Networks of Mobile Elements for Biological Systems

Kanchan Thadani *
Centre for Development of Advanced Computing
Pune University Campus
Pune 411007, Maharashtra, India

Ashutosh
Persistent Systems,
3-B Bhageerath, Senapati Bapat Road
Pune 411008, Maharashtra, India

Abstract

In this paper we present a network model to study the impact of spatial distribution of constituents, coupling between them and diffusive processes in the context of biological situations. The model is in terms of network of mobile elements whose internal dynamics is given by differential equations exhibiting switching and/or oscillatory behaviour. To make the model more consistent with the underlying biological phenomena we incorporate properties like growth and decay into the network. Such a model exhibits a plethora of attributes which are interesting from both the network theory perspective as well as from the point of view of bio-chemistry and biology.

We characterise this network by calculating the usual network measures like network efficiency, entropy growth, vertex degree distribution, geodesic lengths, centrality as well as fractal dimensions and generalised entropy. It is seen that the model can demonstrate the features of both scale free networks as well as small worlds network in different parameter domains. The formation of coherent spatio-temporal patterns is another feature of such networks which makes them appealing for understanding broad qualitative aspects of problems like cell differentiation (e.g. morphogenesis) and synchronization (e.g. quorum sensing mechanisms).

One of the key features of any biological system is its response to external attacks. The response of the network to various attack strategies (isolated and multiple) is also studied.

Key words:
Networks, evolving cellular networks, diffusion, pathways
1 Introduction

Biological systems exhibit tremendous complexity, versatility and robustness in their responses to their environment. Though molecular biology has had spectacular success it is becoming clearer that simple enumeration of genes, proteins and metabolites is not sufficient to understand this complex behaviour. In recent time it has been shown that [Lee et.al,Milo et.al] to capture system level behaviour, it is instructive to search for patterns common to complex systems and networks in general. For example, even if we could calculate the physio-chemical properties of a protein from its structure, to obtain the physiological behaviour from these would be a daunting task without the knowledge about the organisation of the network in which the proteins operate. Networks have been discovered to be important at various levels in biological systems [Hartwell et.al]. These networks of interacting elements are intrinsically dynamic and describe how the system changes in space and time in response to the external stimuli, grows and reproduces, differentiates and dies. Therefore there is a pressing need in biology to develop an alternative language and methodology to deal with such situations.

“Perhaps a proper understanding of the complex regulatory networks making up cellular systems like the cell cycle will require a shift from common sense thinking. We might need to move into a strange more abstract world, more readily analyzable in terms of mathematics than our present imaginings of cells operating as a microcosm of our everyday world” [Nurse P.]

“The best test of our understanding of cells will be to make quantitative predictions about their behaviour and test them. This will require detailed simulations of the biochemical processes taking place within [cells]. We need to develop simplifying, higher-level models and find general principles that will allow us to grasp and manipulate the functions of [biochemical networks].” [Hartwell et.al.]

This modelling of processes in cellular and molecular biology is based upon the assumption that interactions between molecular components can be approximated by a network of biochemical reactions in an ideal macroscopic reactor. Then following the standard methods of chemical reaction kinetics, ordinary

* Email addresses: kanchant@cdac.in (Kanchan Thadani), ashutosh_a@persistent.co.in (Ashutosh).
differential equations are obtained. This modeling approach has been applied to many systems.

However, there are a few considerations which are omitted in above approximation. Firstly, though a few studies consider the spatial aspects of the problem (e.g. [Eldar et. al (2002)] on Drosophila melanogaster), most of the time they are simply neglected. However, it is known that the cell is not a homogenous well stirred reactor, but instead is a highly compartmentalized and heterogeneous environment where phenomena like crowding or channeling are present [Ellis E.(2001)]. In such a situation, the spatial aspects may well play a key role and may need to be incorporated. Such spatial models are particularly relevant for biological networks where the concentration and growth of signalling factors decays with the distance. Thus the probability of establishing a connection with far off nodes is much smaller. Thus the nodes in such networks are more likely to be directly connected to the “nearby nodes” than in the far off nodes. The spatial inhomogeneity observed in real life networks is also indicative of the fact that the values in the network are not uniformly distributed but instead clustered together at various points in space. There may also be bounds and limits in spatial model which forbid or allow connections depending on a cut-off distance.

Another important consideration is that of mobility. In most of the studies the elements are considered to be fixed in space and the structure of coupling is fixed in time i.e. two elements that are initially coupled are coupled forever and they remain at the same point in space. Some models do consider time dependent coupling coefficients [Shibata T. and Kaneko K.] though spatial positions are fixed. However, in real situations the elements move due to diffusion or chemical gradients [Painter K.J and Hillen T.(2002)] and the local interactions determine the connectivity of the element.

There is another way in which such models make restrictive assumptions in context of biological systems. The number of nodes and degrees of freedom are fixed initially and remain fixed. However most of the biological systems show growth, reproduction and death. For example a cell multiplies and transfers its attributes to its offspring and dies. There may be more than one pathway to produce a protein and also more than one way it is degraded. The biological elements also differentiate themselves to allow flexibility with respect to function [Huang S. and Ingber D.E. (2000)]. This differentiation and growth leads to degrees of freedom no longer remaining fixed for a node and diversification of the nature of the elements themselves.

Lastly, most of the models assume either a boolean function for evolution [Shmulevich I. et.al. (2002)] or an arbitrary dynamical rule like dynamical maps(e.g. logistic, circle or Cellular Automata update rules) or some convenient rule like piecewise linear differential equations [de Jong H. et. al.(2003)].
However, such a dynamical rule should have some correspondence (e.g. same kind of non-linearity, continuity properties) to the underlying biological consideration so that it can reproduce behaviour more realistically. Ideally this function should be inferred from experimental data or the detailed knowledge of underlying kinetics. However, as an approximation, at least some motif biological evolution rules which are found in a number of contexts should be used to model the broad qualitative features with a hope that they fall in the same class of systems as the one being modelled and reproduce results with a greater degree of faithfulness to reality.

Thus in light of the above discussion, there is a need to generalise the existing network models of the biological systems. Here we attempt to relax all of the assumptions made above and study the behaviour of such models.

The paper is organised as follows: In the next section we set up the network and elaborate on its features. We also discuss the internal dynamics of the nodes and various choices one can make. The section on analysis studies the mathematical properties of the network and compares them with numerical results. After that we consider various strategies of attack on the network to determine its robustness. Finally we conclude with discussion and interpretation of results.

2 Materials and methods

In this section we set up the network as follows:

2.1 Defining Network Behaviour

A network consists of a set of connected nodes. Each node has an internal state and an intra-dynamics which evolves the state according to a dynamical system. The current state of the node is also dependent on the state of all its neighbours by coupling. In the network that we set up, the number of neighbours a node is coupled to is dependent on the dynamics of the network (inter-dynamics). This is in contrast to the lattice models where the number of neighbours is fixed a-priori. In such lattice models, one either couples to a fixed number of nearest neighbours or has a global coupling to all the other nodes in the network (mean-field models). We feel that in a number of biological contexts all the above assumptions are a little too restricted. During the course of time evolution, a given sub part of the system usually interacts with a variable number of other elements [Shibata T. and Kaneko K.]. Furthermore the network that we set up is an evolving one, where new nodes get added and
redundant nodes get removed. Therefore, the number of neighbours should be
dynamically changing.

Evolving networks have been studied in recent times in great detail
[Erdos et al (1961); Barabasi (2001); Watts (1998); Newman]. Typically the net-
works evolve by adding nodes to a random initial configuration (e.g. Erdos-
Renyi network model). The nodes are usually added by using different heuristic
prescriptions. In particular, scale free networks are obtained by adding
new nodes based on the vertex degree distribution. These networks have been
subject of much study and debate [Albert, R. and Barabasi, A.-L. (2002).] and
model many biological contexts, for example phenomena involving food
webs, biochemical interactions, protein-protein interaction maps for viruses,
prokaryotes and eukaryotes C.elegans etc are described by scale free networks.
A different prescription (e.g. Watt’s Beta model) results in a small worlds net-
work which is found to be useful in modeling co-operative behavior in bi-
ological systems [Erdos et al (1961); Barabasi (2001); Watts (1998); Newman].
However, due to finite size and limited data available in most realistic sit-
uations, it is difficult to establish scale free behavior of a network unambigu-
ously [Albert, R. and Barabasi, A.-L. (2002).]. Secondly, the scale free nature
is often inherent in the prescription to grow a network.

We take a different approach in the present work. Rather than starting from a
random network, we start with a network of few elements and then grow the
network by state-splitting [Lind D. and Marcus B. (1995)]. Such a growth law
does not have a built in prescription for power law behaviour. Thus any scale
free nature found is truly an emergent property rather than a consequence of
a heuristic for network growth.

Diffusion plays a major role in most of the biological processes. Thus any model
which describes a process involving spatial motion or distribution of biological
elements should take into account this important feature. The existing network
models try to incorporate this by varying the connections between the elements
dynamically [Shibata T. and Kaneko K.]. However the assumptions that are
made in these models are too restrictive in many situations. We describe these
assumptions and their possible relaxations below.

(1) **State is mostly considered to be discrete.** For example most com-
monly used network for modelling is a boolean network in which internal
state of the nodes is a discrete (0 or 1) state. This kind of model is,
at best, an approximation for modelling qualitative coarse grained be-
haviour of the system. Some of the limitations of a boolean approach are
the following: firstly since the internal state of an element is evolving
in a discrete manner, it becomes difficult to model situations which in-
volve elements with different times scale being updated asynchronously.
Secondly, we can not have a coupling between elements which is weighted
by the continuous value of the internal state. Such a coupling is desirable particularly in bio-chemical reaction networks. Thirdly, as a dynamical system, the nodes can have more than one attractor i.e., more than one possible final state depending on initial values. To model such situations with boolean networks is cumbersome. On the other hands, using differential equation has advantages of preserving "quantitativity and causal- ity" [De Hoon M. et. al. (2003)] In more realistic models, the nodes have a number of states possible. We assume the state variable describing the internal state to be a continuous variable thus allowing more quantitative approach.

(2) The internal state of the node does not affect the interactions in the networks. In realistic situations this assumption is restrictive. This assumption is actually inherent in the nature of the dynamical systems models where couplings are usually considered as fixed. There are models which consider variable topologies [Erdos et al. (1961); Barabasi (2001); Watts (1998); Newman]. Even in such models the nodes are usually kept fixed and the connections that they make are dynamically changing. However, in most of the situations the elements move (either along a signal gradient or in a diffusive manner) in a very heterogenous medium. Most of the interactions are also dependent on the actual concentrations of active features of these elements. These active features depend on the internal state of the node and change with time. Thus the number of other nodes a given element would interact with depends on the current values of its internal state as well as the current values of internal states of the other elements. Therefore there is a need for introducing an explicit coupling between the internal states and the strength of interaction. We address this issue as follows: The nodes are allowed to move according to either a diffusive motion or a gradient or both. The interaction that takes place depends upon the number of other nodes in its “sphere of influence” and the internal states of the interacting elements have an effect on each other. The mobility of the element is dependent on its internal state and the region it finds itself in. Thus we have an explicit coupling between the number of neighbours that a node has and its internal state.

(3) The interaction range is independent of time. However, for biological systems one can not categorise them as having a fixed interaction range but should have coupling terms which are changing with time. This is because such systems usually have a number of spatio-temporal scales with different elements interacting at different scales. This is modelled in the current system by using a coupling strength that is dependent on the distance between the interacting elements. This dependence can incorporated either by using a threshold value or by using an interaction strength which decays with distance. Thus as an element moves around, its interaction range with other elements is dynamically changing.

(4) The structure of interactions does not change with time i.e. if two
elements are coupled then they have no mechanism to decouple and two
decoupled elements do not couple. We allow for such a mechanism. If an
element is outside the sphere of influence its interaction diminishes and if
it moves within sphere of influence of other element it gets coupled. Thus
the overall network dynamics encompasses many coupling and decoupling
events and overall evolution of network travels through many structures
of interactions.

(5) **Though there is a growth law (fixed by network evolution heuristic) there is usually no decay law.** As mentioned earlier elements
can grow, reproduce and die in many situations of interest. The decay
law, modelling the removal or death of elements in biology plays
an important role in determining behaviour. We incorporate these ef-
fects as follows: the growth is modelled by state splitting of the network
[Lind D. and Marcus B. (1995)]. Such a mechanism is natural for these
contexts. There are two kinds of state splitting that are considered. In-
splitting in which the incoming degree of a node is split between the
node and its offspring and the outgoing degree is shared by both and
the out-splitting in which outgoing degree is split and incoming connec-
tions are shared. This is different from self-organised scale free network
in which nodes get all the degrees inherited. The decay is modelled as
following: once a node is out of the interaction range of all other nodes,
it decouples and is marked as dying. In such a state it would evolve as
an independent dynamical system and reach a attractor of its evolution
state and stay at that state without contributing to the dynamics on the
network. However, since other nodes are also mobile it can happen that
another node with connections to the network comes within the sphere
of influence of this node and get coupled to it. Then the node is “healed”
and starts contributing to the network dynamics again. The nodes which
do not heal within a specified time are considered dead and are removed
from the system. Such a removal does not affect the dynamics of the rest
of the system and is analogous to how biological systems clear dead and
redundant parts.

(6) **The elements are considered identical.** We incorporate diversity and
differentiation of the elements in our system. Firstly, the network is made
up of a number of different types of elements which are representative of
various kinds of dynamics actually observed in biological networks (see
next section).

Thus our network is described by the following: We have a set of coupled nodes.
The number of nodes each element couples to is determined by its state and
its sphere of influence. Every node also has a variable defining its position
in space. This variable depends on the current state of the element, current
position of the element, current position of its neighbours and current state of
the neighbours. Thus the interaction structure of the network is dynamically
dependent on the states, positions and spheres of influence of the nodes.
The network grows by reproduction of its nodes. To model this growth we use state splitting and to model decay we use the following rule: if two nodes are greater than a certain distance apart then they are set to be decoupled. A node which has been decoupled from every node is a “dying” node. However, due to dynamic interactions, such a node may still be revived (“healed”). Any dying node which does not heal for a specified period is considered “dead” and removed from the network.

Thus the network is described by the following set of equations

Calculation of the state of a node.

\[
X_j^{t+1} = X_j^t + \left( \frac{1}{N_{1j}^t} \right) \sum_{i=1}^{N_{1j}^t} (\epsilon_{\text{in}}(i) f_i^j(X_i^t)) - \left( \frac{1}{N_{2j}^t} \right) \sum_{i=1}^{N_{2j}^t} (\epsilon_{\text{out}}(i) f_i^j(X_i^t))
\]  \hspace{1cm} (1)

Calculation of the current position of a node.

\[
r_j^{t+1} = r_j^t + \sum_{i=1}^{N_{1j}^t} \epsilon_{\text{in}}(i) \frac{\sqrt{X_i \cdot X_j}}{|r_i^t - r_j^t|} + \sum_{i=1}^{N_{2j}^t} \epsilon_{\text{out}}(i) \frac{\sqrt{X_i \cdot X_j}}{|r_i^t - r_j^t|}
\] \hspace{1cm} (2)

Calculation of the number of in-neighbours and out-neighbours of a node.

\[
N_{1j}^{t+1} = 2 \sum_{i=1}^{N_{1j}^t} \theta(|r_i^t - r_j^t|) \exp(-c/4)
\] \hspace{1cm} (3)

\[
N_{2j}^{t+1} = 2 \sum_{i=1}^{N_{2j}^t} \theta(|r_i^t - r_j^t|) \exp(-c/4)
\] \hspace{1cm} (4)

where

\[
X_j^t \rightarrow \text{The state vector for a node } j \text{ at time } t
\]

\[
r_j^t \rightarrow \text{The position of a node } j \text{ at time } t
\]

\[
N_{1j}^t \rightarrow \text{The number of in neighbours of a node } j \text{ at time } t
\]

\[
N_{2j}^t \rightarrow \text{The number of out neighbours of a node } j \text{ at time } t
\]

\[
\epsilon_{\text{in}}(i) \rightarrow \text{The in coupling coefficient for node } i
\]

\[
\epsilon_{\text{out}}(i) \rightarrow \text{The out coupling coefficient for node } i
\]
$c(R) \rightarrow$ is a coefficient depending on radius of influence $R$ with value between 0 and 1.

$$\theta(|r_i - r_j|) = \begin{cases} 0 & \text{if } |r_i - r_j| > R \\ 1 & \text{otherwise} \end{cases}$$ (5)

### 2.2 Defining Node Behaviour

Every node has an internal state and an internal dynamics. For the internal dynamics we choose some well known motifs which are frequently seen in dynamics of signalling and regulatory pathways in the cells [Tyson (2003)]. These motifs, which behave as switches and oscillators, serve as building blocks for the bigger, more complex networks. For example, consider a typical situation

The proteins that modulate Cdc2 activity are themselves modulated by Cdc2:Cdc13, through a set of feedback loops

1. Rum1 inhibits Cdc2:Cdc13, but Cdc2:Cdc13 phosphorylates Rum1, thereby targeting Rum1 for degradation.
2. Ste9:APC labels Cdc13 for degradation, but Cdc2:Cdc13 can phosphorylate Ste9, thereby downregulating its activity and targeting it for degradation.
3. Wee1 phosphorylates and inactivates Cdc2:Cdc13, but, at the same time, Cdc2:Cdc13 is trying to phosphorylate and inactivate Wee1.
4. Cdc25 takes the inactivating phosphate group off PCdc2:Cdc13, and Cdc2:Cdc13 returns the favor by phosphorylating and thereby activating Cdc25.
5. Slp1:APC, which also labels Cdc13 for degradation, is itself activated by Cdc2:Cdc13 by an indirect pathway.

The first three feedback loops are examples of mutual antagonism. Under appropriate conditions, the antagonists cannot coexist, i.e. the feedback loop works like a toggle switch. Either Cdc2:Cdc13 has the upper hand and its antagonist (Rum1 or Ste9 or Wee1) is suppressed, or vice versa. The fourth interaction is a positive feedback loop: Cdc2 and Cdc25 activate each other in a mutually amplifying fashion. The last interaction is a time-delayed negative feedback loop, which, under appropriate conditions, can generate oscillations (as Cdc2:Cdc13 concentration rises, it turns on Slp1, which targets Cdc13 for degradation, causing Cdc2:Cdc13 concentration to fall, and Slp1 to turn off).

To study the dynamical consequences of these feedback loops, one must formulate these interactions as a precise molecular mechanism, convert the mechanism into a set of nonlinear ordinary differential equations, and study the
solutions of the differential equations by numerical simulation.

We can set up a network by randomly selecting these elements and connecting them. However, for the present study we choose four elements behaving as switches and two behaving as oscillators.

The equations for the switches and oscillators that we use are as follows

**Perfectly Adapted Switch**

Perfect adaptation means that the steady state response of the element is independent of the signal strength. Such a behaviour is typical of chemotactic systems (including human sense of smell) which initially responds to an abrupt change in interacting attractants and then settles down into a steady response. [Painter K.J and Hillen T.(2002)], [Levchenko A, Iglesias P (2002)] have used a mechanism of this sort to model phosphoinositol signaling in slime mold cells and neutrophils. The equations representing such a system are

\[
\frac{dX_1}{dt} = k_1 S - k_2 X_1 X_2 \\
\frac{dX_2}{dt} = k_3 S - k_4 X_1
\]

\[k_1 = k_2 = 2, k_3 = k_4 = 1, S = 1.2\]

In some situations, like Frog oocyte maturation in response to progesterone and Apoptosis the switching is one way. At a critical signal strength we get an irreversible transition to large response starting from small responses. The transition can be activating or inhibitory giving rise to mutual activation or inhibition

**Mutual Activation Switch**

The equations representing such a system are

\[
\frac{dX_1}{dt} = k_0 E(X_1) - k_2 E(X_1)X_1
\]

\[E(X_1) = G BK(k_3 X_1, k_4, J_3, J_4)\]

\[k_0 = 0.4, k_1 = 0.1, k_2 = k_3 = 1, k_4 = 0.2,\]

\[J_3 = J_4 = 0.05, S = 9.0\]

The Goldbeter- Koshland function [Goldbeter A, Koshland DE(1981)] is graded and reversible. By graded we mean that the response increases continuously with signal strength. A slightly stronger signal gives a slightly stronger response. Reversible means that if we go from an initial value to a final value.
Although continuous and reversible, a sigmoidal response is abrupt. Tyson [Tyson (2003)] compares this to buzzer or a laser pointer, to activate the response one must push hard enough on the button, and to sustain the response one must keep pushing. When one lets up on the button, the response switches off at precisely the same signal strength at which it switched on.

However, even in the presence of such a reversible term, the above switch shows irreversible behaviour due to a feedback loop that is present.

**Mutual Inhibition Switch**

In this switch, if signal is decreased enough, the switch will go back to the off-state. For intermediate stimulus strengths the response of the system can be either small or large, depending on how S was changed. This switch shows hysteresis and is found in the lac operon in bacteria [Laurent M and Kellershohn N (1999)], the activation of M-phase-promoting factor (MPF) in frog egg extracts [Novak B and Tyson J(1993)], the autocatalytic conversion of normal prion protein to its pathogenic form [Kellershohn N and Laurent M (2001)] and budding yeast cell cycle. This behaviour has been observed in a number of experiments. The equations representing this system are

\[
\frac{dX_1}{dt} = k_0 + k_1 S - k_2 X_1 - k_2' E(X_1) X_1 \\
E(X_1) = GBK(k_3, k_4 X_1, J_3, J_4) \\
k_0 = 0.0, k_1 = 0.5, k_2 = 0.1, k_3 = 1, k_4 = 0.2, \\
J_3 = J_4 = 0.05, k_2' = 0.5, S = 10.0
\]

**Negative Feedback Switch** This type of regulation, commonly employed in biosynthetic pathways, is called homeostasis.

The equations representing this system are

\[
\frac{dX_1}{dt} = k_0 E(X_1) - k_2 S X_1 \\
E(X_1) = GBK(k_3, k_4 X_1, J_3, J_4) \\
k_0 = 1.0, k_2 = 1.0, k_3 = 0.5, k_4 = 1.0, \\
J_3 = J_4 = 0.01, S = 0.1
\]

**Activator Inhibitor Oscillator**

The classic example of an activator-inhibitor system is cyclic AMP production in the slime mold, Dictyostelium discoideum [Martiel J and Goldbeter A(1987)].
External cAMP binds to a surface receptor, which stimulates adenylate cyclase to produce and excrete more cAMP. At the same time, cAMP-binding pushes the receptor into an inactive form. After cAMP falls off, the inactive form slowly recovers its ability to bind cAMP and stimulate adenylate cyclase again. This mechanism lies behind all the curious properties of the cAMP signaling system in Dictyostelium: oscillations, relay, adaptation, and wave propagation.

The equations representing this system are

\[
\frac{dX_1}{dt} = k_0 E(X_1) + k_1 S - k_2 X_1 - k_2' X_1 X_2 \quad (10)
\]

\[
\frac{dX_2}{dt} = k_5 X_1 - k_6 X_2
\]

\[
E(X_1) = GBK(k_3 X_1, k_4, J_3, J_4)
\]

\[
k_0 = 4.0, k_1 = k_2 = k_2' = k_3 = k_4 = 1.0, k_5 = 0.1, k_6 = 0.075,
\]

\[
J_3 = J_4 = 0.3, S = 0.2
\]

Substrate Depletion Oscillator

This describes MPF oscillations in frog egg extracts [Novak B and Tyson J (1993)]. MPF is a dimer of a kinase sub-unit, cyclin-dependent kinase 1 (Cdk1), and a regulatory subunit, cyclin B. As cyclin B accumulates in the extract, it combines rapidly with Cdk1 (in excess). The dimer is immediately inactivated by phosphorylation of the kinase subunit X can be converted into active MPF by a phosphatase called Cdc25. Active MPF activates Cdc25 by phosphorylating it.

The equations representing this system are

\[
\frac{dX_1}{dt} = k_1 S - (k_0' + k_0 E(X_2)) X_1 \quad (11)
\]

\[
\frac{dX_2}{dt} = (k_0' + k_0 E(X_2)) X_1 - k_2 X_1
\]

\[
E(X_1) = GBK(k_3 X_1, k_4, J_3, J_4)
\]

\[
k_0' = 0.01, k_0 = 0.4; k_1 = k_2 = k_3 = 1.0
\]

\[
k_4 = 0.3, J_3 = J_4 = 0.05, S = 0.3
\]

where the \( GBK(u, v, J, K) \) is the Goldbeter-Koshland term

\[
GBK(u, v, J, K) = \frac{2uk}{v - u + vJ + uK + \sqrt{v - u + vJ + uK^2 - 4(v - u)uK}} (12)
\]
Thus we can model a number of situations by coupling together these motifs. In the current work, we choose a pair of any of these elements to represent the types of nodes in our network. The parameters specified are fixed at these values following [Tyson (2003)]. The nodes are evolved by integrating the equations using a fourth order Runge-Kutta routine for 100 iterates with time step 0.1 for the switches and 0.001 for oscillators.

2.3 Evolving the Network

We follow the algorithm outlined in flow chart of Fig. 1. The algorithm evolves each node for a fixed time, updates the number of its neighbours Eq. (3), Eq. (4), update the current state by coupling it to the neighbours by Eq. (1), obtain the current position of the node by Eq. (2) and finally update the network by splitting the in and out degree. This constitutes one iteration of the network dynamics. We set the simulation as follows: we choose two of the above six types of nodes and set them up according to an initial golden mean network. The coupling coefficient $\epsilon$ for this particular simulation is considered same for every node, though in principle the equations allow us to have a different value of $\epsilon$ for every node. The offspring of each node is of the same as parent. We choose all possible pairs of the node to do the simulation.

3 Characterisation and Analysis

To understand the behaviour of a network a number of characterizing quantities have been introduced in recent times [Erdos et al(1961); Barabasi(2001); Watts(1998); Newman]. It is important to choose the set of characterisers which reflect the essential features of the network with respect to the problems of interest. We classify the characterisers as the following: Topological characterisers which contain the information about the underlying topological structure of the connections in the network, Geometric characterisers, the growth or Entropic characterisers and characterisers for the dynamics. These characterisers are listed in Table 1 for three representative sets of parameters.

3.1 Topological characterisers

Degree Distributions

The degree is an important characteristic of a node. Based on the degree
of the nodes, it is possible to construct measurements for the network. One of the simplest is the maximum degree which is simply the maximum connections that any node in the network has. The degree distribution, $P(k)$ which expresses the fraction of nodes in a network with degree $k$ contains more information. One may be interested in finding out if there is a correlation between the degrees of different nodes. Such correlations were found to have an important role in many network structural and dynamical properties [Erdos et al(1961); Barabasi(2001); Watts(1998); Newman]. The most common choice is to find correlations between two nodes connected by a link. This correlation can be expressed by the joint degree distribution $P(k, k')$, i.e., as the probability of a link connecting two nodes of degree $k$ and $k'$. Another way to express this is by giving the conditional probability that an arbitrary neighbor of a node of degree $k$ has degree $k'$

$$P(k|k') = \frac{P(k, k')}{P(k)} \quad (13)$$

This distribution gives a very detailed description of node degree correlations but, for fat tailed degree distributions as in scale-free networks, it is difficult to evaluate experimentally, because of the poor statistics. A measure with better statistics is to computing the mean degree of the neighbors of nodes with a given degree, given by

$$k_{nn}(k) = \sum_{k'} k' P(k'|k) \quad (14)$$

In the simulation we use Eq. (14) for calculating vertex degree distribution. The vertex degree distribution can be calculated for the self degrees and neighbour degrees separately. For a random or small worlds network, the vertex degree distribution is a Poisson distribution in a large $N$ limit with a peak at a particular degree being most prominent. In contrast, for the scale free network of Barabasi and Albert, the degree distribution follows a power law for large $k$. In the network model that we consider we can get both types of degree distribution depending on parameter regimes. This is demonstrated in Figs 2(a) and 2(b). the Fig 2(a) shows a typical small worlds behaviour whereas 2(b) shows a power law like decay in large $k$ region. Thus we can model both kinds of phenomena with the same model. As one changes the coupling, we get a transition from small worlds phase to a scale free phase. Further investigations of this transition are in progress.

**Clustering Coefficient**

A characteristic of the Erdos-Renyi model is that the local structure of the network near a node is a tree. More precisely, the probability of loops involving
a small number of nodes tends to 0 in the large network size limit. This is in marked contrast with the profusion of short loops which shows up in many real-world networks. One way to characterise the presence of such loops is through the clustering coefficient. Two different clustering coefficients are frequently used. we can define a coefficient

\[ C = \frac{3N_t}{N_3} \]
\[ N_t = \sum_{i,j,k} a_{ij} a_{jk} a_{ik} \]
\[ N_3 = \sum_{i,j,k} a_{ij} a_{jk} \]

(15)

This coefficient gives the ratio of number of triangles i.e. the triples which are linked to each other to number of connected triples (i.e. three nodes connected to each other directly or indirectly). If we denote by \( l_i \) the number of links between neighbours of node i and \( k_i \) is the number of neighbours of node i then we can also write clustering coefficient as

\[ C_i = \frac{2l_i}{k_i(k_i - 1)} \]
\[ C = \frac{1}{N} \sum_i C_i \]

(16)

The difference between the two definitions is that Eq. (15) gives the same weight to each triangle in the network, while Eq. (16) gives the same weight to each node, resulting in different values because nodes of higher degree are possibly involved in a larger number of triangles than nodes of smaller degree. In the simulation result we calculate the coefficient given by Eq. (16).

**Centrality**

Greater the number of paths in which a node or link takes part, the greater the importance of this node or link for the network. Assuming that the interactions follow the shortest paths between two nodes, it is possible to quantify the importance of a node in this sense by its betweenness centrality

\[ B_i = \sum_{jk} \frac{\sigma(j, i, k)}{\sigma(j, k)} \]

(17)

where \( \sigma(j, i, k) \) is the number of shortest paths between nodes j and k which
pass through node $i$ and $\sigma(j, k)$ is total number of paths between nodes $j$ and $k$. The sum is over all distinct pairs $(j,k)$. Other centrality measures are given by \textit{closeness centrality} $D$, \textit{network centrality} $N$ and \textit{stress centrality} $S$

\begin{align*}
D_i &= \sum_j \frac{1}{d_{ij}} \\
N &= (\max_j d_{ij})^{-1} \\
S &= \sum_{jk} \sigma(j, i, k)
\end{align*}

where $d_{ij} =$ shortest distance between $i,j$

\subsection*{3.2 Geometric characterisers}

\textbf{Geodesic Lengths}

In the general case, two nodes of a complex network are not adjacent. In fact, most of the networks of interest are sparse, in the sense that only a small fraction of all possible links are present. Nevertheless, two non-adjacent nodes $i$ and $j$ can be connected through a sequence of $m$ links $(i, k_1), (k_1, k_2), \ldots, (k_{m-1}, j)$; such set of links is called a path between $i$ and $j$, and $m$ is the length of the path. We say that two nodes are connected if there is at least one path connecting them.

A geodesic path (or shortest path) between nodes $i$ and $j$, is one of the paths connecting these nodes with minimum length. The length of the geodesic paths is the geodesic distance $d_{ij}$ between nodes $i$ and $j$. There can be more than one geodesic paths between two nodes. We take the mean geodesic distance as a characteriser of the network. The definition has a problem if the network contains unconnected nodes therefore this measurement is only performed after cleanup operation. The Dijkstra algorithm is used for calculating the shortest path on the network.

\textbf{Network Efficiency}

Network efficiency measure the ease of communication on a network. This can be defined as

\begin{equation}
\xi = \frac{1}{N(N-1)} \sum_{i \neq j} d_{ij}^{-1}
\end{equation}

This quantity is also useful in characterizing robustness of network to attack
3.3 Entropic characterisers

Growth rate and Entropy

The entropy of a network can be calculated by using the following result based on Frobenius - Perron theory which we state without proof. For proof see [Lind D. and Marcus B. (1995)]

Theorem:

If $G$ is an irreducible graph then $h(X_G) = \log \lambda_{A(G)}$ where $h(X_G)$ is the entropy of the shift dynamical system $X_G$ associated with the graph $\lambda_{A(G)}$ is the Perron eigenvalue of the graph (i.e. in this case the largest real eigenvalue of the adjacency matrix).

In general, however the above result provides an upper bound to entropy and we calculate this by diagonalising the adjacency matrix and finding the largest eigen value. Since our network is an evolving hypergraph, we compute this quantity at every time step and plot it with time in Fig 3. We find that the entropic bound saturates to certain fixed value (see Table 1) for a set of parameters and grows without bound for another set of parameters. This means that the network attains certain optimal size for a set of parameter values and has unrestricted growth for another set. Continuously varying the parameters we have a transition from restrictive to unrestricted growth.

Generalised Entropy

Our network starts with nodes having a degree $\leq 2$, the degrees are added by splitting and by merger and removed by nodes dying. In the parameter regime where we get scale free behaviour the degree distributions can be fitted by

$$P(\geq k) = e^{(k-2)/K}$$

$$e^{q_c} = [1 + (1 - q_c)x]^{\frac{1}{1-q_c}}$$

for

$$1 + (1 - q_c)x > 0$$

(20)

where $K$ is related to the value $q_c$ which best fits the data. This quantity is determined as follows: we take the logarithm of the probability for a series of different values of $q_c$ ranging from -3 to 3 in steps of 0.1. The values obtained from the $q$-exponential form are then fitted to data obtained from the model
by a linear fit. The value of \( q \) which gives the best fit is determined as value \( q_c \).

This form is taken from the formulation of a non-extensive generalised entropic function introduced by Tsallis [Gell-Mann M. and Tsallis C. (2004)]

The entropic function

\[
S_q \equiv \frac{1 - \int_0^\infty dk[p(k)]^q}{1 - q}
\]  

(21)

extremising this as described in [Gell-Mann M. and Tsallis C. (2004)] gives

\[
P(\geq K) = [1 - (1 - q)\beta(k - 2)]^{\frac{(2-q)}{(1-q)}}
\]  

(22)

this equation is identical to Eq.(20) if we assume \( q_c = 1/(2 - q) \) and \( K = 1/(2 - q)\beta \). Thus this non-extensive entropic function reproduces the behaviour of probability distribution. Therefore, it is probable that the correct entropic form for these models is non-extensive Tsallis entropy rather than by Shannon - Boltzmann -Gibbs entropy. The consequences of this conjecture require further investigations.

4 Response to Attack

One of the key features of any network is its ability to withstand various kinds of attacks. An attack or damage to a network typically hampers its performance which leads to a decrease in its efficiency of communication or transfer of signals along various available paths in the network. Self organised networks have various responses depending on the types of attack. Needless to say, this characteristic is of utmost importance for biological contexts where the response to an external attack could determine the survival, recovery or viability of a system. A number of attack strategies are usually considered while evaluating the response. These include targeted attacks (where a particular node or connected segment is attacked based on some of its properties like vertex degree or length), multiple attacks (where random nodes are targeted), isolated attack (where a single node is targeted) etc. In the current simulation we consider isolated and multiple attacks. The measure of response to attack is given by network efficiency as defined in Eq. (19). The response of the network to these attacks is shown in the table where the average change in network efficiency with the number of nodes attacked is shown. This quantity is computed by choosing many nodes either one at a time or in a bunch.
and averaging over the decrease in network efficiency. As can be seen from the table, the network is more susceptible to multiple random attacks rather than to isolated single ones. This is a further manifestation of the fact that the spatio-temporal patterns found in the network are robust with respect to external perturbation such that removal of an element from a local pattern does not impair the network as much as removal of multiple randomly placed elements does. This is also indicative of the fact that there are more than one signalling paths available in any local neighbourhood and unless one blocks a large number of them the efficiency is not significantly impaired.

5 Conclusion

Therefore, in this paper we have proposed a generic model reproducing qualitative behaviour of collective biological systems. The network incorporates a new feature of mobility which has been hitherto missing from such models. This property is shown to be essential in reproducing behaviour in presence of diffusion which many contexts show. The network goes beyond the assumption of boolean elements. It is shown that some of the shortcomings found in boolean networks can be overcome by making the state of an element a continuous variable. Prominent among such enhancements is an ability to model highly heterogenous and diverse environment where the internal state determines and is determined by its surroundings, as in a cellular interaction mechanism. At the same time the network allows us to have a number of states with switching attractors. Thus this model mimics the “robust yet fragile” nature of biological systems.

This explicit coupling between intra and inter dynamics also gives an insight into several topological features of such networks. In particular, we have demonstrated that starting from a particularly simple initial configuration and without explicitly assuming any arbitrarily constructed rule for adding nodes, we can obtain both scale free nature as well as networks containing important hubs. The latter being reminiscent of small world networks. We present evidence for a transition between two regimes upon variation of parameters.

We also show that interesting cooperative phenomena like community structures and synchronisation emerge in these types of networks. This is clearly seen in Fig. 4 showing the spatial profile of internal variables. In this figure the regions of closely linked variations are distinctly separate from other regions.

To investigate robustness of such a network to external perturbations we present a study of two different types of attack on the system. It is found that the multiple random attack is more effective than an isolated one. In current networks the response to the attack is reflected by a decrease in its
efficiency.

The growth and size of the network follows an entropic law which shows signatures of non-extensivity. This leads to the speculation that Tsallis statistics is probably more relevant to biological networks.

Several of these interesting features need further exploration and investigation which are currently under progress. However, in the present paper we only report that many qualitative aspects of the diverse behaviour of cooperation in biological systems can be modeled by a single generic network model. Various aspects of this model can give us an insight into important questions like quorum sensing in bacteria, synchronisation of biological elements, emergence of heterogeniety and morphogeneis, tissue and cell growth, tumour growth, protein interaction pathways and biochemical networks.

For the network theory, this paper presents a network which shows number of different, well known network characteristics in a single model. Therefore, this effort may help in unifying some of different pieces of knowledge which we hope would be useful in understanding general networks better.

Both authors contributed equally to the work. Supplementary materials like programs, data set and additional figures are freely available from the authors upon request.
6 Figure Captions

(1) **Figure 1:** Figure 1 describes the flow of the network evolution. The blocks in the figure describe how various stages of dynamics are handled and how they depend on each other.

(2) **Figure 2:** Figure 2 describes the typical vertex degree distributions obtained. The figures 2(a) and 2(b) are generated by initially setting up a golden mean network with two different kinds of node, namely the Mutual activation switch Eq.(8) and the activator - inhibitor oscillator Eq.(11). The network was evolved with value of $R = 0.35$, and the same values of node parameters as in [Tyson (2003)]. The value of the coupling strength $\epsilon$ was 0.66 for 2(a) and 0.033 for 2(b). Each node was evolve with respect to internal dynamics by using a fourth order unge-Kutta method. The step size for the switch was 0.1 and for the oscillator 0.001.

(3) **Figure 3:** Describes the entropic bound for growth as described in the paper. Fig. 3(a) describes the behaviour where the growth of the network saturates after some time and Fig. 3(b) where it grows without bounds.

(4) **Figure 4:** shows the patterns found in internal profile of the network at various times (a)-(e). The presence of correlated patterns can be easily seen from this figure.

References

[Lee et.al,Milo et.al] Lee T. et al.(2002) Science 298 799
Milo R. et.al(2002) Science 298 824

[Nurse P.] Nurse P.(2000) Cell 100 71

[Hartwell et.al.] Hartwell, L. H., Hopfield, J. J., Leibler, S. Murray, A.W. From molecular to modular biology. Nature 402 (Suppl.), C47C52 (1999),

[Eldar et. al (2002)] Eldar A. et.al.(2002) Nature 419 304

[Ellis E.(2001)] Ellis E.J.(2001) Trends BioChem. Sci. 26 597

[Shibata T. and Kaneko K.] Shibata T. and Kaneko K. arXiv.org : nlin.AO /0204024 and references therein

[Painter K.J and Hillen T.(2002)] Painter K.J. and Hillen(2002) T. Volume filling and Quorum sensing in Models for Chemosensitive Movements Canadian Appl. Math. Quaterly 10(4) 501 and references therein

[Huang S. and Ingber D.E. (2000)] Huang S. and Ingber D.E. Shape dependent Control of Cell growth,Differentiation and Apoptosis: Switching Between Attractors in Cell Regulatory Networks Exp.Cell Research 261 91-103 (2000).
[Shmulevich I. et.al. (2002)] Shmulevich I. et.al. Gene perturbation and Intervention in Probabilistic boolean Networks Bioinformatics 18 1319-1331 (2002).

[Albert, R. and Barabasi, A.-L. (2002).] Albert, R. and Barabasi, A.-L. (2002). Statistical mechanics of complex networks. Rev. Mod. Phys. 74, p47-97 and references therein.

[Lind D. and Marcus B. (1995)] Lind D. and Marcus B. an introduction to Symbolic dynamics and Coding Cambridge university Press (1995)

[Erdos et al(1961); Barabasi(2001); Watts(1998); Newman] Erdos, P. and Renyi, A., On the strength of connect- edness of a random graph, Acta Mathematica Scientia Hungary 12, 261267 (1961).
Jeong, H., Mason, S., Barabasi, A.-L., and Oltvai, Z. N., Lethality and centrality in protein networks, Nature 411, 4142 (2001).
Watts, D. J. and Strogatz, S. H., Collective dynamics of small-world networks, Nature 393, 440442 (1998).

[De Hoon M. et. al.(2003)] De Hoon M. et. al.(2003) Proceedings of the Pacific Symposium on Biocomputing, 8, 1728, 2003.

[de Jong H. et. al.(2003)] de Jong, H., Gouz J.-L., Hernandez, C., Page, M., Sari, T., and Geiselmann, H. (2003). Qualitative simulation of genetic regulatory networks using piecewise-linear models. J. Math. Biol, 66: 301–340

[Levchenko A, Iglesias P(2002)] Levchenko A, Iglesias PA Models of eukaryotic gradient sensing: application to chemotaxis of amoebae and neutrophils. Biophys J 2002, 82 50-63.

[Goldbeter A, Koshland DE(1981)] Goldbeter A, Koshland DE(1981) An amplified sensitivity arising from covalent modification in biological systems. Proc. Natl. Acad. Sci.USA, 78 6840-6844.

[Laurent M and Kellershohn N (1999)]
Laurent M and Kellershohn N: Multistability: a major means of differentiation and evolution in biological systems. Trends Biochem Sci 1999, 24418-422.

[Novak B and Tyson J(1993)] Novak B and Tyson J: Numerical analysis of a comprehensive model of M-phase control in Xenopus oocyte extracts and intact embryos. J. Cell. Sci. 1993, 106 1153-1168.

[Kellershohn N and Laurent M (2001)] Kellershohn N and Laurent M: Prion diseases: dynamics of the infection and properties of the bistable transition. Biophys J 2001, 81 2517-2529.

[Martiel J and Goldbeter A(1987)] Martiel J, Goldbeter A: A model based on receptor desensitiztion for cyclic-AMP signaling in Dictyostelium cells. Biophys J 52 808-828.

[Tyson (2003)] Tyson J. et.al (2003) Current Opinion in Cell Biology 15:221 -231.
[Gell-Mann M. and Tsallis C.(2004)] Gell-Mann M. and Tsallis C. (Editors), Nonextensive Entropy Interdisciplinary Applications (Oxford University Press, New York) 2004
Set up a Golden Mean network and initialize the network and node parameters

Evolve each node by integration using fourth order Runge-Kutta routine

Update the coupling between each node and its neighbors

Update the position and neighbors of each node

Evolve the network by state splitting

Periodic clean up and attack

Are time steps exhausted?

Calculate the network efficiency, vertex degree distribution, entropy and fractal dimension

Fig. 1.
Fig. 2.
Fig. 3.
Fig. 4.