Kinetic Origin of Heredity in a Replicating System with a Catalytic Network

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Abstract

The origin of heredity is studied as a recursive state in a replicating proto-cell consisting of many molecule species in mutually catalyzing reaction networks. Protocells divide when the number of molecules, increasing due to replication, exceeds a certain threshold. We study how the chemicals in a catalytic network can form recursive production states in the presence of errors in the replication process. Depending on the balance between the total number of molecules in a cell and the number of molecule species, we have found three phases; a phase without a recursive production state, a phase with itinerancy over a few recursive states, and a phase with fixed recursive production states. Heredity is realized in the latter two phases where molecule species that are population-wise in the minority are preserved and control the phenotype of the cell. It is shown that evolvability is realized in the itinerancy phase, where a change in the number of minority molecules controls a change of the chemical state.

Key Words: catalytic network, heredity, minority control, origin of life, evolvability
1 Introduction

In recent studies on isologous diversification\cite{1, 2, 3}, we have shown that cell differentiation and developmental processes are general characteristics of systems of interacting and replicating cells that contain mutually catalytic molecules. In such systems, a cell consists of several chemicals that catalyze each other. When the chemicals in a cell exceed a given threshold, the cell divides. Even though detailed mechanisms are not programmed in advance, differentiation of cell types, irreversible processes from multipotent stem cells to commitment, and robust development can be found.

Then one may ask what the role of genes is. As a possible answer to this question, we conjectured that genes come into action to fixate the differentiation provided by the interaction of units with autocatalytic reaction networks. The idea is that some of the chemicals of the network provide a hereditary starting point that plays a role in controlling the phenotype of the cell. Indeed, a related idea was already proposed in the context of evolution, by Waddington\cite{4} and Newman\cite{5}.

Since a gene is a part of the larger structure that is DNA, its action should be represented merely by a component of the catalytic reactions in the chemical network. How is it possible then that some chemicals are able to control cell differentiation?

To answer this question, we have recently proposed a minority-control theory\cite{6}. In a system of mutually catalyzing molecules, a state is selected through reproduction where a minor molecule species starts to control the chemical state of the cell. This state is preserved over many generations of cells. In this chemical state termed the “minority controlled state”, a separation of roles between two sets of molecule species appears. One set has large numbers of molecules, maintaining the diversity of the species, while the other has small numbers of molecules but a larger catalytic activity. The latter set has the following two properties;

[Preservation property]: The molecule species are preserved well over generations. The number of molecules per species exhibits smaller fluctuations compared to other molecule species, and their chemical structure (such as polymer sequence) is preserved over a long time span, even under potential changes caused by thermodynamic fluctuations during the synthesis of these molecules.

[Control property]: A structural change in the molecule species or a change in the number of the molecules has a stronger effect on the behavior of a cell, for example on the growth rate of the cell.
These two properties form the basis for heredity, causing statistical correlations between the phenotypes of ancestor and offspring. Due to the control property, the minority molecules govern the phenotype, and due to the preservation property, they can transfer the information relevant for determining the phenotype to future generations. Indeed, once this minority controlled state is established, a new evolutionary incentive emerges for embedding the representation of heredity into the minority molecule. Thus the emergence of genetic information from the minority-controlled state is expected.

In this earlier study, we considered only two molecule species with different synthesis speeds, one being much higher than the other. Or, equivalently, the catalytic activity of the latter is much higher than the former due to the mutual catalyzation for the synthesis. The minority controlled state is a consequence of this difference in the synthesis speeds. Then, the question remains whether it is a generic property of chemical reaction networks to select states with large differences in synthesis speeds resulting in recursive production controlled by a few minority molecule species. Furthermore, it is necessary to clarify how specific chemical compositions are stabilized as recursive states through control by minority molecules. Here it is interesting to note that cell differentiation in a model with interacting cells\textsuperscript{[1]} was initiated by difference in the concentration of minority molecules by cells.

To answer this question, we consider a protocell system consisting of a large number of molecule species that catalyze each other. By carrying out stochastic simulations of such catalytic reaction networks, we will show in this paper that a biochemical state with minority molecules can establish recursive production. In other words, we will show that the dynamic change of the chemical composition is stabilized by minority molecules that catalyze majority molecules, and that the minority molecule species carries this information through reproduction. Furthermore, we will also show that this minority controlled state has evolvability, since by only a slight change in the number of the minority molecules, the total chemical composition is drastically altered.

2 Model

To study the general features of a system with mutually catalyzing molecules\textsuperscript{[1]}, we consider the following simple model. First, we envision a (proto)cell containing \( k \) molecule species. (Not all of these species are necessarily present.
Some of the molecule species may have zero population. With a supply of chemicals available to the cell, the molecules replicate through catalytic reactions, so that their numbers within a cell increase. When the total number of molecules exceeds a given threshold (here we used $2N$), the cell divides into two, with each daughter cell inheriting half of the molecules of the mother cell, chosen randomly. Each chemical species catalyzes the synthesis of some other randomly chosen chemical as

$$X^i + X^j \rightarrow 2X^i + X^j.$$  \hspace{1cm} (1)

with $i, j = 1, \cdots, k$. The connection rate of the catalytic path is given by $p$ (which will be taken to be 0.2 or 0.1 for most simulations here), and the connection is chosen randomly. (Here we investigated the case without direct mutual connections, i.e., $i \rightarrow j$ was excluded as a possibility when there was a path $j \rightarrow i$, although this condition is not essential for the results to be discussed). Each molecule species $i$ has a given catalytic ability $c(i)$ and its own synthesis speed $g(i)$. Accordingly, the above reaction occurs with the rate $g(i)c(j)$.

During the replication process structural changes may occur that alter the activity of the molecules. Therefore the replicated molecule species can differ from that of the mother. The rate of such structural changes is given by $\mu$, which may not be very small due to thermodynamic fluctuations. This change can consist of the alternation of a sequence in a polymer or other conformational change, and may be regarded as a replication ‘error’. Here, for the simplest case, we take this ‘error’ to affect all molecule species equally, (i.e., with the rate $\mu/(k - 1)$), and could thus regard it as a background fluctuation. Generally in reality of course, even after a structural change, the replicated molecule would keep some similarity with the original molecule, and the replicated species with the ‘error’ would be within a limited class of molecule species. Later we will discuss this case also, but the basic conclusion will not be altered. Hence we use the simplest case for most simulations.

We simulated this model according to the following procedure. In the beginning, we choose connection matrix randomly. Unless otherwise mentioned, the parameters $c(i)$ and $g(i)$ are given from a random number over $[0, 0.5]$, and they are fixed through each simulation. At each step, a pair of molecules, whose species is assumed to be $i$ and $j$, is chosen randomly. If there is a connection between the species $i$ and $j$, then according to the reaction (1), one molecule of the species $i$ is added with the probability
$g(i)c(j)$ if $j$ catalyzes the synthesis of $i$, or $j$ is added with the probability $g(j)c(i)$ if $i$ catalyzes the synthesis of $j$. The replications are subjected to errors with the rate $\mu$ given above. The error rate $\mu$ is fixed at 0.01 through the simulations here, but this specific choice is not so important as long as it is not too large. Since our model deals with rather small numbers of molecules, a stochastic approach was chosen for the simulations rather than the usual ordinary differential equations for chemical concentrations which cannot describe some novel effects [7].

When the number of molecules within a cell is larger than $2N$, it is divided into two, and a new cell is created. The total number of cells, $M_{tot}$, is kept constant so that one protocell, randomly chosen, is removed whenever a (different) protocell divides into two. Often we use $M_{tot} = 1$ here, where only one of the daughter cells remains. In this case, our model is quite similar to a stochastic simulation of the population dynamics of $k$ molecule species, but at each division event large fluctuations arise. Hence a recursive state that continues production has to be selected, resulting in a choice of special initial conditions for the chemical composition. This selection of initial conditions through repetition of divisions is important, especially when $N$ is not large (or $k$ is large). This selection of initial conditions distinguishes our model from standard population dynamics of molecules. With this model we address the question how recursive production is achieved, and how evolutionary change of the recursive state is possible.

3 Recursive States and Itinerancy

In our model there are four basic parameters; the total number of molecules $N$, the total number of molecule species $k$, the mutation rate $\mu$, and the path rate $p$, besides the total cell number $M_{tot}$, the catalytic activity $c(i)$ and growth rate $g(i)$. By investigating many sets of parameters and also by choosing various random networks, we have found that there are roughly three types of behaviors:

1. Fast switching states without recursiveness
2. Itinerancy over several recursive states
3. Achievement of fixed recursive states

In the first phase, there is no clear recursive production and the dominant molecule species changes frequently. At one time step, some chemicals may be dominant but only a few generations later, this information is lost, and the number of the molecules in this species goes to zero. Due to the
autocatalytic nature, the population of one species can be amplified, only to be replaced by another population catalyzed by it. No stable set of catalytic networks is formed that excludes other ‘parasitic’ molecules. (See for example the dynamics of chemicals around the last stage of Fig.1a). In phase (1), this type of dynamics continues forever, without showing any quiescent state as around the middle of Fig.1a).

In the second phase, alternately, recursive states which last over many generations (typically a thousand generations) and fast switching states appear (See for example the dynamics of chemicals around the last stage of Fig.1a).

In the third phase, on the other hand, a recursive state is established, and the chemical composition is stabilized such that it is not altered by the division process. Once reached, this state is permanent and the system remains in it for as long the simulation lasts. (see Fig.1b).

To be precise, the recursive state observed in phases (2) and (3) is not necessarily a fixed point as with regards to the population dynamics of the chemical concentrations. In some cases, as shown in Fig.1b), the chemical concentrations oscillate in time, but the nature of the oscillation is not altered by the process of cell division. In dynamical systems terms, the recursive state here is an attractor, and this attractor can be a limit cycle (or could be chaos or some other low-dimensional attractors), even though often it actually is a fixed point attractor. (In all of these cases there are fluctuations due to the small number of molecules, of course). Generally, all the observed recursive states consist of 5-10 species, except for those species which exist only as a result of replication errors where the number of molecules is one or two.

Which of the states (1), (2), (3) appears depends, of course, on the parameters and the specific choice of the network. First, we discuss how the phase changes when varying $N$ and $k$ if $c(i)$ and $g(i)$ are distributed homogeneously over $[0,0.5]$, $M_{tot} = 1$, and $p = 0.2$, since we have not yet elucidated any clear dependence on $p$ and $M_{tot}$ so far.

Although the behavior of the system depends on the choice of the network, there is a general trend with regards to the phase change, from (1), to (2), and then to (3) with the increase of $N$, or with the decrease of $k$, as is schematically drawn in Fig.2.

We now study what role the catalytic activity plays in maintaining the recursive states. In Fig.3, we have plotted the population of each molecule

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1 Such switching between several recursive states is also studied as chaotic itinerancy
Figure 1: The number of molecules $N(i)$ for the species $i$ is plotted at each successive division event. In both (a) and (b), the same random network with $k=200$ and $p=.2$ was used while $N=2000$ in (a), and $N=6000$ in (b). Only the species with $N(i)>10$ are plotted at each time. The straight lines are artifacts of the drawing occurring when $N(i)$ dips below 10 and then at a later time increases to above 10 again. The parameters $c(i)$ and $g(i)$ are randomly distributed over $[0,0.5]$ and are fixed through the simulation. In (a), there appears a recursive state around $45.8 \times 10^3$-th generation, and another recursive state appears around $47 \times 10^3$-th generation and repeats collapse and reappearance up to $48.3 \times 10^3$-th generation. In (b), a single recursive state lasts through the whole simulation, where 5 species (71, 173, 3, 54, 99 in the order of population size) continue to exist.
species as a function of the catalytic activity. For phase (1), no clear structure is discernible, although a slight tendency for the average population to decrease with an increase in the catalytic activity exists as is shown in Fig. 3a). The maximum of each population, however, depends only minimally on the catalytic activity, implying that almost all species can be the dominant species at some time.

For phase (2), a clear structure can be observed and the maximum populations decrease for increasing catalytic activity. Indeed, this negative correlation between catalytic activity and population is not surprising. If the catalytic activity is higher it can help the synthesis of other molecules. Since the total number of molecules is limited, the fraction of that molecule population itself should be less.

This negative correlation is amplified and fixed in phase (3), as shown in Fig. 3b). Here, only a few molecule species with small populations and high catalytic activity survive. Although these molecules are in the minority population-wise, their existence is essential for maintaining this recursive structure. The major species have lower catalytic activities, and accordingly they are not catalyzed efficiently by the other major molecule species. Indeed, the species with the majority of the population are catalyzed by the minority species. The latter is catalyzed by species with larger populations but with smaller catalytic activities. The surviving molecule species form
Figure 3: Average(∗), maximal(+) and minimal(×) populations of the species $i$, plotted as a function of their catalytic activity $c(i)$. The average, maximum and minimum are taken over the 50000-100000th division. In (b), the five dominant species are discernible, which are species 71, 173, 3, 54, 99 in the order of population size (i.e., in the reverse order of $c(i)$).
a mutually catalyzing reaction network validating the assumptions made in
the minority control theory.

For example, in the recursive state in Fig.1b), there are 5 species whose
population remains in existence for all the time. The network structure of
these molecule species is shown in Fig.4, with other molecules that exist not
over all the time but over relatively long time span. The catalytic activities
of the key species are $c(99) = .48 > c(54) = .39 > c(71) = .063$, while
the respective populations are $N(71) \gg N(54) \gg N(99)$. The recursive
state here is achieved by catalysis of the species 99 whose population is a
minority in the network. When the total population is much smaller, the
species 99 may decrease due to fluctuations and competition with other
molecules catalyzed by the species 71. For example, in the collapse of the
recursive state in Fig.2a), the population of the species 3 increases due
to catalyzation by 71, and a state dominated by 71, 54, and 3 is formed.
Since the catalytic activity of these molecules is not so high, the state is
dynamically unstable and allows for an increase in the populations of other
molecules. For example, the increase of the species 61 leads to the increase
of species 18 and 137 $\cdots$, successively, and the recursive state is replaced by
the fast switching state.

By examining several reaction networks, we came to understand itiner-
cy and the stability of recursive states as follows. As for the recursive
state, one might wonder why the species with higher catalytic activities are
not taken over by parasitic molecule species that have lower catalytic ac-
tivities and are catalyzed by molecules with higher catalytic activities. To
sustain a recursive state, the emergence of such parasitic molecule species
should be suppressed. In our model, molecules with higher catalytic ac-
tivities are catalyzed by a molecule species with lower activities but larger
populations. Hence, the parasitic molecule species cannot easily invade to
disrupt the mutually catalytic core network. Since the minority molecule
(say 99 in the above example) and majority molecule (71 in the above ex-
ample) form a mutual catalytic network (with the aid of another molecule
(54)), a large fluctuation is required to destroy this mutually supporting
network.

For recursive stability, it is important that there are two levels of repli-
cation, i.e., molecular replication and cellular replication[10]. As parasitic
molecules with lower catalytic activity appear, recursiveness is lost, and the
state starts to change, while the total growth speed of the cell generally
decreases. With cellular replication, the selection of initial conditions main-
taining the recursive state is thus favored. In the minority controlled state,
minority and majority molecules catalyze and reinforce each other, and the recursive state is stabilized.

This allows us to explain how the itinerancy over (partially) recursive states appears in phase (2). When the number of molecules is not so large, the numbers of some molecule species that are not in the original core network start to increase. Since the total number of molecules in a cell is limited, the minority molecules may then decrease in number, and, if the total number of molecules is small enough, even become extinct. If that happens, the molecule species with a large population loses the main source that catalyzes it. Hence several molecules mostly of lower catalytic activity start to compete for growth. Consequently, the diversity of molecule species increases, a stable recursive production cycle of cells is not established, and the dominant species change frequently. After the system has been in a fast switching state for a while, another (or possibly the same) core network structure is formed where the existence of a minority molecule species with a higher catalytic activity stabilizes the recursive state.

With the above considerations is it quite reasonable to expect the transition from phase (2) to (3) for increasing $N$. After all, for larger $N$, the number of molecules in the minority species also increases thus protecting them from the extinction. However, the transition from (2) to (3) also strongly depends on the network structure. For some reaction networks, a simple core structure is easily formed and the recursive state is maintained even for small $N$. For some other networks, the itinerancy of phase (2) is observed for large $N$. Although it is not so easy to distinguish these two cases by just examining the network structure, some differences seem to exist in the dynamics of the recursive state. If the state falls on a fixed point, recursiveness is stabilized for smaller $N$. On the other hand, when the population shows oscillatory behavior, dynamic instability is generated,
Figure 5: An example of an overlaid time series for the case with a biased distribution of $c(i)$. $N = 2000$ and $k = 200$.

and fluctuations in the species with small molecule numbers are continuously amplified. Hence the state is subjected to larger overall perturbations and stabilized only at larger $N$.

Here the distribution of catalytic activity is important. We studied the cases that either $c(i)$ or $g(i)$ or both are homogeneous. When $c(i)$ is constant, the itinerancy state of phase (2) is rarely observed. The itinerancy state is common, however, when the distribution of $c(i)$ is biased to having more species with lower catalytic activity. For example, we carried out simulations with log $c(i)$ homogeneously distributed. In this case, the itinerancy over recursive states is a common occurrence as, for example, can be seen in Fig.5. Considering that enzymes with higher catalytic activity are rare, this choice of distribution of $c(i)$ with a bias towards a zero value is rather natural. Hence, the itinerancy over recursive states should be rather common in general.
4 Discussion: Remark on Evolvability

An important consequence of the above itinerancy state is evolvability (see also [3]). With a change in the number of minority molecules, or by a structural change in one of them, the chemical composition of a cell can change drastically and establish a novel recursive state. To investigate this evolvability, we have modified the mutation conditions such that one molecule species can only mutate into a limited range of other molecule species, i.e., a species \( i \) can only mutate to a species in the range \([i-m, i+m]\) with \( m < k/2 \). Starting from an initial condition consisting only of species within a range \([k_1, k_2]\), one can then examine whether the number of molecule species expands to the whole range \([0, k]\). We found that in phase (2), the evolutionary process ‘scans’ over all the molecule species (i.e. \( i \in [0, k] \)), by itinerating over several recursive states. This is a typical feature of the second phase. For phase (3), the chemical state is stuck in the recursive state formed at an early stage, while in phase (1), no recursive state exists.

A recursive state in a mutually catalytic system was also discussed by Lancet [13] as a ‘compositional genome’. Recursive states are furthermore similar to the determined states in the cell differentiation model [1, 3] where the chemical composition of a cell is transferred to its offspring. There, stable states are understood as attractors or as partial attractors stabilized by the interaction. The recursive state resulting from such a partial attractor is stabilized by a single molecule species in the current model.

This recursive state forms a few discrete states among a huge variety of possible chemical compositions, as an attractor. This state gives a basis for heredity. In genomic information, on the other hand, each chemical state of a cell is represented by a single molecule. To bridge the gap between the attractor viewpoint and information viewpoint, each attractor state has to be represented by a single molecule and has to be stabilized. This bridge is provided by the minority controlled theory we proposed. Besides recursiveness, evolvability is an important consequence of the minority controlled state.

The minority-controlled state will generally be important for understanding the ‘bottle-neck’ phenomena ubiquitous in biological systems. Often a process in a cell includes some checkpoints, and each checkpoint has to be passed for a cellular process to continue to the next stage. For example,
consider the timing of a cell division. To start the division, a cell has to complete several processes. Then the slowest dynamics involved in the cell division process forms a bottleneck. With such a bottleneck, a recursive cellular state is guaranteed. (For example, most of the gadgets in the cell, even though some of them require a longer time for synthesis, are doubled before the division). With the minority-controlled state, a dynamics to overcome the bottleneck is provided. Since the minority molecule is necessary for the catalysis of many other molecules, the cell has to wait for it to be replicated before it proceeds to the next stage. If many minority molecules are combined together as a single long polymer sequence, then this polymer will act as the control of all the other cell processes. Probably this is the role of DNA, and the origin of this fascinating information carrier.

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