

# Towards a New Paradigm for Brain-inspired Computer Vision

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**Abstract:** Brain-inspired computer vision aims to learn from biological systems to develop advanced image processing techniques. However, its progress so far is not impressive. We recognize that a main obstacle comes from that the current paradigm for brain-inspired computer vision has not captured the fundamental nature of biological vision, i.e., the biological vision is targeted for processing spatio-temporal patterns. Recently, a new paradigm for developing brain-inspired computer vision is emerging, which emphasizes on the spatio-temporal nature of visual signals and the brain-inspired models for processing this type of data. In this paper, we review some recent primary works towards this new paradigm, including the development of spike cameras which acquire spiking signals directly from visual scenes, and the development of computational models learned from neural systems that are specialized to process spatio-temporal patterns, including models for object detection, tracking, and recognition. We also discuss about the future directions to improve the paradigm.

**Keywords:** Brain-inspired computer vision, spatio-temporal patterns, object detection, object tracking, object recognition.

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## 1 Introduction

Nowadays, computer vision or machine vision, represented especially by deep convolutional neural networks (DCNNs), has achieved great success in many vision tasks<sup>[1, 2]</sup>. Compared to biological vision, however, computer vision is still lagging far behind in both performances and variety of capabilities<sup>[3–5]</sup>. For instance, DCNNs, which mainly mimic the feedforward and hierarchical structure of the ventral pathway of biological vision, has achieved an extremely high accuracy in image classification<sup>[1, 2, 6]</sup>, but in other tasks, such as video analysis and imaging understanding, they are still far from satisfactory<sup>[7]</sup>. Thus, learning from biological vision, the so-called brain-inspired computer vision, is still a promising and efficient way to speed up the development of computer vision<sup>[4, 8, 9]</sup>.

Although the importance of developing brain-inspired computer vision has been widely recognized<sup>[8, 10]</sup>, up to

now, we have not achieved any really breakthrough in the field that can match the achievement of AlphaGo to GO game<sup>[11]</sup> or AlphaFold to protein prediction<sup>[12]</sup>. So, what is the obstacle in the development? We identify that an important issue that is missed in the current practice of brain-inspired computer vision is the ignorance of a key nature of biological vision, i.e., biological vision is targeted on processing spatio-temporal patterns<sup>[10, 13]</sup>. This is fundamentally different from static images which DCNNs are good at. Shortly speaking, the characteristic of having both spatial and temporal structures is the nature of neural signals in every part of the brain<sup>[14, 15]</sup>. At the beginning stage of acquiring visual information from the external world, the signals received by retina are in the form of continuous optical flow; these signals are converted into spike trains by retinal ganglion cells, which are subsequently transmitted layer by layer to the visual cortex, where the visual input is integrated with spikes from other cortical regions conveying the prior knowledge or memory<sup>[16]</sup>; eventually, the visual information is extracted in the form of continuous neuronal responses (see illustration in Fig. 1). The whole process is very complicated with many fine details remaining unknown, nevertheless, the fact that the visual system computes spatio-

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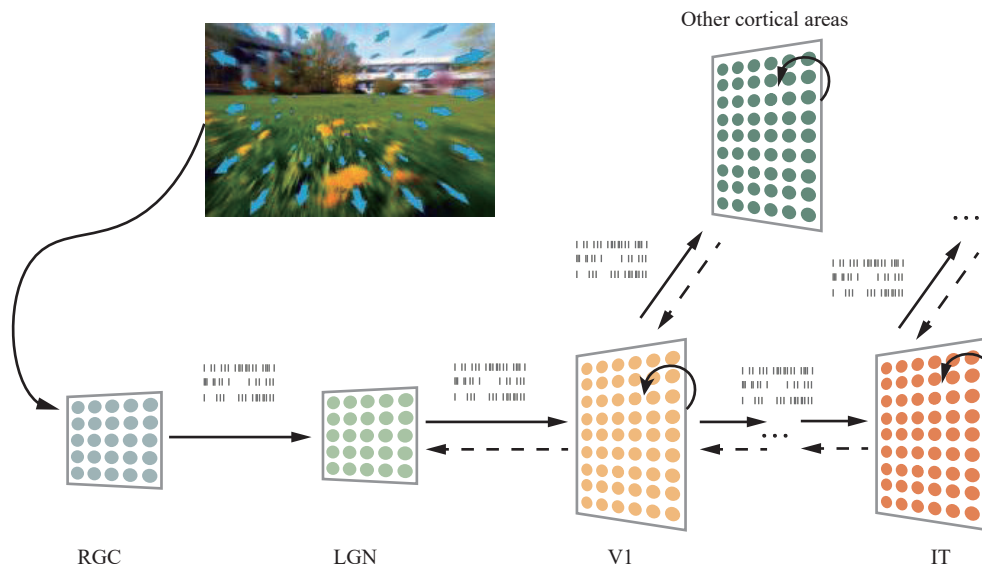


Fig. 1 A schematic of the hierarchical processing of spatio-temporal patterns in the visual system in the form of spike trains. The visual information (in the form of optical flow) of the external world is received by the retina, converted into spike trains by retinal ganglion cells (RGCs), and then processed layer by layer through lateral geniculate nucleus (LGN), V1, V2, V4, IT, etc. There are rich feedback and recurrent interactions within and between visual cortical areas, and interactions with other cortical regions.

temporal patterns in the form of spike trains is fully validated by experiments<sup>[17]</sup>.

In recognition of the aforementioned difference between machine vision and biological vision, a new paradigm which captures this fundamental difference is emerging for developing brain-inspired computer vision. Specifically, in such a paradigm, from beginning the visual information in the external world is expressed in the form of spike trains, which is subsequently processed by computation models inspired by the biological system. Notably, the current popular spiking neural networks (SNNs) in the field do not fulfill this goal<sup>[18]</sup>, and they normally miss many features in biological systems that are crucial for processing spatio-temporal patterns efficiently, such as the differentiation of excitatory and inhibitory neurons, the stochastic neuronal firing<sup>[19]</sup>, the recurrent and feedback interactions between neurons<sup>[15]</sup>, the short-term plasticity of synapses<sup>[20]</sup>, etc.

The organization of the paper is as follows. In Section 2, we will first review the recent develop of spike cameras<sup>[21, 22]</sup>, in particular Vidar<sup>[23]</sup>, which provides a way to represent visual inputs by spike trains. In Section 3, we will review three brain-inspired computational models, which can implement rapid signal detection<sup>[24]</sup>, anticipative object tracking<sup>[25]</sup>, and spatio-temporal pattern recognition<sup>[26]</sup>, respectively based on spike trains. Finally, in Section 4, we will discuss about the future development of brain-inspired computer vision.

## 2 Brain-inspired cameras for sensing visual information

As described above, to develop brain-inspired computer vision, it is critical to represent visual inputs starting from the sensory level to draw analogies with biologic-

al systems. In recent years, brain-like sensing devices have been developed rapidly, which are able to sense visual scenes with high temporal and spatial resolutions, and they convert light signals directly into spike trains. Here, we introduce two of them: one is dynamical vision sensor (DVS)<sup>[27, 28]</sup>, and the other is spike camera<sup>[21, 22]</sup>, e.g., Vidar<sup>[23]</sup>. Both of them were inspired by the retina system of the biological brain.

DVS is a retina-like sensing device. A traditional camera perceives image information frame by frame at a fixed temporal frequency, such as 20fs, and each frame contains the integrated luminance information of the image over the frame interval, while the variance of image luminance between frames is ignored. DVS works in a different way. It asynchronously senses the luminance change at each pixel in the image and outputs a stream of spike events. The output spikes are represented in an address-event manner. Each spike signal contains four elements  $\langle x, y, t, p \rangle$ , which are the horizontal position  $x$ , the vertical position  $y$ , the spiking moment  $t$ , and a variable  $p$  of binary values indicating the direction of luminance change. In other words,  $\langle x, y, t \rangle$  describe the spatio-temporal position of a spike event, and  $p$  describes the way of luminance change. For DVS, a spike is triggered only when the luminance change at the corresponding pixel exceeds a defined threshold, and DVS typically outputs spikes in a spatially sparse and temporally discrete manner. Since DVS only records the relative change of luminance, it is more suitable to sense the motion information in a visual scene, while the detailed texture information of the image is largely ignored. Compared with traditional cameras, DVS has the advantages of broader dynamical range, higher temporal resolution, lower energy

consumption, higher pixel band, etc.<sup>[28]</sup> As a brain-inspired method, DVS has been used recently to simulate a three-layer retina system and implemented a simplified photo receptor-bipolar cell-ganglion cell pathway<sup>[29]</sup>.

If we roughly regard that DVS is mimicking the peripheral region of the retina, then Vidar is mimicking the fovea of the retina<sup>[23]</sup>. Vidar builds a pixel processing array, which contains an analog-to-digital converter (ADC) and an accumulator at each pixel position. ADC converts light intensity into voltage, and this voltage signal is sent to the corresponding accumulator, as shown in Fig. 2. The accumulator integrates the input signal and outputs a spike when the accumulated voltage exceeds a defined threshold, and then resets. The whole process can be regarded as a simplified modeling of photo receptor-bipolar cell-ganglion cell pathway. An ADC simulates a photoreceptor, and an accumulator simulates an integrate-and-fire neuron. Vidar perceives the light intensity information, rather than the change of luminance as done by DVS. The larger the light intensity at a pixel position is, the larger the output of the corresponding ADC generates; consequently, the corresponding accumulator can reach the threshold more rapidly, and generate spikes more frequently. In other words, the light intensity at each pixel is represented by the firing frequency of the corresponding neuron. In Vidar, each accumulator outputs spikes and resets in an asynchronous manner. The spike trains generated by Vidar can then be used to reconstruct the texture features in the visual scene. Vidar is designed to sample data in a temporal resolution as high as millisecond<sup>[21]</sup>. In theory, we can obtain the light intensity information of the image at any time from the recorded spike trains.

### 3 Computational models for brain-inspired computer vision

In the above, we have introduced brain-inspired sens-

ing devices which transform visual inputs directly into spike trains. In this section, we further discuss models for extracting information efficiently from these spike trains. In AI, many image processing algorithms have been proposed, however, these algorithms mainly focus on processing static images, rather than spatio-temporal patterns. Simply transforming artificial neurons in a neural network into spiking neurons does not help much. To develop efficient computational models for processing spatio-temporal patterns, we should learn from biological vision, as the latter is evolved over millions of years to perform this task efficiently<sup>[30]</sup>.

In essence, computer vision needs to perform three fundamental functions: object detection, object tracking, and object recognition. In the below, we will review three type of models inspired by neural systems that are able to perform these three tasks, respectively.

#### 3.1 A brain-inspired model for fast object detection

##### 3.1.1 Biology background

The ability to respond to external stimuli rapidly is critical for animals to survive in natural environments, e.g., to escape from predators. Over millions of years, the biological brain has evolved to process visual information extremely fast. For examples, the response delay of neurons in macaques' visual cortex is only about tens of milliseconds<sup>[31]</sup>, and human brain can complete complex visual scene analysis in around 150ms<sup>[32]</sup>. Computational neuroscience studies have revealed that the capability of fast processing information in the brain is largely attributed to that neural circuits are excitation and inhibition (E-I) balanced, i.e., neurons in a neural circuit on average receive E-I balanced currents. The idea of E-I balanced network was originally proposed by Vreeswijk and Sompolinsky to explain the irregular firings of cortical

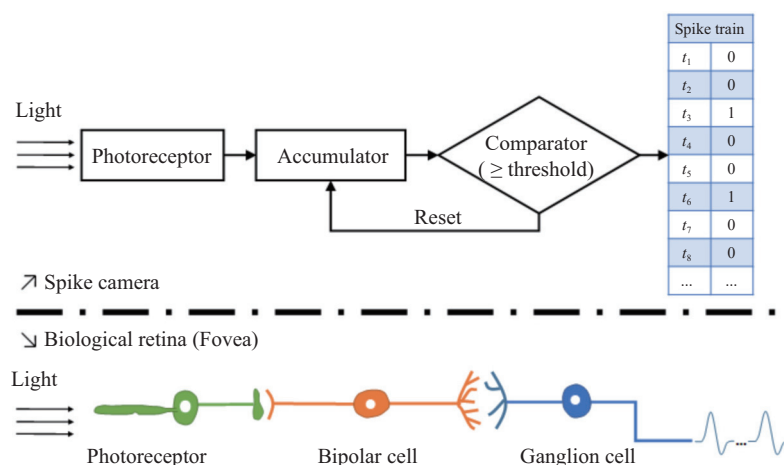


Fig. 2 Workflow of the spike camera Vidar. It contains an analog-to-digital converter (ADC) converting light intensity into voltage, an accumulator integrating inputs, and a comparator generating spike by comparing whether the accumulated voltage exceeds a defined threshold. Because of resetting after spiking, Vidar senses the light intensity change of a visual scene. Vidar can be regarded as modeling the photo receptor-bipolar cell-ganglion cell pathway in the fovea of retina, adapted from [23].

neurons widely observed in the cortex<sup>[33, 34]</sup>. From the computational point of view, they argued that the E-I balanced network has the advantage of responding to external inputs extremely fast, since neuronal activities in such a network are very noisy. The underlying mechanism can be intuitively understood as follows. Without noise, the reaction time of neurons is restricted by the membrane time constant (see illustration in Fig. 3(b)). In an E-I balanced network, however, the system is at a chaotic state with neurons' membrane potentials widely distributed (Fig. 3(b)). As a result, no matter how small the external input change is, there are always a number of neurons responding fast to detect this signal. Thus, at the neuron ensemble level, the network can detect external signals extremely fast. So far, a large amount of experimental data has confirmed that E-I balance is a general property of neural systems<sup>[35-38]</sup>.

**3.1.1.2 The computational model**

Since E-I balanced neural circuits are the substrates of the brain to realize fast computation, it is natural to ask whether such type of model can be used in brain-inspired computer vision. Recently, Tian et al.<sup>[24]</sup> explored this issue and demonstrated the feasibility. In the below, we briefly introduced their work.

Tian et al.<sup>[24]</sup> firstly studied a recurrent network with homogeneous connections, as shown in Fig. 3(a). The network has a large size  $N$ , with  $N_E = q_E N$  and  $N_I = q_I N$

being the numbers of excitatory and inhibitory neurons, respectively. For simplicity of theoretical analysis, they consider no-leaky integrate-and-fire neurons. Each neuron receives recurrent inputs from other neurons and an external input, whose dynamics is written as

$$\tau_a \frac{dv_{a,i}}{dt} = \sum_{b=E,I} \sum_j J_{ij}^{ab} \sum_k \frac{1}{\tau_{b,s}} e^{-(t-t_{j,k})/\tau_{b,s}} + f_a u$$

$$a = E, I$$
(1)

where the subscript  $a$  denotes the neuron population, with  $a = E$  or  $a = I$  representing neurons are excitatory or inhibitory, respectively.  $\tau_a$  is the integration time constant of neurons in population  $a$ .  $v_{a,i}$  is the membrane potential of neuron  $i$  in population  $a$ .  $J_{i,j}^{a,b}$  denotes the connection between neuron  $j$  in population  $b$  and neuron  $i$  in population  $a$ . If a connection exists between them, the connection strength is set to be  $J_{i,j}^{a,b} = j_{ab}/\sqrt{N}$ ; otherwise  $J_{i,j}^{a,b} = 0$ .  $\tau_{b,s}$  is the synaptic time constant of population  $b$ , and  $t_{j,k}$  is the moment of the  $k$ th spike of neuron  $j$ . The probability that neuron  $j$  in population  $b$  connects to neuron  $i$  in population  $a$  is  $p_{a,b}$  for all  $i, j$ .  $u(t)$  denotes the external forward input, i.e., the signal, whose strength is controlled by  $f_a$ . When  $v_{a,i}$  reaches a threshold  $\theta$ , the neuron generates a spike and  $v_{a,i}$  is reset to  $v_0$ .

By using the mean-field approximation, Tian et al.<sup>[24]</sup>

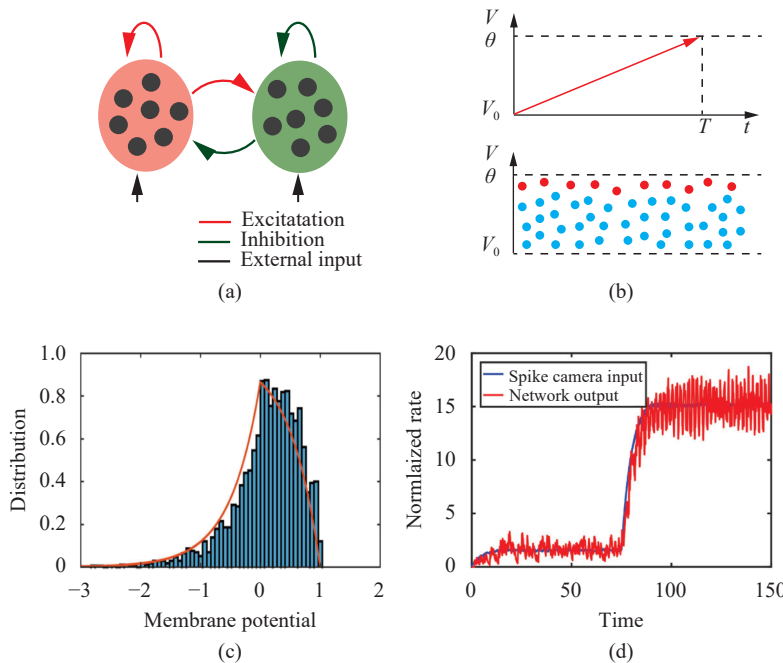


Fig. 3 An E-I balanced neural network for rapid object detection: (a) An example of E-I balanced neural network, in which a neuron receives balanced excitatory and inhibitory recurrent inputs. (b) Illustrating the working mechanism. Upper panel: the reaction time of a non-leaky integrate-and-fire neuron is restricted, which takes time  $T = (\theta - v_0)\tau/I$  to reach the threshold  $\theta$  starting from the resting state  $v_0$ , where  $\tau$  is the membrane time constant and  $I$  the input current. Lower panel: in an E-I balanced network, the membrane potentials of neurons are widely distributed, such that neurons whose potentials are close to the firing threshold (red dots) can always respond rapidly to input changes. (c) Stationary distribution of neuronal membrane potentials in an E-I balanced network when the external input is a constant. The red curve is the theoretical prediction and the blue histogram the simulation result. (d) An E-I balanced network can almost instantly track the change of external inputs generated by a spike camera. Figs. 3(b)–3(d) adapted from [24].

derived the condition for E-I balance in the network, which is  $f_E/f_I > W_{EI}/W_{II} > W_{EE}/W_{IE}$ , where  $W_{a,b} = p_{ab}j_{ab}q_{bb}$ . Under this condition, the network receiving spatially uniform inputs maintains a stable distribution of neurons' membrane potentials (see Fig. 3(c)). By solving the corresponding Fokker-Planck equation, the stable distribution is obtained as

$$p_a(v) = \begin{cases} \frac{1}{\theta} [1 - \exp(-2\tau_a\beta_a)] \exp\left(\frac{2\tau_a v}{\beta_a}\right), & \text{if } v < 0 \\ \frac{1}{\theta} \left[1 - \exp\left(\frac{-2\tau_a(\theta - v)}{\beta_a}\right)\right], & \text{if } 0 \leq v \leq \theta \\ 0, & \text{if } v > \theta \end{cases} \quad (2)$$

where  $\beta_a$  is the variance-to-mean ratio. The derived distribution is confirmed by the simulation (see Fig. 3(c)). This distribution is invariant with respect to the input strength; in other words, the network is always ready to detect the input change.

In practice, however, when the external input is spatially localized, the above condition for E-I balance no longer holds. To solve this problem, Tian et al.<sup>[24]</sup> consider a localized neuronal connections, in term of that each neuron tends to have a higher connection probability to close neighbors than to distal neighbors, with the value decreasing as a Gaussian function of the distance. Tian et al. derived that the E-I balance condition in such a heterogeneous network becomes  $\bar{f}_E/\bar{f}_I > \bar{W}_{EI}/\bar{W}_{II} > \bar{W}_{EE}/\bar{W}_{IE}$ , where the bar denotes spatial average. For more details, see the analysis in [24].

Tian et al.<sup>[24]</sup> applied their model to simulated data of a spike camera and demonstrated that the model can detect the rapid change of the external input (see Fig 3(d)).

### 3.1.3 Future development

Overall, previous studies have demonstrated that the E-I balanced network originated from neuroscience has potential to be applied in brain-inspired computer vision, serving for fast object detection. To fully validate this application, however, there are still a lot of researches to be done. These include, for instances, 1) the reliability of the model in various visual conditions; 2) the robustness of the model to various noise forms; 3) the acceleration of the model to match the high speed of spike cameras; 4) the comparison of the model with other methods; 5) the integration of the model with other modular functions of computer vision; 6) the test of the model in real-world applications.

## 3.2 A brain-inspired model for object tracking

### 3.2.1 Biology background

In navigation tasks, animals need to track their spatial locations and head-directions smoothly<sup>[39]</sup>. A large volume of modelling studies has revealed that the brain

can exploit a type of recurrent neural network called continuous attractor neural networks (CANNs) to achieve this task<sup>[39, 40]</sup>. In the brain, in addition to tracking the movement of an object continuously, there is an extra imposed requirement of tracking the object movement anticipatively, i.e., to predict the future position of the object. This is because in the brain, the transmission of neural signal is significantly delayed. For example, the propagation of visual signal from the retina to the primary visual cortex takes about 50–80ms<sup>[41]</sup>. If this delay is not compensated, our perceived position of the object will lag its true position in the external world considerably, impairing our vision. A strategy to compensate the transmission delay is anticipation, and experimental evidences unveil that the brain does adopt such a strategy. For example, in spatial navigation, the internal head direction encoded by neurons in anterior dorsal thalamic nuclei in a rodent precedes the true head position of the rodent by 25ms<sup>[42]</sup>, i.e., the internal representation of the neural circuit predicts the external input by 25ms. In the below, we will review a computational neural model to realize anticipative object tracking and discuss about its potential applications in brain-inspired computer vision. Non-anticipative tracking will be treated as a special case when the amount of anticipative time is less than zero.

### 3.2.2 Computational model

Motivated by the experimental findings, Mi et al.<sup>[25]</sup> proposed a computational model for anticipative object tracking and applied it to real-world problems<sup>[43, 44]</sup>. Specifically, they consider a CANN with adaptive neuronal responses, which is introduced below. Without loss of generality, a one-dimensional CANN is introduced.

A CANN is canonical network model which has been successfully used to explain the representations of continuous variables in neural system, including motion direction, object orientation, head direction, spatial location, etc.<sup>[45]</sup> Its biological relevance was also supported by recent neuroscience experiments<sup>[40, 46]</sup>. A 1D CANN is illustrated in Fig. 4(a). Denote  $x$  as a 1D continuous stimulus encoded by a CANN, whose value is in the range of  $(-\pi, \pi]$  with a periodic boundary. Denote  $U(x, t)$  as the synaptic input received by neuron at time  $t$  with a preferred stimulus  $x$ , and  $r(x, t)$  as the corresponding neuronal firing rate. The dynamics of  $U(x, t)$  and  $r(x, t)$  are given by

$$\tau \frac{\partial U(x, t)}{\partial t} = -U(x, t) + \rho \int_{x'} J(x, x') r(x', t) dx' - V(x, t) + I_{ext}(x, t) \quad (3)$$

$$r(x, t) = \frac{U(x, t)^2}{1 + \kappa \rho \int_{x'} U(x', t)^2 dx'} \quad (4)$$

where  $\tau$  is the synaptic time constant,  $\rho$  is the neuron



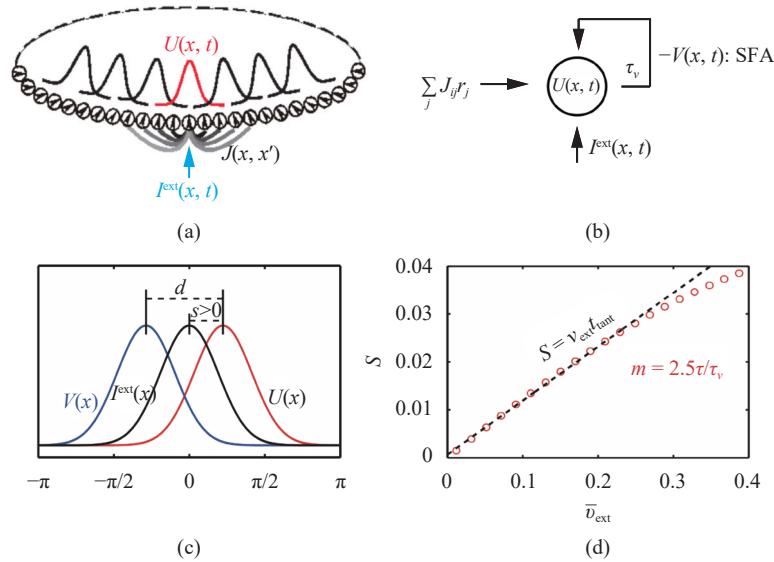


Fig. 4 A CANN with negative feedback for anticipate object tracking: (a) Schematic diagram of 1D CANN encoding the head direction; (b) Diagram of the negative feedback mechanism; (c) An example of anticipative tracking of the moving stimulus (black line). Because of slow spike frequency adaptation (SFA), the bump of  $V(x)$  (blue line) is lagging behind that of  $U(x)$  (red line). (d) Distance  $s$  that the network bump leads the moving object VS. the speed of the moving object  $v_{ext}$ . In anticipative tracking condition, the leading distance of the bump are proportional to the speed of the moving object in a wide speed regime. Figures are adapted from [25].

density, and  $I_{ext}(x, t)$  is the external input.  $J(x, x') = (J_0/\sqrt{2\pi}a)e^{-(x-x')^2/2a^2}$  represents the recurrent connection from neurons  $x'$  to  $x$ , with  $J_0$  controlling the connection strength and the Gaussian width  $a$  controlling the connection range. The relationship between  $r(x, t)$  and  $U(x, t)$  satisfies divisive normalization, with  $\kappa$  controls the inhibition strength.

The current  $-V(x, t)$  denotes the adaptation effect, whose dynamics is written as

$$\tau_v \frac{\partial V(x, t)}{\partial t} = -V(x, t) + mU(x, t) \tag{5}$$

where  $\tau_v$  is the time constant of adaptation and  $m$  controls the amplitude of adaptation.

The adaptation introduces a negative feedback to suppress neuronal activities, as shown in Fig. 4(b). More active a neuron is, stronger the suppression is. Without adaptation, a CANN is known to hold a continuous family of Gaussian-shape stationary states called bumps. Adaptation destabilizes the bump state to induce travelling wave of the bump, i.e., the bump moves spontaneously in the attractor space without relying on external inputs. This reflects the intrinsic mobility of the CANN caused by adaptation, measured by the speed of travelling wave  $v_{int}$ . When the network receives an external moving input, the network tracking behaviour is determined by two competing factors: the intrinsic speed of the network  $v_{int}$  and the speed of the external input  $v_{ext}$ . Interestingly, Mi et al.[25] found that when  $v_{ext} < v_{int}$ , the bump position leads that of the external input, achieving anticipative tracking, as shown in Fig. 4(c). Furthermore, they found that for a wide range of speed values, the leading dis-

tance is proportional to the speed of the external input, implying that the anticipation time is constant, which agrees with experimental data[47], as shown in Fig. 4(d).

The CANN is essentially a computational model employed by the brain to track moving objects. Compared with current machine learning algorithms, it has a number of advantages, including: 1) the computation is performed by the network dynamics, without the need of feature extraction in image frame by frame; 2) the network tracking is robust to noises, a property coming from attractors of a dynamical system; 3) the anticipation time of the model is approximately a constant, independent of the speed of the moving object over a wide range; 4) the model parameters can be defined theoretically according to the task requirement without training by a large dataset; 5) the network computation can be implemented on hardware for neuromorphic computing.

Because of these appealing properties, efforts have been tried to apply CANNs to real-world problems. In [43], a CANN was successfully implemented on the ‘‘Tianjic’’ chip and integrated with other methods in a self-driving bicycle system to track a running object. Recently, a spiking CANN was developed to track moving objects anticipatively[44].

### 3.2.3 Future development

Overall, previous studies have demonstrated that the CANN with adaptation originated from neuroscience has potential to be applied in brain-inspired computer vision, serving for anticipative object tracking. To fully validate this application, however, there are still a lot of researches to be done. These include, for instances, 1) the acceleration of the running speed of a CANN to be compatible with spike camera; 2) the integration of the mod-

el with other feature-based methods in computer vision to improve the tracking accuracy; 3) the implementation of a CANN with adaptive neural responses on hardware.

### 3.3 A brain-inspired model for spatio-temporal pattern recognition

#### 3.3.1 Biology background

A large volume of experimental studies have revealed that the subcortical pathway, which goes from the retina, to superior colliculus (SC), and to higher visual cortex, plays an important role in rapid object recognition<sup>[48, 49]</sup>. For example, the mice study showed that this pathway mediates the rapid innate responses of the animal<sup>[50]</sup>. The capability of the subcortical pathway for rapid motion processing relies on its two main features. Firstly, different from the ventral visual pathway that slowly and hierarchically processes visual information, the subcortical pathway provides a shortcut from the retina to higher visual cortex with no explicit feature extraction, where the retina behaves like a reservoir network. Secondly, SC can linearly read out the retina output and perform fast decision making. The experimental study<sup>[51]</sup> showed that the wide vertical cells in SC, which have large receptive fields and wide-acting inhibition, can realize sampling

over a large retina area and implement winner-take-all computation. The monkey experiment also demonstrated that SC plays a causal role in perceptual decision making<sup>[52]</sup>. In addition to the retina-SC subcortical pathway, the early stages of the auditory pathway also share a similar structure, i.e., a reservoir to decision-making pathway, suggesting that there may serve as a canonical mechanism for fast spatio-temporal pattern recognition.

#### 3.3.2 Computational model

Inspired by the structure and computations of the subcortical pathway, Lin et al.<sup>[26]</sup> recently proposed a computational model for rapid motion patterns recognition. As shown in Fig. 5(a), they built a reservoir module followed by a decision module to mimic the retina-SC pathway, referred to as reservoir decision-making network (RDMN) hereafter. The details of RDMN are introduced below.

In RDMN, a hierarchical reservoir network was employed to simulate the information processing in the retina, which consists of  $L$  feedforwardly connected layers, as shown in Fig. 5(a). Denote  $x_i^l$  as the synaptic input current received by neuron  $i$  in layer  $l$ , for  $i = 1, \dots, N_l$ ;  $l = 1, \dots, L$ , with  $N_l$  as the number of neurons in layer  $l$ . The recurrent dynamics in the reservoir layer  $l$  is given by

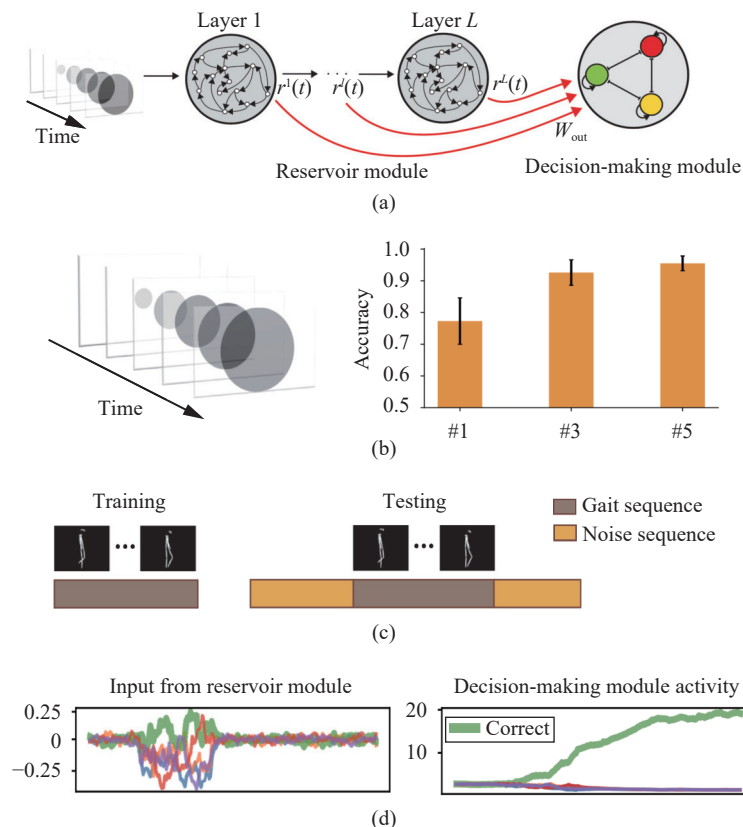


Fig. 5 RDMN performs gait recognition in an event-driven manner. (a) Structure of RDMN, which consists of a reservoir module and a decision-making module. (b) An example of looming pattern (left panel) and the generalization performance in the looming pattern discrimination task (right panel). (c) An example trial of the gait of a subject. (d) Neural dynamics of the decision-making module in the event-based gait recognition task. Figure is adapted from [26].

$$\tau_l \frac{dx_i^l}{dt} = -x_i^l + \sum_{j=1}^{N_{l-1}} M_{i,j}^{l,l-1} r_j^{l-1} + \sum_{j \neq i}^{N_l} M_{i,j}^{l,l} r_j^l + \sum_{j=1}^{N_{in}} M_{i,j}^{l,0} I_j^{ext} \delta_{l,1} \tag{6}$$

where  $r_i = \tanh(x_i)$  is the activation function of neuron  $i$ ,  $M_{i,j}^{l,l-1}$  is the feedforward connections from neuron  $j$  in layer  $l-1$  to neuron  $i$  in layer  $l$ ,  $M_{i,j}^{l,l}$  denotes the recurrent connections from neuron  $j$  to neuron  $i$  in layer  $l$ .  $\tau_l$  is the time constant of layer  $l$ ,  $M_{i,j}^{l,0}$  denotes the input connections.  $I_j^{ext}$  denotes the external input, with  $N_{in}$  is the input dimension.  $\delta_{l,1} = 1$  for  $l = 1$ , and otherwise 0, which indicates that the external input is only sent to layer 1. The connection matrices  $M_{i,j}^{l,0}$ ,  $M_{i,j}^{l,l-1}$ ,  $M_{i,j}^{l,l}$  are all random and sparse, which are sampled from Gaussian distributions and not trained. Notably, the largest eigenvalue of recurrent connections  $M_{i,j}^{l,l}$  in the reservoir module is set to be slightly larger than 1 to enable the the network dynamics to operate on the edge of chaos<sup>[53]</sup>.

After the external visual input is mapped into a special state of the retina network, SC performs temporal information integration and decision making. Thus, a decision-making model is used to model the information processing in SC<sup>[54]</sup>. In RDMN, the decision-making model was greatly simplified and extended to multi-class classification tasks<sup>[26]</sup>. The simplified model is composed of several competing neurons, which receive all the inputs from the neurons in the reservoir module, as shown in Fig. 5(a). And each neuron in the decision-making module represents one category. Denote  $I_i$  as the input summation from all neurons in the reservoir module,  $y_i$  as the total synaptic input of decision neuron  $i$ ,  $r_i$  as the corresponding activation function, and  $s_i$  as the synaptic input due to NMDA receptors. The dynamics of the decision-making network are given as

$$I_i = I_0^* + \sum_{l=1}^L \sum_{j=1}^{N^l} M_{i,j}^{dm,l} r_j^l \tag{7}$$

$$y_i(t) = J_E s_i + \sum_{j \neq i}^{N_{dm}} J_M s_j + I_i \tag{8}$$

$$r_i(t) = \frac{\beta}{\gamma} \ln \left[ 1 + \exp \left( \frac{y_i - \theta}{\alpha} \right) \right] \tag{9}$$

$$\tau_s \frac{ds_i}{dt} = -s_i + \gamma(1 - s_i)r_i \tag{10}$$

where  $M_{i,j}^{dm,l} r_j^l$  is the feedforward connection from neuron  $j$  of layer  $l$  to neuron  $i$  in the decision-making module,  $I_0^*$  is the bias input. The synaptic input  $y_i$  is composed of three parts: 1)  $J_E s_i$  is the self-excitation input, representing the excitatory interactions between neurons encoding the same category; 2)  $\sum_{j \neq i}^{N_{dm}} J_M s_j$  denotes the

mutual inhibition between neurons; 3)  $I_i$  is the feedforward input from the reservoir module. The parameters  $\beta$ ,  $\gamma$  and  $\alpha$  control the shape of the nonlinear activation function. Equation (10) models the the slow dynamics of the synaptic current due to NMDA receptors, with the time constant  $\tau \gg 1$ .  $\tau_s$  controls the time window for integrating input over time by decision-making neurons. The parameters  $I_0^*$ ,  $J_E$ ,  $J_M$  are both chosen optimally based on a thorough mathematical analysis<sup>[26]</sup>.

The only parameters needed to learn in RDMN are the feedforward connection matrix  $M_{l,j}^{dm,i}$  from the reservoir module to the decision-making module, shown by red lines in Fig. 5(a). The loss function is the discrepancy between the actual inputs received by decision-making neurons and the target inputs,  $E = (1/2) \sum_{i=1}^{N_{dm}} \sum_{k=1}^{N_k} \int_0^T dt \times [f_i^k(t) - I_i^k(t)]^2$ , where  $f_i^k(t)$  is the target input.  $M_{i,j}^{dm,l}$  can be optimized by minimizing the loss function using back-propagation through time, or FORCE learning<sup>[55]</sup>, a biologically more plausible method.

When a spatio-temporal input is presented to RDMN, due to the echo properties of the reservoir network, the input is firstly projected from a low-dimensional space to a high-dimensional neural state space, which tends to become linearly separable. Moreover, the hierarchical reservoirs operate at different time and frequency scales, which further enhances the linear separation. With the evidence emerged from the reservoir module, the decision-making module performs a temporal information integration process via its attractor dynamics. Because of the self-excitation and mutual inhibition in the decision-making module, neurons representing different categories integrate information and compete with each other. When a neuron wins the competition, the corresponding classification is made.

Several appealing properties of RDMN are demonstrated through experiments: 1) RDMN successfully reproduces the looming pattern discrimination task as observed in the animal experiment, demonstrating that the model can achieve an excellent generalization performance with only a few trials, see Fig. 5(b); 2) When training data is limited, RDMN outperforms deep learning counterparts, such as LSTM and gated recurrent unit (GRU) on the gait recognition task, and notably, RDMN achieves this by using much fewer numbers of training parameters than long short-term memory (LSTMs) and GRUs. This property is appealing for few-shot learning; 3) Due to the self-excitation and mutual inhibition in the decision-making process, RDMN can perform recognition in an event-based manner, enabling the model to automatically detect and recognize the input pattern, as shown in Figs. 5(c) and 5(d). This property is appealing in real-world applications.

**3.3.3 Future direction**

Overall, previous studies have demonstrated that RDMN originated from mimicking the subcortical path-



way have potential to be applied in brain-inspired computer vision, serving for rapid moving object recognition. To fully validate this application, however, there are still a lot of researches to be done. These include, for instances, 1) the development of temporal structure extraction in RDMN to further enhance its spatio-temporal pattern recognition capability; 2) the deployment of event-based decision module on neuromorphic computing devices for efficient computing on spike signals; 3) the integration of the ventral and subcortical models for integrated local-global and fast-slow visual information processing; 4) the combination of moving object detection, anticipative tracking, and recognition models to build a unified computational framework for spatio-temporal information processing.

## 4 Conclusions and discussions

SNNs were regarded as the “third generation of neural network models” as early as in the 1990s<sup>[56]</sup>. SNNs are biologically more plausible than artificial neural networks, however, up to now, the performances of SNNs are far behind that of artificial neural networks<sup>[18]</sup>. We need to learn more from biological systems to promote the development of brain-inspired computer vision. A key issue that is missed is the fact that biological vision is targeted on processing spatio-temporal patterns. To capture this missing point, recently, a new paradigm for brain-inspired computer vision is emerging, which takes into account the spatio-temporal nature of signals in every part of a vision task, from signal sensing, to object detection, object tracking, object recognition, etc. In recent years, with the rapid development of spike cameras, we are able to collect spatio-temporal spike data from a large and complex scene<sup>[21–23]</sup>; with the rapid advance of computational neuroscience, we are able to build more capable computational models to process spatio-temporal spike data efficiently<sup>[24–26, 57]</sup>; with the fast progress of neuromorphic computing<sup>[58–61]</sup>, we have hope to develop computing platforms that can support the efficient running of brain-inspired neural network models. Combining the above progresses, we are at the edge of entering a new era of practicing brain-inspired computer vision.

In this paper, we have reviewed some recent primary studies on developing spike cameras<sup>[21–23]</sup> and computational models for object detection<sup>[24]</sup>, tracking<sup>[25]</sup>, and recognition<sup>[26]</sup>. This is just the beginning and there are still a lot of researches to be done. In the below, we look ahead to some key issues needed to be solved for the success of the new brain-inspired computer vision paradigm.

1) Developing biologically more plausible brain-like sensing devices. The retina in the brain has rich cell types and wiring structures, which enable the retina to perform smart neural computations<sup>[62]</sup>, such as light adaptation<sup>[63]</sup>, image sharpening, etc. However, recent brain-like sensing devices have only simply converted light patterns

into spike signals, and generated a pixel-level representation of an image. In the future, we need to develop brain-like sensing devices that can mimic more computational features of the retina. For example, DVS records the motion information, which approximately simulates the peripheral information processing of the retina; while the spike camera records the detailed texture information, which approximately simulates the information processing of the fovea. Integrating the advantages of two devices can better simulate the sensory function of the retina of the brain<sup>[21]</sup>.

2) Designing much smarter brain-inspired computational vision models. This paper has only introduced three rather simple computational models for object detection, tracking, and recognition. The biological vision system has much more complicated architectures and richer computational functions, e.g., visual information is processed in parallel through multiple pathways, visual cognition is from global to local, and prior knowledge affects our perception of an image through feedback. We can learn from these characteristics of the biological visual system to develop improved brain-inspired computational models. Also, we can utilize machine learning methods to trained brain-inspired models from data to improve their performances.

3) Exploring more suitable application scenarios for brain-inspired vision models. As the biological vision system is evolved to adapt to natural environments, its computational advantages should be reflected on the efficient and dynamic interactions with the surrounding environments. Thus, for brain-inspired vision, we should also consider suitable application tasks that can fully utilize its advantages, rather than applying it to the tasks, such as static image classification, which deep neural networks are good at. Recently, some interesting researches along this direction have started and they have already demonstrated promising performances, including, for instances, neuromorphic vision-based action detection<sup>[64]</sup>, neuromorphic vision sensor for surveillance (NeuroAED)<sup>[65]</sup>, and event-based neuromorphic vision for autonomous driving<sup>[66]</sup>.

4) Developing more convenient and efficient programming tools for brain-inspired vision models. The fast development of deep learning technologies has benefited from not only large datasets and computing power such as GPU, but also the convenient softwares, such as Pytorch and Tensorflow. These programming tools lower the entry barrier for new comers and boost the whole field. Brain-inspired computational models rely heavily on neural dynamics, event-based computation, and sparse connectivity between neurons. Up to now, simulating and training large-size brain-inspired models are still challenging. Developing efficient and convenient programming tools, similar to Pytorch<sup>[67]</sup> and Tensorflow<sup>[68]</sup> to deep neural networks (DNNs), are urgently needed. Recently, some software platforms towards this goal have been de-

veloped, such as BrainPy<sup>[69]</sup>.

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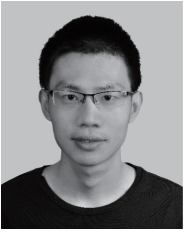
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