Asynchronous cellular automata - special networks local slowdown produces global speedup

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Asynchronous Cellular Automata - Special Networks

Local Slowdown Produces Global Speedup

by

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A thesis submitted in partial fulfillment
of the requirements for the degree of
Master of Arts
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Asynchronous Cellular Automata - Special Networks
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Arash Khani Ardestani

ABSTRACT

Information processing in living tissues is dramatically different from what we see in common man-made computer. The data and processing is distributed into the activity of cells which communicate only with neighboring cells. There is no clock for the global synchronization of cellular activities. There is not even one bit of central memory for globally shared data. The communication network between cells is highly irregular and may change without changing the outcome of the computation. A simple network of automata is introduced and analyzed to represent a mathematical model of special group of cells in an imaginary tissue sample. The interaction between the cells, their communication method, and their level of intelligence is studied. Three different structures of this model are demonstrated. Later on a simplification in the cells’ program and elimination of a beat keeping clock will lead to a finite state automata network that is surprisingly more powerful in achieving the overall network’s goal than its previous generation which had the advantage of more complex programs and a beat keeping clock.
1 Introduction

Let’s begin with some basic definitions and historical information. The word *automaton* is derived from the Greek word *automatos* meaning ”acting of one’s own will”. Automaton is generally referred to as machines that simulate living organisms’ movements and actions, without any electrical parts or components.

The history of automata can be traced back to at least 3000 years ago. There are many evidences of machines in ancient Greece, including the toys and tools built by *Heron*, representing basic scientific principles. Another example is the ancient Chinese mechanical engineer *Yan Shi* also known as the *Artificer*, who demonstrated a life-size human figure of his mechanical handiwork to King Mu of Zhou.

Throughout the centuries the idea of designing life-like machines and toys continued by scientists and enthusiasts all around the world. The main goal behind efforts has been to create a machine that can self operate, based on existing life forms known to scientists. Although the idea behind automata was studied and practiced throughout those years, but it was mainly focused on the high level behavior of the system. It was not until 20th century when automata was viewed as a new tool to study much lower levels of behavior within the smallest elements of a system. This is essentially what is known as *cellular automata* \(^1\) where we study the activity of the basic elements of a system to determine the overall behavior of that system.

In the 1940’s Stanislaw Ulam developed a new method to study growth of crystals, while his colleague John von Neumann studied the very basics of self replicating machines.

\(^1\)A formal definition of cellular automata is provided in chapter 1.
This is believed to be the birth of what we refer to as the cellular automata. Since von Neumann’s original work on self-replicating machines, much work has been dedicated to studying, designing, and analyzing networks of cellular automata.

In the following paper we will introduce definitions, structures, and analysis of some simple networks of automata. The goal of this paper is to provide a general overview of a special kind of network of automata known as asynchronous network of automata.

This paper is intended to demonstrate natural (i.e. biological) computing modeled on networks of automata. As we introduce different generations of this network, the idea is to show that the more relaxed the conditions and the simpler the design of the network, the more powerful its resulting model will be. Ultimately the goal would be to mathematically prove that the model is in fact capable of this natural computing and that its irregularity will lead to a more ideal model.
2 2PartitioN program on a network

Let us begin by introducing the network and how the 2PartitioN program works. The network consists of finitely many cells each taking one of the two values 0 or 1. In this network each cell is actually a finite-state automaton that is connected to finitely many neighbors by a non-directed edge. The 2PartitioN program is then deployed on this network. The 2PartitioN program makes each cell to be able to change its own value based on the input that receives by reading its neighbors’ values. If the cell sees a neighbor with the same value than its own, it will change. Otherwise it will stay the same.

In Figure 1, white indicates a cell-value of 1 and black indicates 0. 0-0 or 1-1 edges cause instability in the network. Unstable cells are marked with ”u” and stable cells are marked with ”s”.

![Figure 1: An example of 2PartitioN network.](image)

In this network cells are represented by vertices and cell-cell communication links are represented by the edges. Initially all the cells have random values.
The cells are active, or not, randomly. When active, a cell’s value is re-computed according to its program. After thousands of changes, the still-living colony stabilizes — as shown in Figure 3. Is there a global meaning to this stability?

To answer this question we shall spend a fair amount of time in the upcoming sections and investigate the network’s behavior in great detail; however the immediate answer will be given in the next few pages while we introduce the basic design of this model.

A mathematical representation of 2Partition program will be as follows: It exists on a network consisting of a set $C$ of cells, and a set $E \subseteq C^2$ of communication edges between
cells. The cells are defined as copies of an automaton, and cell-cell communication is defined by an input function. \((C, E)\) is assumed to be finite, to have more than one cell, to be connected by non-directed edges, and to have no self-edges.

For 2PartitioN, the automata have values \(Q = \{0, 1\}\) and a value-transition function \(\alpha\).

\[
\alpha(value, input) = \begin{cases} 
1 - value & \text{if } input = value \\
value & \text{otherwise.}
\end{cases}
\]

The input is defined for a multi-set\(^2\) \(M\) of neighboring cell-values.

\[
input(M) = \begin{cases} 
0 & \text{if } 1 \not\in M, \\
1 & \text{if } 0 \not\in M.
\end{cases}
\]

The input describes the neighbors’ values — if \(input = 0\) then at least one neighbor has a cell-value of 0, if \(input = 1\) then at least one neighbor has a value of 1. If \(input\) indicates that a neighbor has the same value as the active cell, then \(\alpha\) returns \((1 - value)\) for an active cell’s next value. This means that as long as there are at least two neighboring cells with the same value the network is unstable; therefore the active cells will continue to change their values until the network goes into a state in which there are no two neighboring cells of the same value; thus becoming completely stable. The network is essentially a bipartite graph, so this behavior is basically the ”global stability” that we mentioned in the previous question. Further in this paper we will re-introduce this concept when we define halting.

A small subgraph of the 2PartitioN network, called 2PartitioN square Network will be the focus of this paper from this point forward. Throughout each section we will visit different arrangements of this network and while analyzing the network behavior we will compare each configuration to another. In the end we will reveal some very interesting and unexpected result which is probably the essence of this paper. Figure 4 shows the basic layout of the square Network.

\(^2\)A multi-set is like a set except that elements may be repeated. As sets \(\{0, 1, 1, 1\}\) and \(\{0, 1\}\) are equal, but as multi-sets they are different.
In this network cells are four vertices in the four corners of a square, with the sides of the square representing the edges between the four vertices of a simple graph. Each vertex can take a value 0 or 1. Obviously there are $4^2 = 16$ distinct configurations, since we have four cells and for each cell we have two choices (i.e. 0 and 1). A complete list of these states is shown below:

Figure 5: All possible states of the 2PartitioN square network.

One may notice a symmetry between different states of the model, but it is necessary to point out that while the network architecture is highly symmetric, we make absolutely no use of this symmetry anywhere throughout this paper. In other words we insist on irregular (communication) architecture. The square design was solely chosen due to its simplicity which makes the concept easier to digest; however the entire idea is applicable
to any irregular network such as the one shown in figure-1 on page 3.
More particularly we refer to these configurations as "network states", "global states" or with a more general term states. Next we define a set of simple rules that govern the activity of the cells in the network:

1. two cells are considered neighbors if there is an edge between them.
2. each cell can read the value of its neighbors.
3. a cell changes its value if at least one of its neighbors has the same value as the cell, only if the cell is active. A graphical representation of this network is shown below:

![Graphical representation of a network]

In mathematical language we can represent: Let $C$ be the set of cells such that $C = \{a, b, c, d\}$ and let $E \subseteq C^2$ be the set of edges that denote the communication edges between cells. The values of cells is defined as the set $Q = \{0, 1\}$. The transition function is the same as was defined earlier. If a cell is an active cell then it reads the values of its neighbors and computes its own value at that given moment. Otherwise the cell is inactive.

Figure 6: General state transition.
3 2PartitioN program on a square network with synchronous activity

Suppose we randomly pick one of the states of 2PartitioN network. If all cells are active at the same time, then we call this network a synchronous network. In other words, a cell is only active if all of its neighbors are active at the given moment. Let’s look at an example in more details:

![Figure 7: One-step state transition.](image)

The network flips between two states forever, since it loops between two states. These states are shown in the figure above. Let us look at another example:
In this configuration the activity of each cell is limited to only a set of non-halting configurations since it depends on the activity of all its neighbors. Such behavior makes this configuration of 2PartitioN a rather not interesting network to study. For example given a random initial state [0011] the next state will be [1100]. If the network is activated again, it will immediately go to next state [0011]. Obviously this network in synchronous mode will flip back and forth between maximum two states, except for the two halting states [0101] and [1010]. The two halting states never change to another state, regardless of cell activity. This fact is shown in figure-10 below. The reason for this behavior is that in synchronous mode all the cells are active at the same time. In other words all of the four cells read the values of their neighbors at the same time and change their values based on the rules defined under the transition function. Complete state diagram is demonstrated below:
Figure 9: State transitions of the synchronous square network.
As seen above the number of steps from one state to another is limited to maximum of two step. This criteria of the synchronous 2PartitioN network makes its behavior very simple and thus its next states very predictable. Another interesting fact about the synchronous network is that it never halts, unless it is already in a halting state. What this means for the 2PartitionN synchronous network is that this mode of activity cannot be used to model the partitions of bipartite networks, since it is limited to only a few transition states and it never halts.

Another very important aspect of this network is that in order for all cells to become active at once, there is an inevitable need for a clock. What we mean by a clock in this case is a central process or program which is accessible by all cells and acts as a beat keeper. It is obvious that the need for a central clock in any network will be interpreted as a disadvantage in the efficiency of the communication over the entire network. The lack of efficiency in communication may not be apparent in a small network such as the 2PartitioN square serial; however in a significantly larger network it certainly introduces a real challenge. As an example consider \(10^{12}\) skin cells of the human body trying to read a central clock every 0.050 seconds. One can imagine how this would impact the network in terms of the need for additional communication links between each cell and the central clock, not to mention the extra time needed for each communication transaction.
4 2PartitioN program on a square network with serial activity

Suppose we are given the same 2PartitioN network. Assuming we are given any initial state of the network, but we allow only one cell $c_1, c_2, c_3, c_4$ to be active at any given time. The resulting network is what we generally refer to as 2PartitioN serial network. The network’s activity is serial in the sense that its cells take active role randomly but one cell at a time. Below there is a demonstration of how this network might behave:

![Diagram of 2PartitioN serial activity]

Figure 10: Behavior of the 2PartitioN network with serial activity.

It is worthy to note that the cells do not need to take active role in any specific order. A major difference between the serial network and the synchronous network that was discussed in previous section is that the serial network is a *halting* network, when the network is bipartite and cell activity is random. In other words this network can go from non-halting initial state to a halting state. This criteria gives this network a real advantage over the synchronous network. This is due to the fact that a halting network of this type can solve the problem of bipartite partitioning. Below there is a complete
demonstration of all state transitions from initial states to the next states or halting states:

Figure 11: Global state transitions of 2PartitioN network with serial activity.
We can compute the average halting time for the 2PartitioN square network with serial activity. To do so we need to formulate the expected halting time in relation to the transition probability and the probability of activity for a given initial state:

\[ H_\alpha = 1 + \sum P_\mu H_\mu \]

- \( H_\alpha \) the expected halting time from the state \( \alpha \) for example \( H_{0011} \)
- \( \mu \) the state to which we have a single step transition from state \( \alpha \)
- \( H_\mu \) the expected halting time for the state \( \mu \) for example \( H_{0111} \)
- \( P_\mu \) the probability that state \( \mu \) will have one step transition from state \( \alpha \).

Note: In this case \( P_\mu = 1/4 \) for all \( \mu \). As an example let us write a formula for calculating the expected halting time for state \( \alpha = 0011 \):

\[
H_{0011} = 1 + \left[ \frac{1}{4}(H_{1011}) + \frac{1}{4}(H_{0111}) + \frac{1}{4}(H_{0001}) + \frac{1}{4}(H_{0010}) \right]
\]

To find the average expected halting time we will follow these simple steps:

Step 1: Using global state transition diagram find the expected halting time for each initial state in terms of other

Step 2: Write the equation for each expected halting time in terms of each initial state and other states

Step 3: Set these equations as a system with 16 equations and 14 unknowns and solve it using Maple

Step 4: Find the average of the expected halting times

Let us begin with step 1:

\[
H_{0011} = 1 + [1/4(H_{1011}) + 1/4(H_{0111}) + 1/4(H_{0001}) + 1/4(H_{0010})]
\]

\[
H_{1011} = 1 + [1/4(H_{0011}) + 1/4(H_{1010}) + 1/4(H_{1001}) + 1/4(H_{1011})]
\]

\[
H_{1111} = 1 + [1/4(H_{0111}) + 1/4(H_{1011}) + 1/4(H_{1101}) + 1/4(H_{1110})]
\]

\[
H_{1001} = 0
\]

\(^3\)There are only 14 unknowns because \( H_{0110} \) and \( H_{1001} \) are zero
\begin{align*}
H_{1010} &= 1 + [1/4(H_{0010}) + 1/4(H_{1110}) + 1/4(H_{1000}) + 1/4(H_{1011})] \\
H_{1101} &= 1 + [1/4(H_{0101}) + 1/4(H_{1100}) + 1/4(H_{1001}) + 1/4(H_{1101})] \\
H_{1110} &= 1 + [1/4(H_{1100}) + 1/4(H_{1010}) + 1/4(H_{0110}) + 1/4(H_{1110})] \\
H_{1000} &= 1 + [1/4(H_{1100}) + 1/4(H_{1010}) + 1/4(H_{1101}) + 1/4(H_{1000})] \\
H_{1100} &= 1 + [1/4(H_{0100}) + 1/4(H_{1000}) + 1/4(H_{1110}) + 1/4(H_{1101})] \\
H_{0111} &= 1 + [1/4(H_{0111}) + 1/4(H_{0101}) + 1/4(H_{0110}) + 1/4(H_{0111})] \\
H_{0101} &= 1 + [1/4(H_{1101}) + 1/4(H_{0001}) + 1/4(H_{0111}) + 1/4(H_{0100})] \\
H_{0110} &= 0 \\
H_{0100} &= 1 + [1/4(H_{1100}) + 1/4(H_{0110}) + 1/4(H_{0101}) + 1/4(H_{0100})] \\
H_{0001} &= 1 + [1/4(H_{1001}) + 1/4(H_{0101}) + 1/4(H_{0111}) + 1/4(H_{0001})] \\
H_{0000} &= 1 + [1/4(H_{1000}) + 1/4(H_{0010}) + 1/4(H_{0011}) + 1/4(H_{0001})] \\
H_{0010} &= 1 + [1/4(H_{1010}) + 1/4(H_{0110}) + 1/4(H_{0011}) + 1/4(H_{0010})]
\end{align*}

After solving the system of equations using Maple, the following values are obtained:

\begin{align*}
H_{0011} &= 7.062499992 & H_{1011} &= 6.083333325 \\
H_{1111} &= 7.166666657 & H_{1001} &= 0.0 \\
H_{1010} &= 7.187499990 & H_{1101} &= 6.083333325 \\
H_{1110} &= 6.458333325 & H_{1000} &= 6.124999992 \\
H_{1100} &= 7.187499992 & H_{0111} &= 6.041666658 \\
H_{1011} &= 7.062499988 & H_{0110} &= 0.0 \\
H_{0100} &= 6.083333325 & H_{0001} &= 6.041666658 \\
H_{0000} &= 7.083333323 & H_{0010} &= 6.083333325
\end{align*}

The average expected halting time is 6.553571419

As mentioned earlier the basic property of a serial network is apparent in the way the cell activity takes place. For the purpose of 2Partition square network with serial activity, only one of the four cells can be active at a time. There is an inevitable need for a mechanism to ensure that no more than one cell can be active at a given time. There are different methods to implement this mechanism. One mechanism would be using a
clock just like the clock mentioned in the 2Partition square network with synchronous activity. Another mechanism would be to use a token, similar to the technology used in token ring networks. Depending on the size and structure of the network one mechanism may be preferred over another, but that subject is out of the scope of this article. In any event enforcing serial activity on this model requires a substantial computational effort. The ideal goal with 2PartitioN network -cellular network of automata- is to be able to design the cells in such a way that the memory requirement is minimal to none and that they are programmed in the simplest way possible. In the next section we will describe a special kind of 2PartitioN network that is the closest to the ideal model.
5 2PartitioN program on a square network with asynchronous activity

Considering the same 2PartitioN network that we have been analyzing so far, but the major different this time is that we allow any of the cells to be active at any given time. Even though this change in the cells’ activity may seem minor at first, yet deeper analysis of the network proves that the impact on the network’s behavior and more importantly its halt-ability is absolutely significant. We introduce this network as 2PartitioN square asynchronous. The word asynchronous here is meant to describe the randomness in each cell’s activity. At first glance it might appear as if the asynchronous network design will be more complex but it turns out that the asynchronous network does not need a clock and each cell is designed with a much simpler program, compared to the networks examined in the previous sections. A major difference between the 2PartitioN network with asynchronous activity and the 2PartitioN network with synchronous activity is that the first one will always halt, while the latter may or may not halt.

It is interesting to notice that the state transition of the 2PartitioN square network with serial and synchronous activity are both special cases of the 2PartitioN square network with Asynchronous activity. This also means that the global state diagrams of the 2PartitioN square network with serial and synchronous activity are subsets of the global state diagram for the 2PartitioN square network with asynchronous activity. In 2PartitioN square asynchronous network we relax the rules that govern the activity of the cells; thus giving the network more flexibility and perhaps more complexity.

Before going further let us demonstrate a figure to familiarize ourselves with how this network behaves.
Obviously this process continues until it reaches any of the two halting states. As mentioned earlier the $2$PartitioN square network is a halting network. It is important to mention that $2$Partition must halt on bipartite graphs.

**Theorem:** $2$PartitioN on square network in asynchronous mode will always halt.

**Proof:**

Suppose we are at one of the random initial states. The greatest probability that the next state is not one of the halting states is at most $P$ where $P < 1$. Suppose we go to the next non-halting state and again the probability of going to the next non-halting state is $\leq P$. If we continue in this fashion for $n$ steps, then the probability that we do not reach a halting state in $n$ steps is $\leq P^n$. On the other hand the probability that the network halts in $n$ steps is $\leq 1 - P^n$. As $n$ goes to $\infty$ then $P^n$ approaches 0 therefore $1 - P^n$ approaches 1.

$$\lim_{n \to \infty} (P^n) = 0 \Rightarrow \lim_{n \to \infty} (1 - P^n) = 1$$

This shows that $2$PartitioN on square network in asynchronous mode will always halt with probability 1. To make this point clearer let’s demonstrate the case below:
In the figure on the left the P is calculated as $P = (1 - 1/8) = 7/8$ where in the right figure we have $P = 1 - 1 = 0$.

Figure below represents the global state diagram of the 2PartitioN square asynchronous network:
Figure 14: Global state diagram - 2PartitioN square asynchronous Network.

To demonstrate this more clearly let’s present the next figure:
The figure above captures a portion of the previous diagram and demonstrates how each initial state is related to other states including the halting states. Clearly not all the states are shown, due to lack of space to include all arrows and all states, but the figure delivers the point. Perhaps, the next few paragraphs explain the dynamics of the network better. To obtain the next states from each initial state, we consider all possible
combinations of active cells a, b, c, and d, skipping the repetitions of course. In general we will consider the following 16 cases:

- None of the cells are active
- Case 1: No cell is active
- Just single cell being active
- Case 2: Only cell "a" is active
- Case 3: Only cell "b" is active
- Case 4: ....

.........

All combinations of two cells being active at a time
- Case 6: Cells "a" and "b" are active
- Case 7: Cells "a" and "c" are active
- Case 8: ....

.........

.........

All combinations of three cells being active at a time
- Case 12: Cells "a" and "b" and "c" are active
- Case 13: Cells "a" and "b" and "d" are active
- All four cells active at a time
- Case 16: Cells "a" and "b" and "c" and "d" are active

Mathematically we are basically taking all the subsets of the set of cells \( C = \{a, b, c, d\} \) to obtain the set of active cells. In each case the members of each subset define the active cells for that case.

Let us examine one of initial random states and see how it would go to all other possible states:
Figure 16: State transition from state A to all other possible states.

Now that we have seen this construction one can easily see the relation between each initial state and the next possible states demonstrated in figure-16. One can quickly notice that some of the initial random states can only change to eight of the sixteen possible next states. The reason behind this configuration is that in some of the random initial states one of the cells is what we call a stable cell. A stable cell is a cell that does not have any neighbor with the same value as itself; thus it has no effect on the computation of the next state. According to the transition function defined in Section 1, the stable cell does not need to change its value regardless if it is active or not—alone or in any combination with other cells. Let us demonstrate this behavior in the following figure, where cell "b" is the stable cell:
So far we have established the fact that the 2ParitioN square network with asynchronous activity will always halt. Additionally we have determined a complete global transition graph, which shows the relation between every possible random initial state and all the other states, including the halting states. Now it would be interesting to examine each random initial state to see how the activity of each cell can affect the next transition state. To study and analyze this process we will take advantage of the notion of transition probability. What we mean by transition probability is the probability that a given random initial state would go to the next possible state. We will study the relation between the probability of activity (for each cell in a given random initial state) and the transition probability. Let us begin with the following example:

Figure 18: Example 1 of how transition probabilities are computed.
For each active cell we assign the value $p$ and for each inactive cell we assign the value $(1 - p)$. If there is a stable cell, the process is the same, but the total probabilities will be added together just like in the following example:

In example 1 to compute the total probabilities going from 1010 to 1110 we have:

$$P_{1010 \rightarrow 1110} = (1 - p)^3(p)$$

In example 2 to compute the total probabilities going from 0111 to 0110 we have:

$$P_{0111 \rightarrow 0110} = (1 - p)^3(p) + (1 - p)^2(p)^2$$

By design, we let each cell in the square network with asynchronous activity to be active or inactive at any given time. Therefore the probability of a cell being active is $p = 1/2$. Similarly the probability of a cell being inactive is also $1/2$ (i.e. $1 - p = 1/2$). There are four cells in the square network, so the total probability for going from one state to another is calculated as $1/16$. Of course in some of the states where we have stable cells, the probability would be $2P = 1/8$.

As seen in figure-17 each random initial state will end on at least one of the halting states. This is visual confirmation of the proof of Theorem-1. The fact that this network is a halting network brings up a few interesting questions:
What is the expected halting time of the 2PartitioN asynchronous network?

Does the choice of initial state affect the average expected halting time?

To answer these questions we will use the same method as we used in Section 3 pages 13-14:

\[ H_\alpha = 1 + \sum P_\mu H_\mu \]

- \( H_\alpha \) the expected halting time from the state \( \alpha \) for example \( H_{0011} \)
- \( \mu \) the state from which we have a single step transition from state \( \alpha \)
- \( H_\mu \) the expected halting time for the state \( \mu \) for example \( H_{0111} \)
- \( P_\mu \) the probability that state \( \mu \) will be obtained in one step transition from state \( \alpha \).

It is important to note that in the current case \( P_\mu = 1/16 \) for all \( \mu \). So if we were to write a formula for calculating the expected halting time for state \( \alpha = 0011 \) we would have:

\[
H_{0011} = 1 + \left( \frac{1}{16}(H_{0011}) + \frac{1}{16}(H_{1011}) + \frac{1}{16}(H_{1111}) + \frac{1}{16}(H_{1001}) + \frac{1}{16}(H_{1010}) \right) \\
+ \frac{1}{16}(H_{1101}) + \frac{1}{16}(H_{1110}) + \frac{1}{16}(H_{1000}) + \frac{1}{16}(H_{1100}) + \frac{1}{16}(H_{0111}) + \frac{1}{16}(H_{0101}) \\
+ \frac{1}{16}(H_{1001}) + \frac{1}{16}(H_{0100}) + \frac{1}{16}(H_{0001}) + \frac{1}{16}(H_{0000}) + \frac{1}{16}(H_{0010})
\]

Using the global state diagram and the activity probability \( P \) for each state we can write the equations of halting for each of the random initial state as follows:

\[
H_{0011} = 1 + \frac{1}{16}[H_{0011} + H_{1011} + H_{1111} + H_{1001} + H_{1101} + H_{1110} + H_{1000} + H_{1100} + H_{0111} + H_{0101} + H_{0110} + H_{0001} + H_{0000} + H_{0010}]
\]

\[
H_{1011} = 1 + \frac{1}{16}[2H_{0011} + 2H_{1011} + 2H_{0001} + 2H_{1000} + 2H_{1010} + 2H_{1001} + 2H_{0010} + 2H_{0000}]
\]
\[ H_{1111} = 1 + \frac{1}{16}[H_{0011} + H_{1011} + H_{1111} + H_{1001} + H_{1010} + H_{1101} + H_{1110} + H_{1000} + H_{1100} + H_{0111} + H_{0101} + H_{0110} + H_{0100} + H_{0001} + H_{0000} + H_{0010}] \]

\[ H_{1001} = 0 \]

\[ H_{1010} = 1 + \frac{1}{16}[H_{0011} + H_{1011} + H_{1111} + H_{1001} + H_{1010} + H_{1101} + H_{1100} + H_{0111} + H_{0101} + H_{0110} + H_{0100} + H_{0001} + H_{0000} + H_{0010}] \]

\[ H_{1101} = 1 + \frac{1}{16}[2H_{1101} + 2H_{0101} + 2H_{0001} + 2H_{0100} + 2H_{0000} + 2H_{1001} + 2H_{1000} + 2H_{1110} + 2H_{1100}] \]

\[ H_{1110} = 1 + \frac{1}{16}[2H_{1110} + 2H_{0110} + 2H_{0010} + 2H_{0100} + 2H_{0000} + 2H_{1010} + 2H_{1000} + 2H_{1100}] \]

\[ H_{1000} = 1 + \frac{1}{16}[2H_{1000} + 2H_{1100} + 2H_{1010} + 2H_{1001} + 2H_{1110} + 2H_{1101} + 2H_{1111} + 2H_{1011}] \]

\[ H_{1100} = 1 + \frac{1}{16}[H_{0011} + H_{1011} + H_{1111} + H_{1001} + H_{1010} + H_{1101} + H_{1100} + H_{0111} + H_{0101} + H_{0110} + H_{0100} + H_{0001} + H_{0000} + H_{0010}] \]

\[ H_{0111} = 1 + \frac{1}{16}[2H_{0111} + 2H_{0011} + 2H_{0101} + 2H_{0001} + 2H_{0010} + 2H_{0100} + 2H_{0000}] \]

\[ H_{0101} = 1 + \frac{1}{16}[H_{0011} + H_{1011} + H_{1111} + H_{1001} + H_{1010} + H_{1101} + H_{1110} + H_{1000} + H_{1100} + H_{0111} + H_{0101} + H_{0110} + H_{0100} + H_{0001} + H_{0000} + H_{0010}] \]

\[ H_{1001} = 0 \]

\[ H_{0100} = 1 + \frac{1}{16}[2H_{0100} + 2H_{1100} + 2H_{1110} + 2H_{1101} + 2H_{0110} + 2H_{0101} + 2H_{0011}] \]

\[ H_{0001} = 1 + \frac{1}{16}[2H_{0001} + 2H_{1001} + 2H_{1101} + 2H_{1011} + 2H_{0111} + 2H_{0101} + 2H_{0011} + 2H_{0001}] \]

\[ H_{0000} = 1 + \frac{1}{16}[H_{0011} + H_{1011} + H_{1111} + H_{1001} + H_{1010} + H_{1101} + H_{1110} + \]
\[ H_{100} + H_{110} + H_{011} + H_{010} + H_{100} + H_{001} + H_{000} + H_{0010} \]

\[ H_{0010} = 1 + 1/16[2H_{0010} + 2H_{1010} + 2H_{1110} + 2H_{1011} + 2H_{1111} + 2H_{0110} + 2H_{0111} + 2H_{0011}] \]

We consider this as a system of equations with 14 unknowns where \( H_x \) interprets as the expected halting time for state \( x \). Solving this system of equations will result in a numerical value for the expected halting time for each of 16 possible states.

\[
\begin{align*}
H_{0011} &= 8.031249990 & H_{1011} &= 8.028124995 \\
H_{1111} &= 8.093749991 & H_{1001} &= 0.0 \\
H_{1010} &= 8.031249986 & H_{1101} &= 8.028124997 \\
H_{1110} &= 8.028124997 & H_{1000} &= 8.034374998 \\
H_{1100} &= 8.031249992 & H_{0111} &= 8.028124996 \\
H_{0101} &= 8.031249992 & H_{0110} &= 0.0 \\
H_{0100} &= 8.034374996 & H_{0011} &= 8.034374991 \\
H_{0000} &= 8.031249992 & H_{0010} &= 8.034374998 .
\end{align*}
\]

The average expected halting time is easily calculated to be 8.035714279

Clearly changing the probability of activity for each cell will change the transition probability. This brings up a few more interesting questions:

- How will changing cell activity probability affect the transition probability?
- Will changing the transition probability affect the expected halting time?
- How will the change in transition probability affect the expected halting time?

To answer these questions we write the halting equation for each of the random initial states leaving the cell activity probability as a variable \( p \). To find the minimum halting time we differentiate the equation again with \( p \) staying as a variable. The idea is to write generic equations based on the formula bottom of page 22 and equations shown on pages

\[ ^4 \text{There are only 14 unknowns because } H_{0110} \text{ and } H_{1001} \text{ are zero} \]
23 - 24, except that not the probability of activity for each cell is left as variable p. To deliver a clearer point let’s look at the following example: Suppose we are at the random initial state [0011] and we want to compute the expected halting. The base formula is:

\[ H_\alpha = 1 + \sum P_\mu H_\mu \]

So in this case \( \alpha \) is 0011. To find the \( p_\mu \) we will refer to the global state diagram to see how 0011 will change to other states and find those transition probabilities based on probability of activity:
As explained previously on pages 21 - 22, to write the transition probability we combine the probability of activity of the cells for the random initial state. If the cell is active its probability of activity is \( p \) and if it is inactive the probability of activity is \( 1-p \). The figure above shows all the transition probabilities from initial state \([0011]\) to all other possible next states (including itself). Now it should be very simple to find the expected halting time for the given initial state \([0011]\). To find the expected halting time in terms of \( p \) we can write:

\[
H_{0011} = (1-p)^4[H_{0011}+1]+p(1-p)^3[H_{1011}+1]+p(1-p)^3[H_{0111}+1]+p(1-p)^3[H_{0001}+1]
+p(1-p)^3[H_{0010}+1]+p^2(1-p)^2[H_{1111}+1]+p^2(1-p)^2[H_{1001}+1]+p^2(1-p)^2[H_{1010}+1]
\]
\[+p^2(1-p)^2[H_{0101} + 1] + p^2(1-p)^2[H_{0110} + 1] + p^2(1-p)^2[H_{0000} + 1] + p^3(1-p)[H_{1101} + 1]
+p^3(1-p)[H_{1110} + 1] + p^3(1-p)[H_{1000} + 1] + p^3(1-p)[H_{0100} + 1] + p^4[H_{1100} + 1]\]
Let us demonstrate this point with another figure and example, where now the initial state has a stable cell:

**Figure 21:** Transition probabilities for an initial state with stable cell

Similarly to find the expected halting time in terms of \( p \) for initial state [1101] we have:

\[
H_{1101} = [(1-p)^4 + p(1-p)^3][H_{1101} + 1] + [p(1-p)^3 + p^2(1-p)^2][H_{0101} + 1] \\
+ [p(1-p)^3 + p^2(1-p)^2][H_{1001} + 1] + [p(1-p)^3 + p^2(1-p)^2][H_{1100} + 1] \\
+ [p^2(1-p)^2 + p^3(1-p)][H_{0001} + 1] + [p^2(1-p)^2 + p^3(1-p)][H_{0100} + 1] \\
+ [p^2(1-p)^2 + p^3(1-p)][H_{1000} + 1] + [p^3(1-p) + p^4][H_{0000} + 1]
\]

Now we can write similar equations corresponding to each of the 16 initial states and
find the expected halting time for each initial state in terms of p as a variable. We can then set these equations as a system with 16 unknowns. Once the equation is solved we will be left with one final equation in terms of p as a variable. Clearly we can find the minimum value for that equation by differentiating with respect to p and finding the critical points. Next we devise a Maple program to calculate the halting time with \( P \) being a simple variable and all other factors fixed. A simple loop is setup to change the \( P \) starting from 0.999 going down to 0.001 stepping every 0.001. Then this experimental data set is graphed against the calculated halting time. The result is quite amazing. The following figure shows that as the probability of activity approaches 0.4 the halting time becomes lower.

![Figure 22: Halting time vs. probability of activity.](image)

To see the results more clearly let us present the next figure:
By solving the equations we find that at $P = 0.406$ the halting time is the lowest halting time for the 2PartitioN square asynchronous network being precisely at 7.5838286. Interestingly enough this halting time is certainly lower than the one we calculated using the 16 equations explored earlier. In addition this lower halting time was achieved using a lower probability of activity than $P = 1/2$. It is also important to note that when the probability of activity for the cells is close to 1, meaning each cell is active at all times, the halting time approaches infinity. In other word, it will take this network forever to halt.

Figure 23: Close up of halting time vs. probability of activity.
6 Philosophy and conclusion

The results obtained in previous section are not what we would expect. Imagine with this network was a simple computer with four processors, trying to solve a problem. In reality by increasing the activity of each processor one would expect to solve the problem faster, but in this case we realized that slowing down each processor just enough, would make the simple computer solve the problem faster. Let us consider a scenario where we have finitely many of these 2PartitioN square asynchronous networks in a large grid. Each 2PartitioN square network has four links to four other 2PartitioN network. All the squares are at rest in a random initial state. Randomly we activate one of the 2PartitioN square networks. As soon as this one network reaches the halting state, it activates its four links to other four neighboring networks. Then each of those four neighboring networks become active and begin to work their way towards halting. This process continues for a finite period of time, until all the 2PartitioN squares are halted in the entire grid. This process very much simulates the healing process of an organ or tissue in a living organism. Suppose each 2PartitioN square network in this grid is setup with a probability of activity as $P = 1/2$. Then we can easily predict the average time for given number of cells to reach their goal, in this case healing of the entire tissue. Obviously if there is a special substance that can lower the probability of activity in each cell, thus slowing down the 2PartitioN square networks to the desired level –remember the $P = 0.406$– then we can certainly achieve a significantly faster healing process when working with large tissues consisting of millions of cells. So far quite a few mathematicians have attempted to provide the closest and most ideal mathematical model to cell interaction and growth as seen in nature. The approach to this mathematical model may seem simple; however it provides a bridge between traditional computation models and new approaches to understanding cellular networks in nature. We have all observed the usual computation models in any regular computing machine, anything from a simple Turing machine to a fast super computer. The idea has always been to use highly regular and organized models with strict set of rules governing the model and its elements. Today’s science
however demands one to break free of regularity and order, to look at a model seemingly overwhelmed in chaos, but with just enough order to represent a sophisticated model of cellular activity in nature.
References


Stark W.R. Asynchronous Networks of Automata - A Study of Emergent Phenomena from Mathematical Analysis to Biologically Inspired Applications, Department of Mathematics and Statistics - University of South Florida, Tampa Florida


Appendix A

\( V := \{1, 2, 3, 4\} \)

\( E := \{(0, 1, 0, 1), (1, 0, 1, 0), (0, 1, 0, 1), (1, 0, 1, 0)\} \)

\( V = \{1, 2, 3, 4\} \)

\[
E = \begin{pmatrix}
0 & 1 & 0 & 1 \\
1 & 0 & 1 & 0 \\
0 & 1 & 0 & 1 \\
1 & 0 & 1 & 0 \\
\end{pmatrix}
\]

\( N := \text{proc } (s) \text{ local } t, u, v; \text{ global } V, E; \)
\[ t := \text{copy}(s); \]
\[ \text{for } v \text{ in } V \text{ do} \]
\[ \text{if } s[v] = 1 \text{ then} \]
\[ \text{for } u \text{ in } V \text{ do} \]
\[ \text{if } E[u, v] = 1 \text{ and } s[u] = s[v] \text{ then } t[v] := 1 - s[v]; \]
\[ \text{fi}; \text{od}; \]
\[ \text{fi}; \text{od}; \]
\[ t; \]
\[ \text{end}; \]

\( N(0, 1, 0, 1, 1, 1, 1) \)

\[
\begin{pmatrix}
0 \\
1 \\
0 \\
1 \\
\end{pmatrix}
\]

\( \text{MakeIndex := proc } (s) \)
\[ \text{end}; \]

\( \text{MakeLabel := proc } (\text{ind}) \text{ local } i, \text{lbl}, x; \)
\[ x := \text{ind}-1; \]
\[ \text{lbl} := <0, 0, 0, 0>; \]
\[ \text{for } i \text{ to } 4 \text{ do} \]
\[ \text{lbl}[i-1] := x-2*\text{floor}(1/2*x); \]
\[ x := \text{floor}(1/2*x); \]
\[ \text{od}; \]
\[ \text{lbl}; \]
\[ \text{end}; \]

\( \text{Inds := [n \&\& n=1 .. 16]}; \)
\( \text{Lbsa := map(MakeLabel, Inds)}; \)
\( \text{map(MakeIndex, Lbsa)}; \)
\[ \text{Inds} = [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16] \]

\[ \text{Lhs} = \begin{bmatrix}
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 1 & 1 & 0 & 0 & 1 & 1 & 0 & 0 & 1 & 1 & 0 \\
0 & 1 & 0 & 1 & 0 & 1 & 1 & 0 & 1 & 0 & 1 & 0 & 1 & 0 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
1 & 0 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
\end{bmatrix} \]

\[ \text{map}(a \rightarrow N(<0, 0, 0, a>, \text{lhrs})); \]

\[ \begin{bmatrix}
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 & 0 & 0 & 0 & 0 & 1 & 1 \\
0 & 0 & 1 & 1 & 1 & 0 & 0 & 1 & 1 & 0 & 0 & 1 & 1 & 0 \\
0 & 1 & 0 & 1 & 0 & 1 & 1 & 0 & 1 & 0 & 1 & 0 & 1 & 0 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
\end{bmatrix} \]

\[ a\text{Neighbors} := \text{proc} \{a\} \text{ local } a; \text{ global } lhrs, E, V; \]
\[ \text{map}(a \rightarrow N(a, a), \text{lhrs}); \]
\[ \text{end}; \]

\[ a\text{Neighbors}(<0, 1, 0, 1>); \]

\[ \begin{bmatrix}
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{bmatrix} \]

\[ a\text{Neighbors}(<1, 1, 1, 1>); \]

\[ a\text{Neighbors}(<1, 1, 1, 1>); \]

\[ a\text{Neighbors}(<1, 1, 1, 1>); \]

\[ a\text{Neighbors}(<1, 1, 1, 1>); \]
Transition probabilities are computed from the probability $p$ of cell activity and the stability/unstability of cells.

\begin{equation}
\begin{align*}
\text{unstable} & := \text{proc } (s) \text{ local } i, u, t, v : \\
& \quad \text{unstable} := <0,0,0,0> : \\
& \quad \text{if } s[1] = s[4] \text{ or } s[1] = s[2] \text{ then } u, t, v[1] := 1 ; \\
& \quad \text{unstable}.
\end{align*}
\end{equation}

\begin{equation}
\begin{align*}
\text{tProb} & := \text{proc } (a, t) \text{ local } i, j, c, r, q, u, t, v : \\
& \quad \text{global } p : \\
& \quad q := 1 ; \\
& \quad \text{unstable} := \text{unstable}(a) ; \\
& \quad \text{sum} := 0 ; \\
& \quad c := [1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1] ; \\
& \quad \text{convert}(t, \text{list}) : \\
& \quad \text{for } j \text{ from } 1 \text{ to } 16 \text{ do} \\
& \quad \quad \text{if } \text{not}(\text{Equal}(a, a, \text{neighs}(a)[j], \text{compare} \text{entries})) \\
& \quad \quad \text{then } c[j] := 0 \\
& \quad \quad \text{sum} := \text{sum} + c[j] ; \\
& \quad \quad \text{od} ; \\
& \quad \text{if } \text{sum} = 0 \text{ then } q := 0 ; \\
& \quad \text{else} \\
& \quad \quad \text{for } i \text{ from } 1 \text{ to } 4 \text{ do} \\
& \quad \quad \quad \text{if } a[i] = a[2][i] \text{ and } u, t, v[i] = 1 \text{ then } q := q \cdot (1 - p) ; \\
& \quad \quad \quad \text{elseif } a[i] = a[2][i] \text{ and } u, t, v[i] = 1 \text{ then } q := q \cdot p ; \\
& \quad \quad \quad \text{else } q := q' ; \\
& \quad \quad \quad \text{fi} ; \\
& \quad \quad \text{od} ; \\
& \quad \text{end} ; \\
& \quad \text{tProb}(0, 0, 0, 0, 0, 0, 0, 0, 1) ; \\
& \quad \text{end} ; \\
& \quad \text{map}(\text{tProb}(0, 0, 0, 0, 0, 0, 0, 0, 1), \text{Labs}) ; \\
& \quad \text{map}(\text{tProb}(0, 0, 0, 0, 0, 0, 0, 0, 1), \text{Labs}) ;
\end{align*}
\end{equation}
map(s -> tProb(<0,0,1,0>, s), Lbla);  
map(s -> tProb(<0,0,1,1>, s), Lbla);  
map(s -> tProb(<0,1,0,0>, s), Lbla);  
map(s -> tProb(<0,1,0,1>, s), Lbla);  
\[
\begin{align*}
\end{align*}
\]

\[(8)
T is the matrix of transition probabilities.

\> T := Matrix(16,16,0);
\>
\> for j from 1 to 16 do
\> for i from 1 to 16 do
\> T[i,j] := tProb(Lbla[j], Lbla[i]);
\> od:
\>
\> T[1..4,1..4];
\>
\[
\begin{pmatrix}
(1-p)^4 & 0 & 0 & (1-p)^2 p^2 \\
(1-p)^3 p & (1-p)^3 & 0 & (1-p) p^3 \\
(1-p)^2 p & 0 & (1-p)^3 & (1-p) p^3 \\
(1-p) p^2 & (1-p)^2 p^2 & (1-p)^2 p & (1-p)^3
\end{pmatrix}
\]

\[(9)
Arash, what is the meaning of your dot product?

\> states := Vector(<s1,s2,s3,s4,s5,s6,s7,s8,s9,s10,s11,s12,s13, s14,s15,s16>);  
evals := Vector(T[1..16,3]);
states :=
evals[DotProduct(4, b)];
\[
\begin{pmatrix}
11 & 12 & 13 & 14 & 15 & 16 \\
11 & 12 & 13 & 14 & 15 & 16 \\
11 & 12 & 13 & 14 & 15 & 16 \\
\end{pmatrix}
\]

\[(10)
To avoid worrying about complex values assume that \( p \) is positive real.
\>
\> assume(p > 0);

The transition probabilities from each of \( s_1, s_2, \ldots, s_{16} \) are computed as \( T \cdot v \) (a matrix product).

\> w := Vector(16): v[3] := 1;
\> evals := T \cdot v;

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\( (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + p^3 \)

\[ t = \text{simplify}(4); \]
\[ [0, 0, (1-p)^3, (1-p)^2p, 0, 0, (1-p)^2p, (1-p)p^2, 0, 0, (1-p)^2p, (1-p)p^2, 0, 0, (1-p)^2p, (1-p)p^2] \]
\[ 1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + p^3 = 1 \]  

So the expected halting time from \( s3 \) is the dot product of this vector with states plus 1.

\( v = \text{Vector}(16): v[3] = 1; \)

\[ s3 = 1 + \text{DotProduct}(4, \text{states}); \]
\[ s3 = 1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + s7 + (1-p)^2p^2 + s8 + (1-p)^2p^2 + s9 \]
\[ -p^3s1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + s11 + (1-p)^2p^2 + s12 + (1-p)^2p^2 + s16 + s15 + p^3s16 \]  

The equations defining the expected halting times from the sixteen states in Eqs. Equations 6 and 11 are exceptions.

\[ v = [1]; \]

\[ \text{for } i \text{ from } 1 \text{ to } 16 \text{ do} \]
\[ \text{v} = \text{Vector}(16): v[i] = 1; \]

\[ T. v = \text{states}; \]

\[ \text{Eq} = \text{dotProduct}(v, \text{states}); \]
\[ \text{Eq} = \text{operator}(\text{Eq}), \text{by} : \text{od}; \]

\[ \text{Eq}[6] = (s6 = 0); \]
\[ \text{Eq}[11] = (s11 = 0); \]

\[ \text{for } i = 1 \text{ to } 16 \]  
\[ \text{Eq}[i] = (1-p)^3p^2s + (1-p)^2p^2s + (1-p)p^2s + s7 + (1-p)p^2s + s8 + (1-p)^3p^2s \]
\[ -p^3s1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + s11 + (1-p)^2p^2 + s12 + (1-p)^2p^2 + s16 + s15 + p^3s16 \]
\[ -p^3s1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + s11 + (1-p)^2p^2 + s12 + (1-p)^2p^2 + s16 + s15 + p^3s16 \]

\[ \text{for } i = 1 \text{ to } 16 \]  
\[ \text{Eq}[i] = (1-p)^3p^2s + (1-p)^2p^2s + (1-p)p^2s + s7 + (1-p)p^2s + s8 + (1-p)^3p^2s \]
\[ -p^3s1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + s11 + (1-p)^2p^2 + s12 + (1-p)^2p^2 + s16 + s15 + p^3s16 \]
\[ -p^3s1 + (1-p)^3 + 3(1-p)^2p + 3(1-p)p^2 + s11 + (1-p)^2p^2 + s12 + (1-p)^2p^2 + s16 + s15 + p^3s16 \]

\[ \text{Now these equations can be solved.} \]
\[ \text{Sols} = \text{solve}(\text{Eq}, \text{convert}(\text{states}, \text{list}))[1]; \]
\[ \text{Sols} = \begin{cases} s7 = \frac{1}{2} & \frac{8p^2 - 20p^2 + 18p - 7}{8p^2 - 20p^2 + 18p - 7} \end{cases}, s2 = \begin{cases} 1 & 6p^3 - 15p^2 + 14p - 6 \end{cases}, s3 = \begin{cases} 1 & \frac{2(2p^3 - 7p^2 + 10p - 2)}{2p^3 - 7p^2 + 10p - 2} \end{cases}, s4 = \begin{cases} 1 & \frac{8p^3 - 20p^2 + 18p - 7}{8p^3 - 20p^2 + 18p - 7} \end{cases}, s5 = \begin{cases} 1 & \frac{2(2p^3 - 7p^2 + 10p - 2)}{2p^3 - 7p^2 + 10p - 2} \end{cases} \]
\begin{align*}
&-1 \quad \frac{6 p^3 - 15 p^2 + 14 p - 6}{2 (2 p^4 - 7 p^3 + 10 p^2 + 7 p - 2)} p^2, s_{6} = 0, s_{7} = 1 \\
&+1 \quad \frac{8 p^3 - 20 p^2 + 18 p - 7}{6 p^2 - 15 p^2 + 14 p - 2}, s_{8} = 1 \\
&+1 \quad \frac{6 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)}{6 p^2 - 15 p^2 + 14 p - 6}, s_{9} = 1 \\
&+1 \quad \frac{2 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)}{6 p^2 - 15 p^2 + 14 p - 6}, s_{10} = 1 \\
&+1 \quad \frac{2 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)}{6 p^2 - 15 p^2 + 14 p - 6}, s_{11} = 0, s_{12} = 1 \\
&+1 \quad \frac{6 p^3 - 15 p^2 + 14 p - 6}{2 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)} p^2, s_{13} = 1 \\
&+1 \quad \frac{8 p^3 - 20 p^2 + 18 p - 7}{6 p^2 - 15 p^2 + 14 p - 6}, s_{14} = 1 \\
&+1 \quad \frac{2 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)}{6 p^2 - 15 p^2 + 14 p - 6}, s_{15} = 1 \\
&+1 \quad \frac{2 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)}{6 p^2 - 15 p^2 + 14 p - 6}, s_{16} = 1 \\
&+1 \quad \frac{2 (2 p^4 - 7 p^3 + 10 p^2 - 7 p + 2)}{6 p^2 - 15 p^2 + 14 p - 6}, s_{17} = 1
\end{align*}

The initial state may be taken from states with a uniform probability.

\begin{verbatim}
> su := add(rhs(Sols[i]), i = 1..16);
\end{verbatim}

\begin{verbatim}
> Sols[1];
\end{verbatim}

\begin{verbatim}
> unapply(rhs(%, p));
S1 := %;
\end{verbatim}

\begin{verbatim}
> Sols[2];
\end{verbatim}

\begin{verbatim}
> unapply(rhs(%, p));
S2 := %;
\end{verbatim}

\begin{verbatim}
> Sols[6];
\end{verbatim}

\begin{verbatim}
> unapply(rhs(%, p));
S6 := %;
\end{verbatim}

\begin{verbatim}
> unapply(su, p);
Su := %;
\end{verbatim}

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\[ s_1 = -\frac{1}{2} \frac{8 p^2 - 20 p + 18 p - 7}{p^3 - (2 p - 7 p^2 + 10 p^2 - 7 p^4 + 2)} \]
\[ p = -\frac{1}{2} \frac{8 p^3 - 20 p^2 + 18 p - 7}{p^3 - (2 p - 7 p^2 + 10 p^2 - 7 p^4 + 2)} \]
\[ S_1 = p = -\frac{1}{2} \frac{8 p^3 - 20 p^2 + 18 p - 7}{p^3 - (2 p - 7 p^2 + 10 p^2 - 7 p^4 + 2)} \]
\[ S_2 = p = -\frac{1}{2} \frac{8 p^3 - 20 p^2 + 18 p - 7}{p^3 - (2 p - 7 p^2 + 10 p^2 - 7 p^4 + 2)} \]
\[ S_6 = p = 0 \]
\[ \text{To see the relationship between cell-activity-frequency and expected hailing time, functions are plotted for } 0.05 < p < 0.9. \]
\[ > \text{plot}([S_1(p), S_2(p), S_6(p), S_u(p)], p = .05..0.85, y = -1..35, \text{color = red, blue, green, plum}, \text{axes = boxed}); \]
Minimum expected halting times are computed using derivatives

\[ dS1 = D(S1); \]
\[ ds2 = D(s2); \]
\[ dsu = D(su); \]

\[ dS1 \approx p \rightarrow \frac{1}{2} \frac{24 p^2 - 40 p + 18}{p - \left(2 p^4 - 7 p^3 + 10 p^2 - 7 p - 2\right)} \]
\[ + \frac{1}{2} \frac{1}{p^2 \left(2 p^4 - 7 p^3 + 10 p^2 - 7 p - 2\right)^2} \]
\[ + \frac{1}{2} \frac{1}{\left(2 p^4 - 7 p^3 + 10 p^2 - 7 p - 2\right)^2} \]
\[ + \frac{1}{2} \frac{1}{p \left(2 p^4 - 7 p^3 + 10 p^2 - 7 p - 2\right)^2} \]

\[ ds2 \approx p \rightarrow \frac{1}{2} \frac{1}{18 p^2 - 30 p + 14} \]
\[ + \frac{1}{2} \frac{1}{18 p^2 - 30 p + 14} \]
\[ + \frac{1}{2} \frac{1}{6 p^2 - 15 p + 14} \]
\[ + \frac{1}{2} \frac{1}{6 p^2 - 15 p + 14} \]

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\[ dS_u = p_e \rightarrow - \frac{3}{16} \frac{24 p_e^2 - 40 p_e + 16}{p_e^5 (2 p_e^4 - 7 p_e^3 + 10 p_e^2 - 7 p_e + 2)} + \frac{3}{16} \frac{8 p_e^5 + 15 p_e^4 - 7}{p_e^3 (2 p_e^4 - 7 p_e^3 + 10 p_e^2 - 7 p_e + 2)} \]
\[ + \frac{3}{16} \frac{8 p_e^5 - 20 p_e^4 + 18 p_e^3 - 7}{p_e^3 (8 p_e^3 - 21 p_e^2 + 20 p_e - 7)} \]
\[ - \frac{1}{16} \frac{16 p_e^2 - 20 p_e + 14}{(2 p_e^4 - 7 p_e^3 + 10 p_e^2 - 7 p_e + 2) p_e} \]
\[ + \frac{1}{4} \frac{16 p_e^5 - 15 p_e^4 + 14 p_e - 6}{8 p_e^3 - 21 p_e^2 + 20 p_e - 7} \]
\[ + \frac{1}{4} \frac{2 p_e^5 - 7 p_e^4 + 10 p_e^3 - 7 p_e + 2}{p_e^3 (2 p_e^4 - 7 p_e^3 + 10 p_e^2 - 7 p_e + 2)} \]

\[ p_1 = \text{fsolve}(0 = dS_1(p), p = 0.2 \ldots 0.5) \]
\[ p_2 = \text{fsolve}(0 = dS_2(p), p = 0.2 \ldots 0.5) \]
\[ p_u = \text{fsolve}(0 = dS_u(p), p = 0.2 \ldots 0.5) \]

\[ S_1(p_1) \approx S_1(p_1) \]
\[ S_2(p_2) \approx S_2(p_2) \]
\[ S_u(p_u) \approx S_u(p_u) \]

\[ p_1 = 0.4209828778 \]
\[ p_2 = 0.5049132591 \]
\[ p_u = 0.406932440 \]
\[ S_1(p_1) = 7.999997770 \]
\[ S_2(p_2) = 7.488842975 \]
\[ S_u(p_u) = 6.675843786 \]