ILIF: Temporal Inhibitory Leaky Integrate-and-Fire Neuron for Overactivation in Spiking Neural Networks

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Abstract

The Spiking Neural Network (SNN) has drawn increasing attention for its energy-efficient, eventdriven processing and biological plausibility. To train SNNs via backpropagation, surrogate gradients are used to approximate the non-differentiable spike function, but they only maintain nonzero derivatives within a narrow range of membrane potentials near the firing threshold—referred to as the surrogate gradient support width γ . We identify a major challenge, termed the dilemma of γ : a relatively large γ leads to overactivation, characterized by excessive neuron firing, which in turn increases energy consumption, whereas a small γ causes vanishing gradients and weakens temporal dependencies. To address this, we propose a temporal Inhibitory Leaky Integrate-and-Fire (ILIF) neuron model, inspired by biological inhibitory mechanisms. This model incorporates interconnected inhibitory units for membrane potential and current, effectively mitigating overactivation while preserving gradient propagation. Theoretical analysis demonstrates ILIF's effectiveness in overcoming the γ dilemma, and extensive experiments on multiple datasets show that ILIF improves energy efficiency by reducing firing rates, stabilizes training, and enhances accuracy. The code is available at github.com/kaisun1/ILIF.

1 Introduction

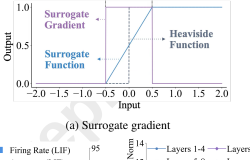
The Spiking Neural Network (SNN), recognized as the third generation of neural networks, is distinguished by the simulation of neuronal message passing through spike-based activations [Maass, 1997]. Unlike the traditional Artificial Neural Network (ANN), which process information with continuous values, SNN, driven by the dynamic accumulation of membrane potential to trigger discrete activation events, effectively mimics the behavior of biological neurons by propagating discrete spike signals (0 or 1) between neurons [Tavanaei et al., 2019]. This spike-based processing endows SNN with remarkable energy efficiency, especially when deployed on neuromorphic hardware platforms such as Intel's Loihi, IBM's TrueNorth, and Tianjic chips [Cai and Li, 2021].

A key challenge in gradient-based optimization for training SNNs is its inherent non-differentiability due to the discrete nature of spike transmissions between neurons. To resolve this issue, surrogate gradient (SG) has been introduced to enable backpropagation in SNNs [Neftci et al., 2019]. However, these approximations can cause or worsen two major issues: overactivation and gradient vanishing. Specifically, overactivation occurs when the accumulated membrane potential surpasses twice the threshold, causing excessive spiking that drives up energy consumption and masks essential temporal information. Strategies such as adaptive thresholding, residual membrane potential modulation, and normalization have been proposed to regulate neuronal activity and mitigate overactivation [Wei et al., 2023; Wang and Yu, 2024; Jiang et al., 2024]. Gradient vanishing, on the other hand, stems from the mismatch between surrogate gradients and discrete spikes, as well as membrane potential decay, resulting in ineffective backpropagation. Current solutions, such as residual learning and adaptive mechanisms [Fang et al., 2021a; Yao et al., 2022], enhance gradient propagation and improve training efficiency.

Existing research often treats overactivation and gradient vanishing as separate issues, overlooking the conflicting effects of the SG on both phenomena. As shown in Figure 1a, given the support width γ in the SG, Figure 1b demonstrates that a larger γ results in elevated firing rates and reduced accuracy (more details will be presented in Section 4.1). However, as [Huang *et al.*, 2024] points out, when the threshold remains constant, a smaller γ risks gradient vanishing. In this paper, this contradiction is referred to as **the dilemma of** γ , which highlights the need to balance the mitigation of overactivation with the preservation of gradient flow. Inspired by the brain's efficient spike regulation through feedforward and feedback inhibition, which controls excessive activation, our objective is to design a module that mitigates the γ -induced conflict between overactivation and gradient vanishing.

This study conducts a theoretical examination of the relationship between γ and both excessive activation and the disappearance of gradients. To address the difficulty of balancing overactivation and gradient vanishing solely by adjusting $\gamma,$ we propose a temporal Inhibitory Leaky Integrate-and-Fire (ILIF) neuron model with two inhibitory units, namely the membrane potential inhibitory unit (MPIU) and the current inhibitory unit (CIU), which are designed to mimic the

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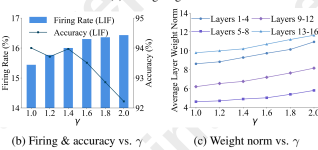


Figure 1: Effect of SG support width (γ) on network performance: (a) Surrogate gradient method. (b) Changes in firing rate and accuracy with γ . (c) Average layer weight norm variation with γ .

inhibitory mechanism of the human brain. The major contributions of this work are as follows:

- Theoretical Analysis: We conduct an in-depth mathematical investigation to clarify how the configuration of γ affects overactivation and its conflict with gradient vanishing, while analyzing how ILIF effectively mitigates both issues.
- **SNN Modeling:** The proposed MPIU and CIU in the ILIF neuron model function as feedforward and feedback inhibition, temporally interacting to reduce excessive activation and mitigate gradient vanishing.
- Experimental Validation: Comprehensive experiments demonstrate that the ILIF model's temporal inhibition mechanism generates fewer spikes while achieving a more stable training process and higher accuracy.

2 Related Work

2.1 Overactivation Control

To mitigate overactivation in SNN, prior works have proposed adaptive thresholds, residual potential modulation, and normalization. Adaptive threshold methods dynamically raise or lower the firing threshold based on recent activity patterns [Ding et al., 2022; Wei et al., 2023; Fang et al., 2020; Zhang et al., 2019]. Residual modulation methods, such as TC-LIF and CLIF, reduce post-spike membrane potential to suppress reactivation, mimicking the effect of afterhyperpolarization (AHP) [Wang and Yu, 2024; Niu and Wei, 2023]. Normalization techniques standardize input distributions across layers and time to stabilize spiking activity [Jiang et al., 2024; Guo et al., 2023; Duan et al., 2022]. However, these methods often rely on handcrafted rules or instantaneous signals, lacking temporal adaptability and biological interpretability. In

contrast, our ILIF model integrates inhibitory units that accumulate both short- and long-term activity, and crucially, adjusts inhibition based on the post-spike membrane potential, providing more precise, causal, and biologically aligned suppression than methods like TC-LIF and CLIF, which depend on pre-spike estimates and risk over-inhibition.

2.2 Improving Gradient Propagation

Due to the intrinsic properties of spiking neurons—namely, spike-based activation and membrane potential decay—temporal gradients often vanish over time. A key factor contributing to this vanishing is the leakage of membrane potential, which diminishes the impact of earlier inputs as time progresses. To mitigate this, models such as PLIF [Fang et al., 2021b] adaptively adjust the leakage rate, preserving critical temporal information. Similarly, gating mechanisms employed in models like GLIF [Yao et al., 2022], STC-LIF [Wang and Yu, 2024], and SpikGRU [Dampfhoffer et al., 2022] enhance temporal dependencies by regulating information flow within neurons. However, these methods typically incur additional computational overhead and are limited to intra-neuron dynamics. In contrast, our approach introduces biologically inspired temporal connections between inhibitory units, serving as shortcuts that simultaneously facilitate backward gradient propagation and forward inhibitory signaling, thereby enhancing temporal dependencies without increasing model parameters.

3 Preliminary

3.1 Vanilla LIF Neuron Model

As shown in Figures 2b and 2c, the LIF serves as a fundamental computational framework, encompassing membrane potential integration, leakage, and spike firing upon exceeding the threshold. The model is described by the following equations:

$$\boldsymbol{U}^{l}[t] = \lambda \boldsymbol{m}^{l}[t-1] + \boldsymbol{I}^{l}[t] \tag{1}$$

$$\boldsymbol{I}^{l}[t] = \boldsymbol{W}^{l} \boldsymbol{S}^{l-1}[t] \tag{2}$$

$$\mathbf{S}^{l}[t] = \mathbb{H}(\mathbf{U}^{l}[t] - V_{th}) = \begin{cases} 1, & \mathbf{U}^{l}[t] \ge V_{th} \\ 0, & \text{Otherwise} \end{cases}$$
(3)

$$\boldsymbol{m}^{l}[t] = \boldsymbol{U}^{l}[t] - \boldsymbol{S}^{l}[t]V_{th} \tag{4}$$

In the LIF model, $U^l[t]$ is the membrane potential in the l-th layer at time step t, combining the postsynaptic current $I^l[t]$ and the residual membrane potential from the previous time step $m^l[t-1]$ with a decay factor λ . The postsynaptic current $I^l[t]$ is computed as the product of the synaptic weight W^l and the input spike $S^{l-1}[t]$. The output spike $S^l[t]$ is generated using the Heaviside function $\mathbb{H}(\cdot)$, which outputs a spike $(S^l[t]=1)$ when the membrane potential $U^l[t]$ exceeds the threshold $V_{\rm th}$ and no spike $(S^l[t]=0)$ otherwise. After firing, the membrane potential $m^l[t]$ is softly reset by subtracting $V_{\rm th}$.

3.2 Spatio-Temporal Backpropagation

Training SNN directly involves implementing Backpropagation Through Time (BPTT) [Werbos, 1990] and using SG to

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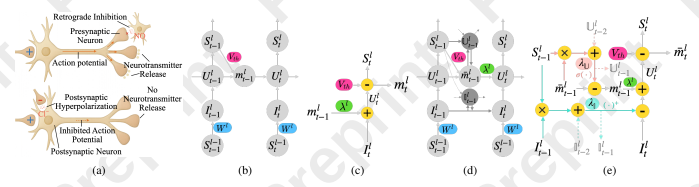


Figure 2: (a) Diagram of the inhibition mechanism. (b) Structure of the vanilla LIF model. (c) Internal operations of the vanilla LIF model. (d) Structure of the ILIF model. (e) Internal operations of the ILIF model.

handle the non-differentiability of spike signals. The gradient of the loss function \mathcal{L} with respect to the weight \mathbf{W}^l at layer l is calculated across all time steps T. It is given by

$$\frac{\partial \mathcal{L}}{\partial \mathbf{W}^{l}} = \sum_{t=1}^{T} \frac{\partial \mathcal{L}}{\partial \mathbf{U}^{l}[t]} \cdot \frac{\partial \mathbf{U}^{l}[t]}{\partial \mathbf{W}^{l}}, l = L, L - 1, \dots, 1, \quad (5)$$

This gradient is computed by decomposing it into spatial and temporal components:

$$\frac{\partial \mathcal{L}}{\partial \boldsymbol{U}^{l}[t]} = \underbrace{\frac{\partial \mathcal{L}}{\partial \boldsymbol{S}^{l}[t]} \cdot \frac{\partial \boldsymbol{S}^{l}[t]}{\partial \boldsymbol{U}^{l}[t]}}_{\text{Spatial Term}} + \underbrace{\frac{\partial \mathcal{L}}{\partial \boldsymbol{U}^{l}[t+1]} \cdot \frac{\partial \boldsymbol{U}^{l}[t+1]}{\partial \boldsymbol{U}^{l}[t]}}_{\text{Temporal Term}},$$

The **Spatial Term** represents the gradient contribution from the current time step, while the **Temporal Term** accounts for the influence of future time steps on the current gradient. It is calculated recursively as follows:

$$\frac{\partial \mathcal{L}}{\partial \boldsymbol{U}^{l}[t]} = \frac{\partial \mathcal{L}}{\partial \boldsymbol{S}^{l}[t]} \frac{\partial \boldsymbol{S}^{l}[t]}{\partial \boldsymbol{U}^{l}[t]} + \sum_{t'=t+1}^{T} \frac{\partial \mathcal{L}}{\partial \boldsymbol{S}^{l}[t']} \frac{\partial \boldsymbol{S}^{l}[t']}{\partial \boldsymbol{U}^{l}[t']} \prod_{t''=t+1}^{t'} \lambda \epsilon^{l}[t''-1] \quad (7)$$

$$= \sum_{t'=t}^{T} \frac{\partial \mathcal{L}}{\partial \boldsymbol{S}^{l}[t']} \frac{\partial \boldsymbol{S}^{l}[t']}{\partial \boldsymbol{U}^{l}[t']} \prod_{t''=t+1}^{t'} \lambda \epsilon^{l}[t''-1]$$

where the term $\epsilon^{l}[t]$ for LIF model is as:

$$\epsilon^{l}[t] \triangleq \frac{\partial U^{l}[t+1]}{\partial U^{l}[t]} + \frac{\partial U^{l}[t+1]}{\partial S^{l}[t]} \frac{\partial S^{l}[t]}{\partial U^{l}[t]}$$
(8)

Detailed derivations are provided in Appendix A. The derivative $\frac{\partial \mathbf{S}^l[t]}{\partial U^l[t]}$ is approximated with the SG $H'(\mathbf{U}^l[t])$, where $H(\cdot)$ provides a smooth approximation of the Heaviside function. A common choice for the SG is the rectangular function, defined as:

$$\frac{\partial \mathbf{S}^{l}[t]}{\partial \mathbf{U}^{l}[t]} = H'(\mathbf{U}^{l}[t]) = \frac{1}{\gamma} \mathbb{1}\left(\left|\mathbf{U}^{l}[t] - V_{\text{th}}\right| < \frac{\gamma}{2}\right), \quad (9)$$

where $\mathbb{1}(\cdot)$ is the indicator function, and γ controls the SG support width. γ is typically set to $V_{\rm th}$ [Meng *et al.*, 2023]. In this case

$$\epsilon^{l}[t] \triangleq 1 - V_{\text{th}}H'(\boldsymbol{U}^{l}[t])$$
 (10)

The gradient $\frac{\partial \mathcal{L}}{\partial S^I[t]}$ in Eq. (7) varies depending on the layers:

$$\frac{\partial \mathcal{L}}{\partial \mathbf{S}^{l}[t]} = \begin{cases} \frac{\partial \mathcal{L}}{\partial \mathbf{S}^{L}[t]}, & \text{if } l = \mathcal{L}, \\ \frac{\partial \mathcal{L}}{\partial \mathbf{U}^{l+1}[t]} \frac{\partial \mathbf{U}^{l+1}[t]}{\partial \mathbf{S}^{l}[t]}, & \text{if } l = \mathcal{L} - 1, \dots, 1. \end{cases}$$
(11)

where
$$\frac{\partial \boldsymbol{U}^{l+1}[t]}{\partial \boldsymbol{S}^{l}[t]} = \boldsymbol{W}^{l+1}$$
.

3.3 Inhibitory Mechanism

In neural systems, various inhibitory mechanisms ensure stable and efficient signal processing. For instance, after a neuron fires an action potential, its membrane potential undergoes afterhyperpolarization. This causes the potential to drop below the resting level, reducing excitability and preventing immediate re-firing. The effect of afterhyperpolarization accumulates over successive spikes, intensifying suppression over time. Additionally, activated downstream neurons send retrograde inhibitory signals to presynaptic neurons by modulating ion channel activity and suppressing neurotransmitter release. This process, mediated by GABAergic interneurons and commonly referred to as retrograde inhibition, suppresses presynaptic currents and limits excessive signal transmission [Bellec et al., 2018; Zenke and Ganguli, 2018]. Inspired by these mechanisms, we enhance the LIF model by incorporating inhibitory processes to reduce overactivation and facilitate more effective gradient flow. The corresponding inhibitory mechanisms are illustrated in Figure 2a.

4 Methodology

4.1 The Dilemma of γ

The dilemma of γ refers to the challenge of balancing overactivation and gradient vanishing in neural networks. The correlation between γ and overactivation has been underexplored, motivating our investigation into this correlation.

Lemma 1. The likelihood of experiencing overactivation is positively correlated with γ .

Proof. When the neural network converges to an optimum such that no further weight updates occur, we consider the system to have reached an equilibrium. Let W_1 be the equilibrium weights when $\gamma = \gamma_1$ and W_2 be the equilibrium weights when $\gamma = \gamma_2$, with $\gamma_2 > \gamma_1$. Suppose that there are

neurons in the firing state (requiring $\mathbf{W}^{l}\mathbf{S}^{l-1}[t] > 0$ to accumulate sufficient membrane potential), whereas others remain inactive (requiring $\mathbf{W}^{l}\mathbf{S}^{l-1}[t] < 0$ to suppress activation).

In the output layer, the gradient terms related to the loss function are: $\frac{\partial \mathcal{L}}{\partial S^L[t]} = \mathbf{Y}^L[t] - \hat{\mathbf{Y}}^L[t], \quad \frac{\partial \mathbf{U}^L[t]}{\partial \mathbf{W}^L} = \mathbf{S}^{L-1}[t],$ where $\mathbf{Y}^L[t]$ represents the average output (within [0,1]), and $\hat{\mathbf{Y}}^L[t]$ represents the predicted values encoded in a one-hot format (taking values of either 0 or 1). As derived from Equations (5), (7), (9) and (10), the expected gradient with respect to W^L is

$$\frac{\partial \mathcal{L}}{\partial W^{L}} = \sum_{t'=t}^{T} \left((\mathbf{\hat{Y}}^{L}[t'] - \hat{\mathbf{\hat{Y}}}[t']) \underbrace{H'(U^{L}[t'])}^{\geq 0} \right) \\
\prod_{t''=t+1}^{t'} \lambda \underbrace{(1 - V_{\text{th}} H'(U^{L}[t'']))}_{\mathbf{\hat{Y}}} \right) \mathbf{\hat{S}}^{L-1}[t] \begin{cases} \leq 0, & \hat{\mathbf{\hat{Y}}}[t'] = 1 \\ \geq 0, & \hat{\mathbf{\hat{Y}}}[t'] = 0 \end{cases} \tag{12}$$

Under the SG framework, the gradient for a given sample becomes zero whenever the membrane potential moves outside the range $[V_{\rm th}-\frac{\gamma}{2},\,V_{\rm th}+\frac{\gamma}{2}]$. Neurons whose membrane potentials exceed or fall below this interval stop receiving weight updates. As λ increases from λ_1 to λ_2 , neurons previously outside the range may re-enter it, allowing further updates during the transition from W_1 to W_2 .

According to Equation (12), Neurons expected to fire $(\hat{Y}[t'] = 1)$ receive negative gradients, which increase positive weights. Conversely, neurons expected to remain inactive $(\hat{Y}[t'] = 0)$ receive positive gradients, which decrease negative weights. This process contributes to an overall increase in the weight norm $\|W\|$. Additionally, based on Equation (11) and the chain rule, changes in the final layer's weights propagate to earlier layers, causing their weights to increase or decrease accordingly.

During activation, the membrane potential increment is $\mathbf{W}^l \mathbf{S}^{l-1}[t]$. A larger $\|\mathbf{W}\|$ amplifies this increment, increasing the chance of exceeding twice the threshold and causing overactivation. As shown in Figure 1c, increasing γ leads to a higher average layer weight norm, which amplifies the membrane potential further, thereby heightening the risk of overactivation.

Lemma 2. The likelihood of experiencing gradient vanishing is inversely correlated with γ .

The gradient in Eq. (7) includes the product $\prod \epsilon^l[t]$, where $\epsilon^l[t] = 1 - V_{\text{th}}/\gamma$ when the membrane potential is near threshold and $\gamma > V_{\text{th}}$. As γ decreases toward V_{th} , this factor becomes smaller, reducing the gradient magnitude over time. At $\gamma = V_{\text{th}}$, $\epsilon^l[t] = 0$, and gradients are completely blocked. See Appendix B for details.

As seen from Lemmas 1 and 2, the impact of γ is twofold: increasing γ exacerbates overactivation, while decreasing it increases the risk of gradient vanishing. This trade-off demonstrates the limitation of adjusting γ alone to improve SNN performance. Thus, specialized modules are needed to strike an optimal balance between these competing effects.

4.2 ILIF Neuron Model

To address overactivation and gradient vanishing in the vanilla LIF model, we propose the ILIF model with two biologically-inspired inhibitory mechanisms. The MPIU provides long-term inhibition by mimicking afterhyperpolarization, stabilizing neuronal excitability. The CIU delivers short-term inhibition via retrograde-like feedback, regulating presynaptic currents. The ILIF structure and internal operations are illustrated in Figures 2d and 2e.

MPIU Each spike contributes to the integration of the post-spike membrane potential $\bar{m}^l[t]$ into $\mathbb{U}^l[t]$, which decays slowly $(\lambda_{\mathbb{U}} \approx 1)$ to maintain a long-term memory of spiking activity:

$$\mathbb{U}^{l}[t] = \lambda_{\mathbb{U}} \left(\mathbb{U}^{l}[t-1] + \mathbf{S}^{l}[t] \cdot \bar{\mathbf{m}}^{l}[t] \right)$$
 (13)

This unit accumulates historical firing patterns, retaining the neuron's firing history and mimicking the afterhyperpolarization phenomenon in biological systems. As firing activity accumulates, the accumulated inhibition $\mathbb{U}^l[t]$ grows, progressively reducing excitability and preventing excessive firing. This inhibition directly affects the membrane potential before subsequent firings:

$$\boldsymbol{m}^{l}[t] = \bar{\boldsymbol{m}}^{l}[t] - \boldsymbol{S}^{l}[t] \cdot \sigma\left(\mathbb{U}^{l}[t]\right)$$
 (14)

where $\sigma(\cdot)$ is a bounded sigmoid function that allows inhibition to increase with historical firing, in contrast to linear mappings or unbounded nonlinear functions that may lead to instability. This mirrors synaptic plasticity mechanisms that adjust synaptic strengths based on past activity, preventing excessive suppression and maintaining learnability over extended timescales. Moreover, the temporal linkage within MPIU mitigates vanishing gradients by offering a direct pathway for backpropagation, as further analyzed in Section 4.3.

CIU Complementing the MPIU's long-term inhibition, the CIU provides rapid, short-term feedback. With a decay coefficient $\lambda_{\mathbb{I}}$ close to 0, primarily inhibiting the incoming current based on the previous time step's current and the current spikes from the next layer:

$$\mathbb{I}^{l}[t] = \lambda_{\mathbb{I}} \left(\mathbb{I}^{l}[t-1] + \boldsymbol{S}^{l}[t] \cdot \boldsymbol{I}^{l}[t] \right)$$
 (15)

The incoming current is adjusted as follows:

$$\mathbf{I}^{l}[t] = \mathbf{S}^{l-1}[t] \cdot W^{l} - \mathbb{I}^{l}[t-1]^{+}$$
 (16)

Here, $\mathbb{I}^l[t-1]^+$ ensures non-negative inhibition, mimicking the rapid feedback mediated by GABAergic interneurons in biological systems. By delivering rapid inhibition tied to the prior spikes and current, the CIU stabilizes excitability in tandem with MPIU.

Combining these mechanisms, the ILIF model evolves according to the following equations, with the corresponding pseudocode provided in Appendix E:

$$\mathbb{I}^{l}[t-1] = \lambda_{\mathbb{I}} \left(\mathbb{I}^{l}[t-2] + \boldsymbol{S}^{l}[t-1] \cdot \boldsymbol{I}^{l}[t-1] \right)
\boldsymbol{I}^{l}[t] = \boldsymbol{S}^{l-1}[t] \cdot W^{l} - \mathbb{I}^{l}[t-1]^{+}
\boldsymbol{U}^{l}[t] = \lambda^{l} \boldsymbol{m}^{l}[t-1] + \boldsymbol{I}^{l}[t]
\boldsymbol{S}^{l}[t] = \mathbb{H} \left(\boldsymbol{U}^{l}[t] - V_{th} \right) = \begin{cases} 1, & \boldsymbol{U}^{l}[t] \geq V_{th} \\ 0, & \text{otherwise} \end{cases}$$

$$\boldsymbol{\bar{m}}^{l}[t] = \boldsymbol{U}^{l}[t] - \boldsymbol{S}^{l}[t]V_{th}
\mathbb{U}^{l}[t] = \lambda_{\mathbb{U}} \left(\mathbb{U}^{l}[t-1] + \boldsymbol{S}^{l}[t] \cdot \boldsymbol{\bar{m}}^{l}[t] \right)
\boldsymbol{m}^{l}[t] = \boldsymbol{\bar{m}}^{l}[t] - \boldsymbol{S}^{l}[t] \cdot \sigma \left(\mathbb{U}^{l}[t] \right)$$

4.3 Analysis of the ILIF Model

The ILIF model builds directly upon the vanilla LIF model by introducing inhibitory decay factors $\lambda_{\mathbb{U}}$ and $\lambda_{\mathbb{I}}$. Setting these factors to zero removes all inhibitory effects, reducing the ILIF model to the standard LIF model. This demonstrates that the ILIF model's inhibitory mechanisms are natural extensions that refine the neuron's behavior without altering its fundamental structure. In this analysis, we explore the relationship between the ILIF and LIF models and the underlying principles that enable the ILIF model to overcome the limitations of the LIF model.

Theorem 1. W' is the equilibrium weight in the vanilla LIF, and W'' is the new equilibrium weight in the ILIF, ||W''|| > ||W'||.

Proof. In the vanilla LIF model, the gradient of the loss \mathcal{L} with respect to $U^l[t]$ consists of both a spatial and a temporal term (denoted as $a(\boldsymbol{W})$, highlighted in blue). In contrast, the ILIF model introduces additional gradient terms (denoted as $b(\boldsymbol{W})$, highlighted in red), as shown in Equation (18). A detailed derivation is provided in Appendix C.

$$\frac{\partial \mathcal{L}}{\partial \boldsymbol{U}^{l}[t]} = \frac{\partial \mathcal{L}}{\partial \boldsymbol{S}^{l}[t]} \frac{\partial \boldsymbol{S}^{l}[t]}{\partial \boldsymbol{U}^{l}[t]} + \sum_{t'=t+1}^{T} \left(\frac{\partial \mathcal{L}}{\partial \boldsymbol{S}^{l}[t']} \frac{\partial \boldsymbol{S}^{l}[t']}{\partial \boldsymbol{U}^{l}[t']} \prod_{t''=t+1}^{t'} \lambda \epsilon^{l}[t''-1] \right) + \sum_{t'=t+1}^{T} \left(\phi^{l}[t'] \prod_{t''=t+1}^{t'} \lambda \epsilon^{l}[t''-1] \right) + \phi^{l}[t] \tag{13}$$

where $\phi^l[t]$ encapsulates the derivatives associated with MPIU and CIU:

$$\phi^{l}[t] = \frac{\partial \mathcal{L}}{\partial \mathbb{U}^{l}[t]} \left(\frac{\partial \mathbb{U}^{l}[t]}{\partial \boldsymbol{\bar{m}}^{l}[t]} \epsilon^{l}[t] + \frac{\partial \mathbb{U}^{l}[t]}{\partial \boldsymbol{S}^{l}[t]} \frac{\partial \boldsymbol{S}^{l}[t]}{\partial \boldsymbol{U}^{l}[t]} \right) + \frac{\partial \mathcal{L}}{\partial \mathbb{I}^{l}[t]} \left(\frac{\partial \mathbb{I}^{l}[t]}{\partial \boldsymbol{S}^{l}[t]} \frac{\partial \boldsymbol{S}^{l}[t]}{\partial \boldsymbol{U}^{l}[t]} \right)$$

$$(19)$$

Consider the vanilla LIF model without inhibition, where the equilibrium weight W' satisfies a(W') = 0. The function a(W) includes terms that are products of W and typically oppose the sign of W. Introducing an additional gradient b(W), which shares the same sign as W (see Appendix

D), modifies the total gradient to $a(\mathbf{W}) + b(\mathbf{W})$, and the new equilibrium \mathbf{W}'' must satisfy $a(\mathbf{W}'') + b(\mathbf{W}'') = 0$.

If $\mathbf{W}'' > 0$, then $b(\mathbf{W}'') > 0$ and thus $a(\mathbf{W}'') < 0$ to maintain equilibrium. Since $a(\mathbf{W})$ becomes more negative as \mathbf{W} increases, this implies $\mathbf{W}'' > \mathbf{W}'$. Similarly, if $\mathbf{W}'' < 0$, then $b(\mathbf{W}'') < 0$ and $a(\mathbf{W}'') > 0$, and because $a(\mathbf{W})$ becomes more positive as \mathbf{W} decreases, it follows that $\mathbf{W}'' < \mathbf{W}'$.

Therefore, in both cases, the magnitude of the new equilibrium increases: $\|W''\| > \|W'\|$. Empirical results supporting this conclusion are provided in Section 5.3.

Theorem 2. r' is the equilibrium firing rate in the vanilla LIF, and r'' is the new equilibrium firing rate in the ILIF. It satisfies that r'' < r' even $\|\mathbf{W}''\| > \|\mathbf{W}'\|$.

Proof. According to Lemma 1, we define the excitatory input function $F(\|\mathbf{W}\|)$, which increases with $\|\mathbf{W}\|$, indicating that larger weight magnitudes result in higher excitatory input necessary for neuron firing, and the response function R(x), which maps input x to firing rate r, reflecting that higher inputs lead to increased activation. Additionally, based on Equations (13) and (15), we define the inhibition function J(r), which increases with the firing rate r and satisfies $J(r) \geq 0$ for all $r \geq 0$, ensuring that higher firing rates induce stronger inhibitory effects.

In the vanilla LIF without inhibition, the equilibrium firing rate is $r' = R(F(\|\mathbf{W}'\|))$. When inhibition is introduced, the new equilibrium rate satisfies $r'' = R(F(\|\mathbf{W}''\|) - J(r''))$. By Theorem 1, we know that $\|\mathbf{W}''\| > \|\mathbf{W}'\|$.

To prove r'' < r', suppose instead that $r'' \ge r'$. Since R is increasing, we have

$$F(\|\mathbf{W}''\|) - J(r'') \ge F(\|\mathbf{W}'\|),$$
 (20)

which implies $F(\|\mathbf{W}''\|) - F(\|\mathbf{W}'\|) \ge J(r'') > 0$. Define the function $H(r) = R(F(\|\mathbf{W}''\|) - J(r))$. Since both R and J are strictly increasing, we have

$$\frac{dH}{dr} = R'(F(\|\mathbf{W}''\|) - J(r))(-J'(r)) < 0,$$
 (21)

so H(r) is strictly decreasing. Note that J(r') = 0, so

$$H(r') = R(F(\|\mathbf{W}''\|)) > R(F(\|\mathbf{W}'\|)) = r'.$$
 (22)

By the fixed-point theorem¹, since H(r') > r' and H is decreasing, the fixed point r'' = H(r'') must satisfy r'' < r', contradicting our assumption.

Therefore, r'' < r'. This shows that even though $\|W''\| > \|W'\|$, the resulting firing rate is strictly lower, meaning that inhibition effectively balances the excitatory input and prevents overactivation.

Theorem 3. *ILIF introduces additional gradient propagation pathways, mitigating the vanishing gradient problem.*

¹Let f(x) be strictly decreasing. If $f(x_1) > x_1$, then any fixed point x^* such that $f(x^*) = x^*$ must satisfy $x^* < x_1$.

Proof. According to Lemma 2, the vanishing gradient problem in LIF is caused by the term $\epsilon^l[t]$, which gradually accumulates to zero with lower γ . In ILIF, however, membrane potential inhibitory units introduce a direct shortcut that connects all time steps. Because each inhibitory unit is interconnected and the decay coefficient is set to 1 in our experiments, there exists a shortcut in the backward pass given by $\frac{\partial \mathbb{U}^t[t]}{\partial S^t[t]} \frac{\partial S^t[t]}{\partial U^t[t]}$ enabling each time step's inhibitory unit to transmit the gradient directly to the preceding time step. Consequently, as inhibition stops after T with the output produced, the complete chain of partial derivatives from time step T-1 back to t can be written as:

$$\frac{\partial \mathcal{L}}{\partial \mathbb{U}^{l}[T-1]} \frac{\partial \mathbb{U}^{l}[T-1]}{\partial \mathbb{U}^{l}[T-2]} \dots \frac{\partial \mathbb{U}^{l}[t+1]}{\partial \mathbb{U}^{l}[t]} \frac{\partial \mathbb{U}^{l}[t]}{\partial \mathbf{S}^{l}[t]} \frac{\partial \mathbf{S}^{l}[t]}{\partial \mathbf{U}^{l}[t]}
= \frac{\partial \mathcal{L}}{\partial \mathbb{U}^{l}[T-1]} \frac{\partial \mathbb{U}^{l}[T-1]}{\partial \mathbf{S}^{l}[t]} \frac{\partial \mathbf{S}^{l}[t]}{\partial \mathbf{U}^{l}[t]}$$
(23)

Since there is no decay in these shortcuts, the gradient can flow from the time step T-1 back to any earlier step without attenuation. This mechanism mitigates the vanishing gradient problem and facilitates effective gradient propagation. \Box

5 Experiments

5.1 Experimental Settings

In this research, we conduct experiments on two types of datasets without data augmentation: standard image classification datasets (CIFAR10 and CIFAR100) and neuromorphic

datasets (DVSCIFAR10 and DVSGesture). For a comprehensive and unbiased evaluation, the proposed ILIF model is implemented under various settings, including different network architectures, neuron models, and training time steps. Further details are provided in Appendix F.

5.2 Evaluation of Accuracy

As shown in Table 1, our ILIF model consistently achieves superior top-1 accuracy compared to existing methods. Specifically, our method achieves 95.49% on CIFAR10 and 78.51% on CIFAR100 using the ResNet-18 architecture with 6 time-steps. For the neuromorphic dataset, ILIF achieves an accuracy of 78.60% on DVSCIFAR10 and matches the top-1 accuracy of 97.92% achieved by other models on DVSGesture, further demonstrating its effectiveness. More experiments are provided in Appendix J.

5.3 Effectiveness of Overactivation Inhibition

We quantified the proportion of neurons activated for more than half the time. As shown in Figure 4, the ILIF model consistently exhibits lower firing rates and fewer instances of continuous activation across all layers compared to LIF model on DVSCIFAR10 and DVSGesture. Furthermore, the reduction in continuous activation rates leads to a lower overall average firing rate, especially with the activation rate by over 30% on DVSGesture. Additional figures for all datasets are provided in the Appendix G.

To better understand the firing patterns across time steps and layers, we compared the spike counts of the LIF and ILIF

Dataset	Method	Network Architecture	Time Step	Accuracy (%)
CIFAR10	STBP-tdBN [Zheng et al., 2021]	ResNet-19	4/6	92.92 / 93.16
	Dspike [Li <i>et al.</i> , 2021]	Modified ResNet-18	4/6	93.66 / 94.05
	GLIF [Yao et al., 2022]	ResNet-18	4	94.67
	SML [Deng et al., 2023]	ResNet-18	6	95.12
	CLIF [Huang <i>et al.</i> , 2024]	ResNet-18	4/6	94.89 / 95.41
	Ours	ResNet-18	4/6	95.24 / 95.49
CIFAR100	STBP-tdBN [Zheng et al., 2021]	ResNet-19	4/6	70.86 / 71.12
	Dspike [Li <i>et al.</i> , 2021]	Modified ResNet-18	4/6	73.35 / 74.24
	GLIF [Yao et al., 2022]	ResNet-18	4/6	76.42 / 77.28
	SML [Deng et al., 2023]	ResNet-18	6	78.00
	CLIF [Huang <i>et al.</i> , 2024]	ResNet-18	4/6	77.00 / 78.36
	Ours	ResNet-18	4/6	77.43 / 78.51
DVSCIFAR10	STBP-tdBN [Zheng et al., 2021]	ResNet-19	10	67.8
	Dspike [Li <i>et al.</i> , 2021]	ResNet-18	10	75.40
	OTTT [Xiao et al., 2022]	VGG-11	10	76.27
	SLTT [Meng et al., 2023]	VGG-11	10	77.17
	Ours	VGG-11	10	78.60
DVSGesture	STBP-tdBN [Zheng et al., 2021]	ResNet-17	40	96.87
	OTTT [Xiao et al., 2022]	VGG-11	20	96.88
	SLTT [Meng et al., 2023]	VGG-11	20	97.92
	CLIF [Huang et al., 2024]	VGG-11	20	97.92
	Ours	VGG-11	20	97.92

Table 1: Comparisons with other SNN neuron models on CIFAR10, CIFAR100, DVSCIFAR10 and DVSGesture

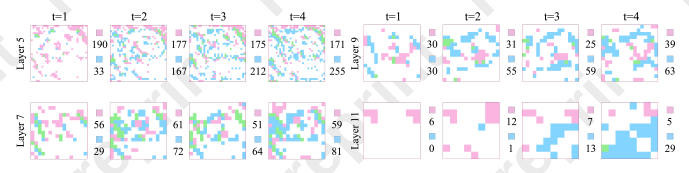


Figure 3: Comparison of firing rates between LIF and ILIF in Layers 5, 7, 9, and 11 on CIFAR10. Blue pixels indicate LIF spikes, red pixels indicate ILIF spikes, and green pixels indicate simultaneous spikes. Spike counts are displayed beside each time step.

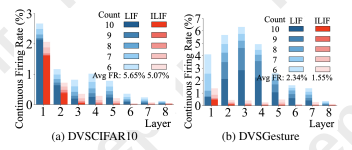


Figure 4: Continuous firing rate comparison on neuromorphic data.

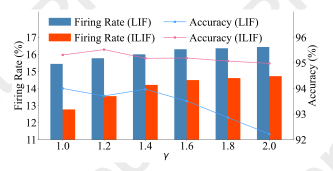


Figure 5: Firing rate and accuracy comparison w.r.t. γ on CIFAR10

models. According to Theorems 1 and 2, inhibitory units lead to larger synaptic weights but also induce strong inhibition. As shown in Figure 3, the ILIF model exhibits a higher firing rate in the first time step due to the absence of immediate inhibition and larger synaptic weights. This efficiently captures critical information early, avoiding prolonged accumulation and delayed activation as observed in the LIF model, where the number of activations rises rapidly across time steps. In later time steps, firing rates decrease as inhibition suppresses redundant activations, ensuring efficient information transfer.

5.4 Evaluation of the Effect of γ on ILIF

Figure 5 illustrates the impact of varying γ on the firing rate and accuracy of the LIF and ILIF models. As γ increases, the LIF model experiences a significant decline in accuracy due to heightened activation, accompanied by a relatively higher firing rate. In contrast, the ILIF model consistently demonstrates a lower firing rate, highlighting its inhibitory effect.

Settings		Accuracy (%)			
MPIU	CIU	CIFAR10	CIFAR100	DVS-C10	DVS-Ges
	V	95.49	78.51	78.60	97.92
	X	95.11	78.06	78.21	97.56
X	\	94.63	76.84	77.13	97.22
X	X	93.76	75.46	75.50	96.88

Table 2: Ablation study of ILIF model components.

Method	CIFAR10	CIFAR100	DVS-C10	DVS-Ges
PLIF	94.25	75.65	75.32	95.49
IPLIF	95.61	78.66	78.41	96.55

Table 3: Impact of adding inhibitory unit to PLIF model.

Furthermore, the ILIF model works synergistically with the backpropagation shortcut provided by its inhibitory units, ensuring more stable accuracy across all γ values.

5.5 Ablation Study

We performed ablation experiments to evaluate the contributions of each inhibitory unit in the ILIF model. As shown in Table 2, removing either unit results in a noticeable drop in accuracy, confirming their necessity. To explore the generalizability of our proposed inhibitory mechanism, we integrated it into the PLIF model [Fang *et al.*, 2021b], a LIF variation that adjusts decay parameters. This integration (Table 3) significantly enhances performance, demonstrating the value of our inhibitory mechanism given that PLIF only modifies decay parameters and inherits the limitations of LIF neurons.

6 Conclusion

This paper analyzes the limitations of the LIF model, specifically overactivation and gradient vanishing caused by the SG support width γ . Inspired by biological inhibitory mechanisms, we propose the ILIF model, which integrates MPIU and CIU to regulate neural activity and enhance gradient propagation simultaneously. Experiments show that ILIF achieves state-of-the-art performance, reduces continuous firing rates, and enhances result stability, providing an efficient and reliable solution for advancing neuromorphic computing.

Ethical Statement

There are no ethical issues.

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