. The computer-generated visual stimuli are categorised into two groups: 1) an approaching object that has an expanding edge and 2) a translating object that moves from left to right with constant speed, then reverse. Each group of visual stimuli is set with different object-background contrasts $C_{obj}$, which is defined by:

$$C_{obj} = B_{obj} - B_{back}$$ (24)

where $B_{obj}$ is the brightness value of moving object, $B_{back}$ is the brightness of background. The brightness values are within the range [0,1]. No additional noise is added to the image stimuli. Experiment stimuli, explanation and sample data logged from the robot are shown in figure 11.
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FIGURE 11. The experiments of VMPNs challenged by computer-generated visual stimuli. (a) & (b) The output of both LGMD1, LGMD2 DSNL and DSNR when the virtual stimulus is on an approaching trajectory. (c) & (d) The output of all VMPNs when the virtual stimulus is translating from one right to left, then left to right. (e) The explanation of applied virtual visual stimuli. The colour combination of both background and foreground are tested in groups.

FIGURE 12. Analysis of experiments on virtual stimuli. (a) The TTC of both LGMD1 and LGMD2, when challenged with approaching objects under different object/background contrast settings. Notice that the LGMD2 only respond to dark objects in front of a bright background; (b) The peak value of DSNs when challenged with translating objects from left to right. Showing the standard errors of logged data during the progress; (c) The TTC of LGMD1 and LGMD2 when the moving speed of approaching object changes. (d) The peak value of DSNs when challenged with translating objects with different motion speeds, where $C_{\text{obj}} = -1$. Showing the standard errors with errorbar.

responding time for an agent to take actions to avoid an impending collision.

The relationship between DSN outputs and the translating motion speed is also tested, as shown in figure 12d. The results show that both the DSNs output increase along with the moving speed, but selectively gradually saturates when the speed of the object increases.

B. EXPERIMENTS WITH REAL-WORLD STIMULI

Comparing to computer-generated virtual stimuli, the visual scene in real-world is noisy, caused by unwanted flashings or shadows cast by poor illumination. Moreover, the object’s motion speed cannot be strictly maintained as video do. Therefore, the performance of proposed neural models should be tested in the “real” dynamic scenes.

1) CHALLENGE WITH ROLLING BALLS
To test the postsynaptic visual cues fusion neurons and demonstrate the ability to distinguish different motion patterns successfully, a rolling ball is used as the visual stimulus in the following experiments.

The experimental configuration used in this test is shown in figure 13. Several billiard balls with four different pure colours are used as visual stimuli, which the variety in colours contribute to different brightness levels in the luminance channel ranging from totally black to very light grey. The object’s trajectory is defined by the approaching angle against the robot’s median plane $\theta$ and a determined safe range $d$. The object’s speed $v$ is approximately 20 cm/s when crossing the robot’s median plane. Sets of experiments are conducted in nine groups of motion trajectories, which are listed in table 3. For each group of motion trajectories, experiments are repeated for 20 times. During the object moving process, the recognised visual motion cues from the compound model are logged and compared in figure 14. The result of motion pattern detection is shown in figure 15 and figure 16.

The neural outputs during the tests are measured, and the response details in some typical scenarios are illustrated in figure 14. In the results, the performances of designed VMPN models are consistent with our expectations. Both LGMD1 and LGMD2 can recognise dark approaching objects and produce spikes accordingly, but the LGMD2 produces slight excitation when a bright object is observed. When the object is near-miss but not on a head-on collision trajectory, both LGMD1/LGMD2’s response cannot reach
TABLE 3. Experiment configurations of looming balls.

| scene | d (cm) | v (cm/s) | θ (°) | description |
|-------|--------|----------|-------|-------------|
| 1     | -      | 20       | 0     | head on collision |
| 2     | 50     | 20       | ±15   | near miss |
| 3     | 10     | 20       | -45   | cutting through (right) |
| 4     | 10     | 20       | 90    | translating (right) |
| 5     | 10     | 20       | 135   | overtaking (right) |
| 6     | 10     | 20       | -45   | cutting through (left) |
| 7     | 10     | 20       | -90   | translating (left) |
| 8     | 10     | 20       | -135  | overtaking (left) |
| 9     | 35     | 20       | 90    | translating far away (right) |

FIGURE 14. Model variables and outputs during a typical experiment with real-world stimuli, showing the VMPNs’ outputs (upper plot), TMCIS output $d_{\text{TMCIS}}$ (middle plot) and spiking outputs (lower plot). In the output plots, horizontal dashed lines indicate the threshold for shown models. Vertical dashed lines indicate the time when a collision happens, or the stimulus cross the median plane. The decision of recognised motion is made by the uppermost spiking neuron shown in the spiking plot.

FIGURE 15. The performance of motion pattern recognition evoked by a head-on colliding object with varied colours. (a) The counted results of recognised motion patterns of the head-on colliding object. The CORRECT decision is marked with a blue bar. (For white and pink objects, the bright approaching results are considered as “correct”. For red and black objects, dark approaching are considered as “correct”). The maximum chance of correctly recognise a motion pattern is 1 (100%). (b) The ratio of detected bright approaching and dark approaching results vs. different ball colours.

In figure 16, the results of challenges with a single black ball but on different translating paths are shown. For objects moving not far away from the robot, most of the translating motion can be recognised and separated correctly. However, for the motion happening far away from the robot, the scene has a higher chance to be treated as safe rather than a translational motion. Notice that in some cases, the ball suddenly appears and passed by, an unknown scene might be prompted due to the excitation of FFI cell in LGMD1 brought by the fast brightness fluctuations.

2) EXPERIMENT WITH ROBOT BEHAVIOUR CONTROL

In the experiments above, it has been demonstrated that each of the VMPNs can reliably detect specific motion patterns, and the visual cues fusion neurons, including the TMCIS, can integrate and fuse the visual cues and make a final decision. Now the motor commands are associated with each recognised visual motion situation to test whether the integration of VMPNs in a miniature mobile robot can trigger responding actions robustly.
An experiment arena is built for the Colias-IV robot to run freely inside. The arena is about 70x60 cm² surrounded by low barriers which leave the robot’s view unblocked to the open laboratory environment. An obstacle (a billiard ball described in previous experiments) is used as the visual stimulus. The path of the obstacle can be controlled when repeated.

The robot is set free from a fixed position inside the arena. Before it detects an obstacle in front or translating in front, it remains a certain course. Four kinds of visual events are employed to challenge the robot: 1) fixed position dark loom; 2) fixed position bright loom, 3) object translating from left to right and 4) object translating from right to left. Testing scenarios are repeated 50 times each. The obstacle used in the experiment is a black ball as dark loom and a white ball as the bright loom. The motion command rules responding to the visual obstacle is kept simple but clear enough to be recognised, as described in table 4.

The robot’s path with correct responding to the testing event in this experiment are illustrated in figure 17, and the data conclusion of the experiment is illustrated in figure 18. A top-down camera tracks the robot’s path by the Whycon [73] tracking marker carried on top of the robot.

From the observed results, the following analysis can be given, that 1) the proposed ViMDNN model can detect most of the dynamic visual motion events and corresponding trigger reactive motor commands. 2) The average success rate (as indicated in figure 18) of detecting a visual stimulus correctly with our proposed systems is slightly more than 85%. 3) The Distance to Collision (DTC) is nearly constant, which demonstrates the robustness of the ViMDNN model. However, evidence from the previous study proves that DTC would increase as the moving speed gets faster [42], while the TTC decreases that allowing less time to respond to a certain event, according to the results depicted in figure 12c.

V. DISCUSSION

In biology, both LGMD1 and LGMD2 are believed to be involved in triggering collision avoidance behaviours in locusts; however, their exact roles and actual neural connections are still unknown. A bio-plausible computational model could be a useful tool to unravel the mechanisms behind these fascinating animal behaviours. One good source of such speculation stems from the anatomy and behavioural studies of young locusts [21], in which the LGMD2 grows to a higher state of maturity than the LGMD1 in younger locusts, especially for newborn and first instars. Meanwhile, the “startling” behaviour is observed more frequently in
TABLE 4. Actions to be taken by each recognised visual motion.

| Recognised Visual Motion | Action Neuron  | Speed (cm/s) | Turn angle (°) |
|--------------------------|---------------|--------------|----------------|
| safe                     | forward       | 6            | 0              |
| dark loom                | U-turn to left | -3           | 175            |
| bright loom              | U-turn to right| -3           | -175           |
| translating left         | turn left     | 0            | 100            |
| translating right        | turn right    | 0            | -100           |
| unknown hazard           | wait 1 sec    | 0            | 0              |

FIGURE 18. The ratio of actions taken by the robot in responding to different visual stimuli, in which each action indicates a recognised motion pattern described in table 4. Each set of experiment is repeated for 50 times.

younger instars when tested with dark looming stimuli. By assuming that LGMD2 plays the dominant role in detecting and reacting to dark approaching objects during locust’s young period, with slighter and delayed reactions ("startle") as opposed to stronger and immediate reactions ("hiding"), the proposed visual cue fusion neurons provide a compelling explanation of the biological behavioural studies.

Implementing complex visual algorithms into embedded processors is not an easy task. In the proposed model, the scale of neurons is tailored to fit the resources available, while methods are used to ensure real-time specifically to ensure the computation is almost constant and within the duration of a whole frame. If the requirements of compactness increase, there will be further room for optimisation through the use of a sparse matrix to store and access data, expanding loops into linear calculations, using bit-shift calculations instead of multiplying or dividing, and taking advantage of faster accessing RAM space or using Direct Memory Access (DMA) channels etc.

The parallelism of ViMDNN can be achieved at two levels. Firstly, for each of the presynaptic VMPNs, the layers formed by cell arrays are implemented by low-level operations such as multiplication and accumulation. This internal consistency benefits from parallel processing. Secondly, VMPNs employed in the ViMDNN are involved in minimum data exchange during their processing except for the signal input from the shared P layer. Therefore, the order of calculation of different VMPNs does not affect the result which is advantageous for multithreading. Meanwhile, unique functions of different VMPNs, such as the selectivity to dark objects and the direction selectivity, are achieved by parameterised and re-configurable parts of the E-LGMD. This guarantees the ease of accessibility to specific functions. On the platform of Colias-IV, due to the constrain of hardware resources, only four VMPNs have been implemented. When a platform with better support for parallel computing can be utilised, such as Field Programmable Gate Array (FPGA) or even Application Specific Integrated Circuit (ASIC), a ViMDNN with more complex motion selectivity may be conveniently designed.

This work provides a possible solution for low-cost and reliable visual motion perception on micro robot platforms with constrained computational resources. There are two methods to implement the ViMDNN model on other robot platforms. Firstly, the ViMDNN model may be applied via a mainstream microcontroller which has been widely adopted by other robot platforms. Therefore, the model can be easily migrated onto these robots with minor tailoring and modifications. Secondly, thanks to the modular design of the CSU module which contains all the necessary hardware resources for data acquisition and processing, the CSU can be regarded as a “plug and play” module which can be easily attached and integrated within other robot platforms to achieve the desired functions. This is highly advantageous for multi-robot systems such as the bio-inspired swarm robotics research [74].

VI. CONCLUSION

In this paper, we have proposed a bio-inspired visual motion discrimination neural network (ViMDNN) for micro mobile robots to enable them to effectively react to or interact with the visual events in dynamic visual environments. Equipped with the proposed ViMDNN, a micro mobile robot with a tiny camera and an embedded processor can robustly detect different visual motion cues in real-time at a rate of 30 frames per second and respond with different behaviours accordingly. The compact modularised ViMDNN is realised with a bio-plausible fusion method to integrate multiple motion cues based on E-LGMD. The functionalities and the robustness of the proposed system have been demonstrated and verified with systematic experiments. The proposed embedded visual system shows the possibility of introducing rapid and reliable reactive control strategies to micro-robots in spite of their constrained computational resources. In the future, more integration methods for sequential visual cues under constrained computing resources will be investigated.
APPENDIX

A. THE NOTATION OF SIGNALS AND VARIABLES

1) The neuron layers are grid-shaped 2D gray-scale images. Each pixel (neural cell) in most layers is stored as 8-bit unsigned value (0-255), except in the G layer which each cell is 16-bit unsigned value (0-65535). The input image format is YUV and is stored by YUYV format, that every two pixels take 32-bit in total.

2) The processing sequence for each frame is labeled with index \( f \). A neuron layer at frame \( f \) from the processing sequence are labelled in the form of \( P(f) \), \( S(f) \), \( G(f) \) and so on.

3) Individual cell element at coordination \((x, y)\) of a certain layer is described in the form such as \( S(x, y, f) \).

B. THE BOUNDARY ISSUE OF CONVOLUTION OPERATION

Considering the boundary issue, the calculation of convolution for image border ring is omitted out. For example, we have input layer \( I \) with size \( M \times N \) and a kernel \( K \) with square size \((2r + 1) \times (2r + 1)\). Their convolution operation:

\[
C(x, y) = \sum_{i=-r}^{r} \sum_{j=-r}^{r} I(x+i, y+j) \cdot K(i+r, j+r) \]

is defined as:

\[
C(x, y) = \begin{cases} 
0 & \text{for } x = 0, \ldots, r - 1, M - r, \ldots, M - 1, \\
0 & \text{otherwise}
\end{cases} \quad (25)
\]

C. THE PRE-S LAYER CALCULATION

B. The operation of \([x, y]^B\) and \([x, y]^C\) in calculating pre-S layers are defined as follows:

\[
[x, y]^B = \begin{cases} 
\min(x, y) & \text{if } x > 0 \text{ and } y > 0 \\
0 & \text{otherwise}
\end{cases} \quad (27)
\]

\[
[x, y]^C = \begin{cases} 
-\max(x, y) & \text{if } x < 0 \text{ and } y < 0 \\
0 & \text{otherwise}
\end{cases} \quad (28)
\]

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C. Hu et al.: Coping With Multiple Visual Motion Cues Under Extremely Constrained Computation Power

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