SPIRAL WAVE IN NETWORKS OF HODGKIN-HUXLEY NEURONS: CHANNEL NOISE-INDUCED PHASE TRANSITION

Pratibha Rana*, Jyotsna Singh
Department of computer science, The Northcap University India
Department of computer science, The Northcap University India

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ABSTRACT
The phase transition of spiral waves in networks of Hodgkin-Huxley neurons impelled by channel noise is examined in point of interest. All neurons in the networks are combined with little world associations, and the outcomes are contrasted and the case for normal networks, in which all neurons are totally combined with closest neighbor associations. A statistical variable is characterized to contemplate the aggregate conduct and phase transition of the spiral wave because of the channel noise and topology of the network. The impact of little world connection networks is depicted by neighborhood normal networks and long-go connection with certain probability p. The numerical results affirm that (1) a stable rotating spiral wave can be created and keep up powerful with low p, where the breakup of the spiral wave and turbulence result from expanding the probability p to a specific threshold; (2) suitable intensity of the advanced channel noise can build up a spiral wave among turbulent states in little world association networks of H-H neurons; and (3) regular connection networks are more hearty to channel noise than little world connection networks. A spiral wave in a little world network experiences insecurity all the more effectively as the membrane temperature is expanded to a specific high threshold.

KEYWORD: Access control, AODV, storage node, Optimized Link State Routing, Topology Control, RREP, RERR, HopCount.

INTRODUCTION
Aggregate electrical practices of neurons and oscillators in networks regularly have spatiotemporal examples [1–11]. A spiral wave is one such spatial design and is regularly seen in volatile and oscillatory media. In trial concentrates, the vast majority of the works used to examine the chemical wave in the Belousov-Zhabotinsky reaction [12], and numerous other hypothetical and numerical chips away at spiral waves have been accounted for [13–18]. The significance of examining spiral waves is that it gives vital intimations with reference to how to uproot spiral waves in cardiac tissue and forestall ventricular fibrillation [19] and permits a superior comprehension of the nonlinear flow from a spiral wave to turbulence. There is proof that a spiral wave in cardiac tissue is hurtful, and hence, numerous successful plans have been proposed to take out spiral waves in media. For instance, the plan of periodical compelling is proposed to dispose of the spiral waves and turbulence by creating an objective wave or voyaging wave in the media [20, 21]. Move from a spiral wave to different states actuated by a polarized field [22], twisting of media [23, 24], and the synchronization of spiral waves [25] have additionally been examined widely. Specifically, the clamor actuated arrangement and improvement of spiral waves in a response dissemination framework was examined by Hou and Xin in subtle element [26]. The motion of spiral waves and control pattern determination in response dissemination frameworks have been concentrated broadly while few works have been accounted for on the improvement and phase transition of the spiral wave in the networks of neurons, and its part in signal correspondence inside network is unknown.

A neuronal system comprises of an extensive number of neurons [27] with complex associations. Ordinary electric action of neurons is basic for signal communication among neurons. Stable pivoting spiral waves in rat neocortical slices envisioned by voltage-delicate dye imaging were found in investigations [6, 7]. As noted in [6, 7], spiral waves may serve as pacemakers for the new populace to produce periodic activity in a no oscillatory network.
without the requirement for individual cellular pacemakers. It is fascinating to reenact and explore the arrangement and separation of spiral waves in networks of neurons with various topologies. Perc et al. gained awesome ground in the arrangement of spiral waves in networks [8–10, 28, 29]. He et al. [30] displayed brilliant results on the development of spiral waves in little world systems of FitzHugh-Nagumo neurons and affirmed that the ruinous impact of an inhomogeneous medium on spiral waves can be diminished or uprooted by proper little world associations. A few parts of this theme stay vague; for instance, the dynamics of spiral waves in little world networks of neurons and the impacts of channel noise and the measure of the network on a spiral wave. It is ideal to think about spiral wave dynamics in networks of Hodgkin-Huxley (H-H) neurons than in systems of Hindmarsh-Rose (H-R) neurons [4, 5] on the grounds that the H-R neuron model is a streamlined form of the practical H-H model. Channel noise can change the dynamics of H-H neurons significantly [31, 32]. White et al. [31] brought up that the probabilistic gating of voltage-ward ion channels is a wellspring of electrical ‘channel noise’ in neurons. Schmid et al. [32] reported the capacitance fluctuations lessening channel noise in stochastic H-H frameworks. Fox et al. [33] displayed the autocorrelation functions of channel noise to assess the impact of channel noise. The present work explores the robustness and separation of a spiral wave in little world systems of H-H neurons in the vicinity of channel noise. A statistical variable is characterized to examine the phase transition of spiral waves, and the outcomes are contrasted and those for standard networks. The outer compelling current at all destinations (neurons) is set at zero, which makes every single H-H neuron quiet. It is found that fitting channel noise effectively creates spiral waves and keeps up its power so that signal communication still can go through these calm region.

**MATHEMATICAL MODEL AND DISCUSSION**

The H-H neuron mode is more practical than other displayed neuron models. Little world networks of H-H neurons are portrayed as takes after:

\[
\frac{dm_{ij}}{dt} = am(v_{ij})(1-m_{ij}) - \beta m(V_{ij})m_{ij} + \xi m(t) \tag{2}
\]

\[
\frac{dh_{ij}}{dt} = ah(v_{ij})(1-h_{ij}) - \beta m(V_{ij})h_{ij} + \xi h(t) \tag{3}
\]

\[
\frac{dn_{ij}}{dt} = am(v_{ij})(1-n_{ij}) - \beta m(V_{ij})n_{ij} + \xi n(t) \tag{4}
\]

\[
am = 0.1(V_{ij} + 40)\phi(T)/(1 - \exp(-(V_{ij} + 40)/10)) \tag{5}
\]

\[
\beta m = 4\phi(T)\exp(-(V_{ij} + 65)/18)
\]

\[
\beta h = \phi(T)/(1 - \exp(-(V_{ij} + 35)/10)) \tag{6}
\]

\[
ah = 0.07\phi(T)\exp(-(V_{ij} + 65)/20)
\]

\[
nm = 0.125\phi(T)\exp(-(V_{ij} + 65)/80)
\]

\[
\phi(T) = 3(T - 6.3°C/10°C)
\]

Here, Dm, Dn and Dh depict the intensity of noise, capacity \(\delta(t-t') = 1\) at \(t = t'\) and \(\delta(t-t') = 0\) at \(t \neq t'\), and NNa and NK are the aggregate quantities of sodium and potassium channels present in a given patch of the membrane, individually. On account of homogeneous ion channel density, \(\rho Na = 60 \mu m^{-2}\) and \(\rho K = 18 \mu m^{-2}\), the aggregate channel number is chosen by \(NNa = \rho Na s\) and \(NK = \rho K s\), and \(s\) portrays the membrane patch. Utilizing mean-field hypothesis, a statistical variable [5, 34, and 35] is characterized to examine the aggregate practice.

\[
Cm \frac{dV_{ij}}{dt} = gK n^4(VK - V_{ij}) + gNa m3ij h_{ij}(V_{Na} - V_{ij}) + gL(VL - V_{ij}) + I_{ij} + D \sum_{kl} E_{ijkl}(V_{kl} - V_{ij}) \tag{1}
\]

Here \(R\) is a component of synchronization, \(N2\) is the quantity of neurons and \(V_{ij}\) is the membrane capability of the neuron. It is important to characterize the statistical variable \(R\) to describe the framework's standardized variety and in this way synchronization. \(R\) may not be suitable to portray the synchronization of the example of spiral waves, while it could be valuable in identifying the basic bifurcation parameter inducing separation or disposal of the spiral wave in networks of neurons as beforehand said. As already said [34,35], the curve of the element of
synchronization versus bifurcation parameter delineates the phase transition of the spiral wave through purposes of sudden change. In [34], the creator of the present work reported the additive Gaussian-hued clamor instigated breakup in a regular network of H-R neurons, and What’s more, multiplicative noise in the improvement of a spiral wave in customary systems of neurons (H-R, H-H) has likewise been researched in point of interest [35]. Further numerical results have affirmed that a spiral wave can create in networks (customary or little world sort) of neurons regardless of the fact that there is no external forcing current. The accompanying area displays a numerical examination of the vigor and period of spiral waves in the little world networks of H-H neurons in the vicinity of channel noise where there are no outer constraining currents following up on neurons.

NUMERICAL RESULTS AND DISCUSSION
The numerical examines have a time step h = 0.001, external forcing current I_{ij} = 0, 40000 neurons in a two-dimensional array of 200 × 200 sites, and a no-flux limit condition. The little world connection network can be depicted by nearby consistent networks (finish closest neighbor connections) and a long-extend connection (alternate way) with a certain probability p. In the first place, the instance of no channel noise is considered, and the previews of the membrane possibilities of neurons under various probabilities (p = 0.02, 0.03, 0.04 and 0.05) are plotted with a transient period of around 500 time units. The numerical results introduced in Figure 1 demonstrate that a stable rotating spiral wave can grow totally with proper long-extend probability, and no normal spiral wave is created when the long-run probability surpasses a specific threshold. Note that the patterns in the figure are transient snapshots at t = 500 time units, and the shape and contour of a stable spiral wave frequently stay unaltered a new portions of the spiral wave rise for broken waves. The transient depictions for any altered length of time demonstrate the dissemination of membrane possibilities of neurons, and stable spiral waves are kept up despite the fact that the membrane capability of a neuron at a site in the network changes with time. The comparing component of synchronization R is given in Table 1.

Figure 1 Spatiotemporal patterns developed within a transient period of about 500 time units for long-range probability p = 0.02 (a), 0.03 (b), 0.04 (c) and 0.05 (d). The snapshots are plotted in grayscale from black (about −80 mV) to white (about −40 mV) and the coupling coefficient D = 1.

It is found that a spiral wave can develop and cover more region of the system if a lower long-go association probability is utilized, and a littler component of synchronization is frequently utilized. A littler variable of synchronization additionally demonstrates a shorter transient period required to add to a spiral wave in a system. It is essential to contemplate the impact of channel noise on the phase transition of a spiral wave. Figure 2 shows the connection of the synchronization element and the membrane patch, which depicts the intensity of channel noise, and the previews of membrane possibilities of neurons for various altered membrane patches (intensities of channel noise).

The outcomes in Figure 2(b) affirm that the spiral wave separates when the intensity of channel noise increments to a specific threshold. A spiral wave rises and covers a more noteworthy territory on account of frail channel noise, as seen by looking at the outcomes in Figure 2(b) with those in Figure 1(a).
Table 1  Factors of synchronization under different long-range probability

<table>
<thead>
<tr>
<th>Parameter</th>
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<tbody>
<tr>
<td>$P$</td>
<td>0.02</td>
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![Graph showing calculated factor of synchronization vs. channel noise described by the membrane patch (a)](image)

**Figure 2** Calculated factor of synchronization vs. channel noise, described by the membrane patch (a) and spatiotemporal patterns developed within a transient period of about 2000 time units for $s = 2$ (b1), 3 (b2), 4 (b3), 15 (b4), 17 (b5), 20 (b6), 28 (b7), 30 (b8) and 36 (b9). The snapshots of the development of a spiral wave are plotted for a transient period of about 200 time units for $s = 13$ (c1), 15 (c2), 17 (c3) and 20 (c4). The long-range probability is fixed at $p = 0.02$, the coupling intensity is $D = 1$ and the membrane temperature $T = 6.3^\circ C$. The snapshots are plotted in grayscale from black (about $-80$ mV) to white (about $-40$ mV).

The bend in Figure 2(a) demonstrates that the variable of synchronization abatements with expanding intensity of the channel noise (littler membrane being utilized). There are two unmistakable crests ($s = 15$ and 20) in the curve in Figure 2(a), and the improvement of the spiral wave under channel noise near the two tops is explored by checking the development rate of a spiral wave in the systems. Figure 2(c) affirms that a more drawn out transient period is required for a spiral wave to rise and cover a greater area of a system for the situation that the membrane patch (channel noise) compares to the two crests in the curve. A spiral wave can rise and cover the whole framework with low long-range association probability [36]. It is intriguing to check the dynamic part of channel noise in supporting a spiral wave in a system of neurons. As delineated in Figure 1(c), no general and unmistakable spiral wave involves the systems with long-range association probability $p = 0.04$. Channel commotion is chosen with various intensities to check the impact of channel noise on the arrangement of the spatiotemporal pattern.
Figure 3 Calculated factor of synchronization vs. channel noise described by the membrane patch (a) and a spatiotemporal pattern developed within a transient period of about 2000 time units for $s = 17$ (b1), 23 (b2), 25 (b3) and 28 (b4) at fixed long-range probability $p = 0.04$, coupling intensity $D = 1$, and membrane temperature $T = 6.3\degree C$. The snapshots are plotted in grayscale from black (about $-80$ mV) to white (about $-40$ mV).

Figure 4 Calculated factor of synchronization vs. membrane temperature (a) and spatiotemporal patterns developed within a transient period of about 2000 time units for $T = -4\degree C$ (b1), $0\degree C$ (b2), $2\degree C$ (b3), $8\degree C$ (b4), $10\degree C$ (b5), $12\degree C$ (b6), $16\degree C$ (b7), $18\degree C$ (b8) and $20\degree C$ (b9) at fixed long-range probability $p = 0.02$ and coupling intensity $D = 1$. The snapshots are plotted in grayscale from black (about $-80$ mV) to white (about $-40$ mV).

The outcomes in Figure 3 demonstrate that suitable channel noise can induce and add to a spiral wave in the networks of neurons at a specific long-extend association likelihood despite the fact that the channel noise regularly
affects separation of the spiral wave. Looking at the outcomes introduced in Figure 3(b) with those exhibited in Figure 1(c), it is seen that the spiral wave covers a more noteworthy territory when suitable channel noise is brought into networks of neurons. Obviously, channel noise can optimize the request of the spatiotemporal pattern in a network, and the optimized intensity of channel noise is near the peak of the curve of the variable of synchronization versus membrane patch. As is understood, a high likelihood of long-range connection and noise regularly crush the request of the spatiotemporal pattern and separate the spiral wave. A requested state can be produced when suitable channel noise is brought into media with little world associations. It is the channel noise that improves the request of little world networks, in spite of the fact that it can likewise devastate the request. The membrane temperature regularly has a critical part in deciding the elements of neurons. In this manner, it is intriguing to examine the aggregate practices of spiral waves in networks with little world connections. Figure 4 gives the elements of synchronization at various membrane temperatures and an altered probability of a long-go association. The outcomes in Figure 4 demonstrate that the components of synchronization decline with expanding membrane temperature and separation of the spiral wave is induced in the little world networks of H-H neurons with altered long-go probability $p = 0.02$. It is the little world impact that crushes the requested condition of networks, which varies from the case for consistent networks, in which a specific high membrane temperature just synchronizes all neurons with complete closest neighbor couplings (the media get to be homogeneous at a specific membrane temperature). To make an unmistakable comparison, the components of synchronization for different membrane patches in normal networks are computed and the outcomes are appeared in Figure 5. The outcomes in Figure 5 demonstrate that the element of synchronization changes gradually with the membrane patch size, and there are no sudden changes in the curve of the synchronization component versus channel noise (membrane patch). This shows no stage transition happens as the membrane patch size expands (diminish in the power of the channel noise), and a stable turning spiral wave at last rises to cover the system of neurons. Then again, breakup of the spiral wave is induced by expanding the intensity of channel noise (or diminishing the membrane patch size). These announcements are affirmed by the depictions of membrane possibilities of neurons in the networks. Looking at the outcomes for the normal networks with those for little world networks of neurons, it is found that a consistent network effectively bolsters the spiral wave and keeps up its robustness against channel noise while the little world network regularly instigates the breakup of a spiral wave when the long-extend association likelihood surpasses a specific threshold.
Channel noise can assume a dynamic part in building up a spiral wave in little world networks of H-H neurons just when a proper intensity is chosen. To date, most works have guaranteed that little world connections better depict the mind boggling connections of neurons than consistent networks, in which a neuron is just combined with the four closest adjoining neurons. As far as anyone is concerned, a normal association supporting a spiral wave and long-run connections in little world networks regularly demolishes the spiral wave in homogeneous media. The reason could be that a customary connection results in general variety in the possibilities of the (five adjoining) neurons in the neighborhood area attributable to solid local coupling, and the long-go connection with high probability essentially keeps neurons from evolving all the while.

CONCLUSION
In this work, the channel clamor instigated arrangement and changes in spiral waves in networks of H-H neurons were researched and some intriguing results were found. A statistical variable alluded to as the element of synchronization was characterized to quantify the phase transition of the spiral wave. The little world networks are portrayed by the blend of nearby standard association and long-run connection with certain probability p. Long-go connections with high probability frequently keep the arrangement of a spiral wave, and a created spiral wave can cover a network of neurons just when proper intensity of the channel noise is chosen. The comparing bend of the variable of synchronization versus channel noise (membrane patch) shows coherent resonance-like conduct, odder determination and streamlining with the channel noise. Separation of the spiral wave in a little world network happens more effectively than that in customary networks of neurons as the membrane temperature builds; that is, higher membrane temperature can induce breakup of the spiral wave all the more effortlessly inferable from the impact of little world connections. The component of synchronization changes gradually as the membrane patch size increments (or the power of channel noise diminishes), and the spiral wave keeps up its robustness against certain channel noise. Subsequently, selecting improved channel noise is useful in adding to a stable spiral wave in the little world networks of neurons through measuring and distinguishing the basic element of synchronization versus channel noise bend attributable to its dynamic part in engendering the electrical signal in the quiescent domain.

REFERENCES