Networks of catalyzed reactions with nonlinear feedback have been proposed to play an important role in the origin of life. We investigate this possibility in a polymer chemistry with catalyzed cleavage and condensation reactions, studying the properties of a well-stirred reactor driven away from equilibrium by the flow of mass. Near equilibrium, the distribution of material is uninteresting; it favors short polymers but is otherwise homogeneous. However, under appropriate non-equilibrium conditions, the situation changes radically: The nonlinear feedback of the reaction network focuses the material of the system into a few specific polymer species, whose concentrations can be orders of magnitude above the background. Like a metabolism, the network of catalytic reactions “digests” the material of its environment, incorporating it into its own form. For this reason we call it an *autocatalytic metabolism*. We vary the diet of an autocatalytic metabolism, and demonstrate that under some variations it persists almost unchanged, while in other cases it dies. We argue that the dynamical stability of autocatalytic metabolisms gives them regenerative properties that allow them to repair themselves and to propagate through time.
1. MOTIVATION

1.1 SETTING THE STAGE FOR AN ORIGIN OF LIFE

When Miller and Urey discovered that amino acids could be formed under conditions that might be similar to those of the prebiotic earth, the spontaneous synthesis of proteins seemed just around the corner. However, this turns out to be much more difficult than the spontaneous synthesis of individual amino acids. Equilibrium conditions tend to favor dissociation, and generate a concentration profile that is fairly uniform. Except for occasional fluctuations, for long polymers the population of any given molecular species is typically zero. The population distribution of polymers is homogeneous, nonspecific, and uninteresting. This is in contrast to living organisms, which have high concentrations of a few specific polymer species.\[1\]

Contemporary organisms achieve specificity through a codependent relationship between templates and enzymes. Proteins and nucleic acids synthesize each other through a replication mechanism in which none of the components synthesizes itself. Even for the simplest organisms, this process is highly complex. There seems to be a minimum level of complexity below which a replicating machine based on proteins and nucleic acids simply cannot function. While it is easy to understand how such a replicating machine perpetuates itself, it is difficult to understand how the necessary initial conditions ever arose on their own. The probability that both enzymes and templates could be created through a statistical fluctuation is effectively nil. This suggests that other processes preceded contemporary life.

The idea that enzymatic activity might have set the stage for the origin of life was developed by Oparin, who suggested that conserves may have played a major role. Early experiments unsuccessfully attempted to use clay and other materials as nonspecific catalysts for polymerization. Calvis studied several different scenarios through which catalytic activity could provide a selection mechanism, even without self-replication. In 1971 Rössler and Kaufman developed this idea further. In particular, Rössler envisioned a form of chemical evolution similar to that studied here. He emphasized the importance of specific catalysts which catalyze only a small fraction of all possible reactions. Along these same lines, Kaufman later modeled the problem in terms of random graphs, and showed that under reasonable assumptions the probability of catalytic closure is quite high. The random graph model was developed into a kinetic model that could be simulated on a computer by Farmer et al. This line of investigation, which attempts to find possible precursors facilitating the emergence of life should be contrasted with

An exception is provided by the experiments of Sidney Fox, who by heating a mixture of amino acids demonstrated the formation of polypeptides, called proteinoids.\[2\] The structure of proteinoids is not random; some subsequences, such as certain hexapeptides, occur much more frequently than others. In contrast, our goal is to increase the concentration of entire molecules, so that it is several orders of magnitude above equilibrium.

\[1\] Another toy model investigating the possibility that a metabolism might have spontaneously emerged without a replicator is due to Dyson.\[8\]
other work that addresses the (also very interesting) question of the early evolution of life once replication has already begun.\textsuperscript{1,5,10,39}

In this paper we study the behavior of a network of catalyzed chemical reactions, along the lines laid out by Farmer et al.\textsuperscript{11} We make several enhancements of the model and analytically study a few simple cases to gain better intuition about the dynamics. We also improve the simulation so that it is several orders of magnitude faster. This allows us to simulate the kinetics of a complicated reaction network in a matter of seconds. As a result, we are able to widely explore the parameter space and answer many of the questions originally raised in earlier papers.

Our main result is that, under appropriate conditions, a catalytic reaction network can focus most of the material of its environment into a few chemical species. For this to happen the system must be driven by the appropriate distance from equilibrium, polymerization must be favored, and it must have diverse kinetic parameters. Favoring polymerization may require the addition of energy, for example, through pyrophosphates energized by light. In spite of these restrictions, there is a wide range of parameters in which the material of the system is focused into only a few species, which dominate over the background.

Focusing radically alters the material composition of the environment. The species that emerge reinforce each others' production and largely take over the reaction vessel, excluding other possibilities. Since this behavior is analogous to that of a metabolism, we call the resulting set of species and reactions an \textit{autocatalytic metabolism}. Under appropriate conditions autocatalytic metabolisms can evolve out of a simple, undifferentiated initial state, generating a sequence of complex, highly differentiated, final states. Like contemporary organisms, these final states are composed of a highly focused, specific set of long polymers. While the autocatalytic metabolism does not replicate itself in the usual sense, it propagates itself by taking over any medium with suitable properties, sustaining itself as long as the appropriate conditions are met. Furthermore, it may generate a lineage of related autocatalytic metabolisms.

The model that we study here applies to any system in which polymers can catalyze the formation of other polymers through cleavage and condensation reactions. If the basic building blocks are amino acids, then the polymers are called polypeptides or proteins. Such reactions are common among proteins, forming the basis for many of the functions of living organisms. If the basic building blocks are nucleic acids, then the polymers are called nucleic acids (RNA or DNA). It is well known that polypeptides possess a large repertoire of catalytic activities of this type; the recent discovery of specific catalytic reactions in RNA suggests that nucleic acids may also possess the necessary properties.\textsuperscript{7} Whether the polymers are polypeptides or nucleic acids changes the parameters but not the basic form of the model.

Even if the model we discuss here has nothing to do with the actual origin of life on earth, it might provide a possible origin of life in the laboratory. Although at present we cannot predict the outcome of experiments in detail, we can make qualitative predictions that provide broader experimental guidelines. The accumulation of more experimental knowledge can be used to determine the unknown parameters of our model, which in turn should sharpen its predictive value for experiments. Although many important experimental details are still unknown, and some important questions await further study, our numerical simulations suggest that it may be possible to synthesize autocatalytic metabolisms in the laboratory.

### 1.2 NONTRIVIAL DISSIPATIVE STRUCTURES

The problem of the emergence of life is embedded in the broader problem of understanding self-organizing phenomena, which from the point of view of a physicist may be more interesting anyway. Many non-living systems exhibit self-organizing properties, albeit much weaker than those of living systems. Is there a sharp distinction between living and nonliving systems? Or can there exist levels of organization that are between those of present living and non-living systems? Can evolution and other self-organizing properties of living systems be viewed as manifestations of a general law that describes the tendencies of matter to organize itself?

There are many simple examples that have been cited as instances of self-organizing phenomena in nature. For example, when a fluid is heated from below, under appropriate conditions, patterns of convecting cells form. The macroscopic structure of these cells is internally generated by the system itself, and is not apparent in its initial conditions. Such patterns are often called \textit{dissipative structures}, because they occur when energy flows into a system and then is dissipated.\textsuperscript{32}

Several researchers have asserted that life is a non-equilibrium phenomenon, associated with dissipative structures.\textsuperscript{32} This is certainly true, but it is a very weak statement. While deviation from equilibrium is a necessary condition for life, it is far from sufficient. Driving a system away from equilibrium does not necessarily cause the emergence of order—in fact, it often has precisely the opposite effect. A central question that must be addressed in a theory of self-organizing phenomena is: Why do some non-equilibrium situations foster the spontaneous emergence of organization, while others do not?

There is a big gap between the dissipative structures of simple non-living systems, such as patterns in fluid convection, and the much richer dissipative structures associated with living systems. The model discussed here is intended to bridge this gap, at least to some extent, by showing the possibility for dissipative structures that are intermediate in complexity between living and non-living systems. Autocatalytic metabolisms are more complex than convection patterns, in that they propagate specific information through time. One autocatalytic metabolism can seed the formation of another, similar metabolism. The autocatalytic metabolisms of this model can be viewed as proto-life forms, since they have a metabolism, they evolve and store information, and they reproduce (although more continuously and with less fidelity than contemporary organisms). They are also dynamically stable, and so capable of self-repair. They, thus, have many of the essential properties of living systems, albeit in a much less sophisticated form.

Besides demonstrating the possibility for the spontaneous generation of autocatalytic metabolisms, one of our main purposes in this paper is to discover under
what conditions they can be expected to form. How does their formation depend on the parameters of the system, such as the flow of energy, or the inherent diversity of the underlying dynamics? Although our results are specific to this model, they nonetheless suggest several rules that may pertain to the more general problem of self-organization.

1.3 A SIMPLE MODEL FOR STUDYING EVOLUTION WITH AN EMERGENT NOTION OF FITNESS

In principle, it is possible to describe biological systems at a fundamental level in terms of their dynamics. At this level of description, "selection" is an emergent property of the dynamics. In practice, however, for most systems this is hopelessly intractable. As a result, studies of evolution are typically couched in terms of the fitness function, which is an empirical construct, disconnected from the laws of physics. Even so, in most systems the fitness function is known only in very special circumstances where all but a few relevant factors are neglected. In general, fitness is a complicated function of the external environment, which includes other organisms. As a result, most theoretical models for evolution make many ad hoc assumptions, postulating fitness functions that may be qualitatively different from those in the real world.

As pointed out by Eigen, Rössler, and others, chemistry provides an excellent forum for studying evolution. The laws of chemical kinetics are well understood, and make it possible to model the behavior of the system at a fundamental level. These laws determine population levels and therefore determine fitness. As in biological systems, fluctuations are always present, generating random variation. Thus, for chemical networks we can describe the fitness at the fundamental level of dynamics.

Even though autocatalytic metabolisms do not have templates or a genetic code, because of their specificity they are nonetheless capable of evolution. This is discussed in a companion paper. Autocatalytic metabolisms, therefore, provide an interesting alternative for studying evolution in a chemical setting. It is also interesting to note that autocatalytic structures analogous to those we study here occur spontaneously in more abstract environments, as observed by Fontana and Rasmussen et al.

2. BACKGROUND

In this section we discuss some of the properties of catalyzed reaction networks, providing a background for the development of the simulation in Section 3. We discuss the reactions we are going to consider, and show why they are uninteresting at equilibrium. We then explain how the situation is altered as we move away from equilibrium, and how catalysis can play an important role in focusing the material of the system into just a few chemical species. We define autocatalytic sets and the related notion of autocatalytic metabolisms.

2.1 SPONTANEOUS REACTIONS

We are interested in irreversible polymerization reactions, in which two polymers either condense to form a single longer polymer, or a single polymer cleaves into two shorter polymers. Cleavage and condensation can be considered together as a single reversible reaction. The reaction in which polymers A and B join together to form C, giving off water, or equivalently, in which C hydrolyzes into A and B, can be written

$$A + B \rightarrow C + H,$$

where H represents water.

Providing the concentrations are sufficiently high and the solution is well stirred, the law of mass action provides a good approximation of the kinetics. Let $k_f$ be the rate constant for the forward reaction, $A + B \rightarrow C + H$, and $k_c$ be the rate constant for the backward reaction $C + H \rightarrow A + B$. The rate equation for C is then

$$\dot{C} = \frac{dC}{dt} = k_f AB - k_c CH.$$

For convenience, whenever the meaning is unambiguous, we use the same symbol to represent both a polymer and its concentration. Similar equations apply for A and B.

2.2 EQUILIBRIUM DISTRIBUTION OF POLYMERS

At equilibrium the concentrations of the polymers of a given length can be computed analytically using the classical theory of polycondensation reactions developed by Flory and Stockmayer. For simplicity we assume that all the reactions have the same forward and backward rate constants, and that the reaction vessel is well stirred. Furthermore, we assume that the monomers are oriented so that each monomer has two sites, which we arbitrarily designate as the "+" site and "-" site.

We follow the treatment of Macken and Perelson. Rather than solving for the concentrations of each polymer, it is more convenient to use an aggregate variable $y$, which is the concentration of free sites of a given kind (either "+" or "-"). We assume that the polymers are unbranched, and that they cannot form rings. For reactions of the form of Eq. (2),

$$\dot{y} = -k_f y^2 + k_c (m_0 - y) H,$$

where $m_0$ is the number of monomers available.
where \( n_0 \) is the total concentration of monomers, which is equal to the concentration of free sites if nothing is bound. At steady state \( \dot{y} = 0 \) and the concentration of free sites is

\[
y = \frac{(1 + 4\kappa y_0)^{1/2} - 1}{2\kappa},
\]

where \( \kappa = k_f / H k_r \) is the equilibrium constant.

We now compute the concentration of polymers of length \( n \). At equilibrium, let \( \rho \) be the probability for the formation of a bond. This is the ratio of bound sites to the total number of sites, i.e.,

\[
\rho = \frac{m_0 - y}{y_0}.
\]

Assume that each binding event is independent. For a given free site, the probability that it is attached to \( n - 1 \) bonds followed by another free site is \( \rho^{n-1}(1 - \rho) \). Solving Eq. (5) for \( y \) shows that the concentration of free sites for a given value of \( \rho \) is \( y = m_0(1 - \rho) \). Thus, the concentration of polymers of length \( n \) is

\[
x_n = m_0(1 - \rho)^2 \rho^{n-1}.
\]

At equilibrium, inserting Eq. (4) into Eq. (3) gives

\[
\rho = 1 - \frac{(1 + 4\kappa m_0)^{1/2} - 1}{2\kappa m_0}.
\]

Note that \( \rho < 1 \). Thus, Eq. (6) implies that the concentration of polymers of length \( n \) decreases exponentially with \( n \), at a rate that depends only on the product of the equilibrium constant and the concentration of monomers. In a system with \( m \) distinct monomers, present initially at equal concentrations, the concentration of any particular polymer species of length \( n \) is further decreased by a factor of \( m^{-n} \). For \( m > 1 \), even if \( \rho \approx 1 \), so that polymerization is favored, for large \( n \) the concentration of any particular species is quite small. For example, for polypeptides \( m = 20 \); the concentration of a polypeptide of length \( n = 30 \) is roughly \( 20^{-30} \) less than that of a monomer. For a container of finite size, this implies that, except for occasional fluctuations, most longer species are not present.

2.3 CATALYZED REACTIONS

The presence of a catalyst (enzyme) \( E \) can accelerate a reaction.

\[
A + B \xrightarrow{E} C + H.
\]

At equilibrium the rate of the forward reaction equals that of the backward reaction, so that \( A = \dot{B} = C = 0 \). Catalysis speeds up the rate at which the system approaches equilibrium, but does not change the concentrations at equilibrium.

However, when the reaction is driven away from equilibrium, for example by externally supplying one of the participants in the reaction, catalysis can shift the steady state. This is the basis for the effect we study here.

Catalysis increases both the forward and backward rate constants by the same amount. This can be taken into account by defining a quantity \( \nu \) that we call the catalytic efficiency. For fixed concentration of the reactants, the increase in the velocity of the reaction is proportional to the product of \( \nu \) and the concentration \( C \) of the catalyst. The kinetic equation for \( C \) can be crudely approximated as

\[
\dot{C} = (1 + \nu E)(k_f AB - k_r HC).
\]

Similar equations apply to \( \dot{A} \) and \( \dot{B} \). When the catalytic efficiency \( \nu = 0 \), this reduces to the kinetic equation for a spontaneous reaction.

Note that, for a population of polymers, this reaction is just one reaction in a network of many. A given polymer may play the role of \( A \) in some reactions, and the role of \( C \) in others. To compute the rate of production of any given species, it is necessary to sum all the relevant reaction terms.

The approximation made in Eq. (8) neglects the effect of saturation, which comes about because the enzyme and the reactants are bound together for a finite time. During this time they cannot participate in new reactions, which lowers the effective reaction rates. If this is a dominant effect, so that most of the enzyme or product is bound at any given time, the reaction is saturated. To take this into account we do not use Eq. (9), but rather use a more accurate approximation. We keep track of the concentration of any given species \( x_i \) that is bound into complexes through an auxiliary variable \( x_i \), which is equal to the sum of the concentrations of all the complexes in which \( x_i \) is bound. To keep the simulation tractable, we assume that all complexes unbind at the same rate \( k_u \). This approximation is described in more detail in the Appendix.

2.4 DRIVING FROM EQUILIBRIUM

To make anything interesting happen in a reaction network it must be driven away from equilibrium. In this model we investigate two different mechanisms. The first involves a flux of mass, and the second involves the formation of energetic pyrophosphate molecules, driven by light.

2.4.1 MASS FLOW We model a reaction vessel with a steady input flux of monomers or short polymers, and an output flux due either to diffusion or overflow of the reaction vessel. This might correspond to a prebiotic environment, or it might correspond to a chemostat in a laboratory experiment. The chemical species that are input are collectively called the food set. For simplicity, we assume an inflow rate \( \delta \) of concentration per unit time, and an outflow that is proportional to concentration, with rate constant \( K \).
For an element of the food set, the kinetic equations are of the form

\[
\frac{d\bar{x}_i}{dt} = k_0\bar{x}_i + \sum \text{(reaction terms)} + \delta - K\bar{x}_i, \tag{10}
\]

\[
\frac{d\bar{x}_i}{dt} = -k_0\bar{x}_i + \sum \text{(reaction terms)} - K\bar{x}_i. \tag{11}
\]

The (reaction terms) are defined in the appendix in the discussion following Eq. (28). For a species outside the food set, the kinetic equations are of the same form, except that \(\delta = 0\). There is no net flow of mass from the food set to the other elements of the system, which drives it away from equilibrium.

Because the reaction (within the biomass), the total mass in the reaction vessel always goes to a fixed point, independent of initial concentrations. To see this, note that only the last two terms in Eq. (10) and the last term in Eq. (11) change the total mass. The total mass concentration is proportional to \(m = \sum n(i)x_i\), where \(n(i)\) is the length of the ith species. Letting \(N_f\) be the number of elements in the food set, the rate of change of the total mass concentration is given by a simple differential equation,

\[
\frac{dm}{dt} = N_f\delta - Km, \tag{12}
\]

which has a global fixed point \(m_0 = N_f/K\delta\). This means the initial mass is irrelevant anyway, and so for convenience in our simulations, we choose \(m(0) = m_0\). Thus, there are effectively only two parameters relating to the flow of mass through the system, which can be \(\delta\) and \(K\), or equivalently \(\delta\) and \(m_0\).

Since \(\delta\) and \(K\) are not intuitively easy to interpret, it is sometimes useful to quote results in terms of the mean reaction number \(r\). This definition is the mean number of times a given monomer participates in a reaction, on average, from the time it enters the vessel until the time it is flushed out. At equilibrium, \(r\) is infinite, and when the other parameters are fixed, it decreases monotonically as \(\delta\) increases.

2.4.2 PYROPHOSPHATES As we demonstrate in subsection 2.6, catalytic focusing requires conditions that favor polymerization. The tendency to polymerize can be enhanced by an appropriate input of energy. The mechanism that we investigate here involves pyrophosphate molecules, which play a role analogous to that of \(ATP\) in contemporary organisms. This mechanism is supported by early experiments. The detailed sequence of reactions was suggested by Ron Fox, and is illustrated in Table 1.

It proceeds as follows. Light causes the formation of pyrophosphate molecules \(p_2\), which is balanced by hydrolysis. When a pyrophosphate molecule binds to polymer \(A\), it creates the energized form \(A^*\) and releases a phosphate atom in the process. \(A^*\) may hydrolyze, releasing the other phosphate atom, or it can bind to another polymer \(B\) (in the presence of the catalyst \(E\)). This occurs with a rate constant \(k_{r}\), which is greater than the unenergized rate constant \(k_f\). Thus, the addition of energy favors polymerization.

Based on simulations involving the full reaction scheme shown in Table 1, we found that when the concentration of pyrophosphate and the input of light are sufficiently high, the behavior is roughly equivalent to that obtained by simply using the equations given in the Appendix, with the effective forward rate constant equal to \(k_f\). For convenience, in the numerical experiments described here we simply assume that the parameters quoted correspond to \(k_f\), and use the simpler equations which do not involve pyrophosphate.

2.5 REACTION GRAPH

Each distinct monomer can be assigned a character from a fixed alphabet, \(a, b, c, \ldots\). A polymer can then be represented as a character string, for example \(\text{acabace} \ldots\). We assume that the polymers are oriented, so that \(abc\) and \(cba\) are different strings. The topological structure of a network of reactions, each of the form of Eq. (8), can be represented as a polygraph with two types of nodes and two types of connections, as illustrated in Figures 1 and 2. One type of node represents the
polymer species and is labeled by the corresponding string. The other type of node represents the catalyzed reaction and is labeled by a black dot. The polymers that participate in a reaction are connected to the corresponding reaction node by reaction links (black arrows), which point in the direction of condensation. Each polymer is connected to the reactions it catalyzes by a catalytic link (dotted line). Each reaction has at least four links: three reaction links, and one or more catalytic links.

2.6 CATALYTIC FOCUSING

Under appropriate conditions catalysis can focus most of the material of a reaction network into only a few species. The basic idea can be grasped by considering the simple reaction network

\[ ba + H \rightleftharpoons a + b \rightleftharpoons ab + H, \]

as shown in Figure 1. Assume \( a \) and \( b \) are supplied at rate \( \delta \), and diffuse out of the container with rate constant \( K \), as described in subsection 2.4. For simplicity, assume the concentrations of \( E \) and \( H \) are maintained at fixed values. Neglecting saturation, according to the approximation of Eq. (9), the rates of change of \( ab \) and \( ba \) are

\[
\frac{[ab]}{[ba]} = \gamma \left( k_1[a][b] - k_2[H(ab)] - K[ab] \right),
\]

\[
[ba] = k_f[a][b] - k_2[H[ba] - K[ba]],
\]

where \([ab]\) is the concentration of polymer \(ab\). Setting the derivatives to zero and using the mass conservation condition of Eq. (12) gives

\[
\frac{[ab]}{[ba]} = \frac{1 + \beta}{1 + \frac{\gamma}{\gamma}},
\]

\[ \beta = \delta/m_0k, \]

\[ H \] is a dimensionless parameter related to the deviation from equilibrium, where \( m_0 = a(0) + b(0) \) is the total concentration of monomers. Note that \( \beta \geq 0, \gamma \geq 1 \) is a dimensionless parameter that characterizes the strength of catalysis. \( \gamma \geq 1 \), and \( \gamma = 1 \) corresponds to an uncatalyzed reaction.

Under what circumstances is the concentration of \( ab \) much greater than that of \( ba \)? At equilibrium \( \beta = 0 \) and the ratio of \([ab]\) to \([ba]\) is one. This ratio can become large only when \( \beta \gg 0 \), i.e., only when the system is driven well away from equilibrium. When \( \beta \gg \gamma \) this ratio approaches \( \gamma \); when \( \beta \ll \gamma \) it approaches \( \beta \). Thus, by varying \( \gamma \) and \( \beta \) the concentration of \( ab \) relative to \( ba \) can be made arbitrarily large.

![Figure 1](image)

FIGURE 1 A simple network illustrating how steady-state concentration can be boosted by catalytic activity. \( a \) and \( b \) are driven at a fixed rate \( \delta \), and the enzyme \( E \) is maintained at a fixed concentration.

Note that the ability to focus comes about because the formation of one species is catalyzed, while that of the other is not. If all reactions were catalyzed equally, with equal kinetic parameters, there would be no focusing; the concentration of \( ab \) would equal that of \( ba \). Focusing thus requires specific catalysis, in which some reactions are catalyzed more strongly than others.

2.7 AUTOCATALYTIC SETS AND METABOLISMS

To achieve catalytic focusing the enzyme \( E \) must be maintained at high concentration. One way for the system to accomplish this by itself is through an autocatalytic reaction, in which one of the products catalyzes its own formation. A simple example is

\[ A + B \rightleftharpoons C + H. \]

If we set \( C = ab = E \) in reaction (13), then the enzyme is produced automatically, and the focusing maintains itself.

Simple autocatalytic reactions such as reaction (16) are obviously very special. A more common situation occurs when autocatalysis involves a cooperation between reactions, in which one species catalyzes the formation of another. An autocatalytic set is defined as a set of chemical species such that each member of the set is produced by at least one catalyzed reaction involving only members of the set. This notion was introduced by Calvin,\(^9\) Eigen,\(^10\) Kauffman,\(^23\) and Rössler.\(^27\) Since the reactions we are considering are reversible, a species can be produced either by cleavage or condensation. Thus reaction (16) is an autocatalytic set, and so is

\[ A + B \rightleftharpoons C + H. \]
To make this distinction, we define an \textit{autocatalytic metabolism} as an autocatalytic set whose concentrations make significant departures from the values they would have if none of the reactions were catalyzed. The phrase "significant departures" is subjective, and is admittedly rather vague. However, from an operational point of view, in our simulations we often see a clear distinction between autocatalytic sets that can function as metabolisms and those that cannot, as shown in Section 6.

3. SIMULATION

In principle the kinetic equations are all we need to know in order to simulate the behavior of a reaction network. In practice, however, there are two major problems: The first is that the kinetic parameters cannot be determined from first principles. To deal with this we construct an artificial chemistry, as discussed in subsection 3.1. The second problem is that there are an infinite number of possible reactions, and it is intractable to solve all of them; we must focus our computational resources on only the most relevant ones. Our method for doing this is discussed in subsection 3.2.

3.1 ARTIFICIAL CHEMISTRY

In a real chemical system the efficiencies and rate constants of the reactions depend on detailed properties of chemical composition, as well as on thermodynamic parameters such as temperature and pressure. While a computation of these constants from quantum mechanics and statistical mechanics is possible in principle, from a practical point of view, at this point in time it is hopelessly intractable.

To circumvent this problem, the approach introduced by Kauffman, Farmer et al.,11,12 and Bagley et al.2 is to invent an artificial chemistry, a set of rules stating which catalyzed reactions occur, and with what strength. An artificial chemistry cannot reproduce the behavior of a real chemistry in detail, but it may reproduce many of the correct qualitative properties. An artificial chemistry can produce complex behavior, even though it is simple from a computational point of view. By exploring different artificial chemistries, we can discover which properties cause significant changes in behavior, and which do not. We can begin with simple chemistries and move toward more complex chemistries, adding layers of realism as needed. The knowledge gained in this way can be useful in guiding experimental investigations of real systems, by pinpointing the essential quantities that need to be measured in experiments in order to make the model more realistic.

Since our primary interest is in understanding the effect of catalysis, we first address the problem of assigning a catalytic efficiency to each reaction. We do this using two different methods. In the first, we construct a completely disordered artificial chemistry, by assigning catalytic efficiencies at random, and in the second, we
construct a highly ordered artificial chemistry, assigning them with a string matching algorithm. The random method is more disordered than real chemistry, and the string matching method is more regular than real chemistry. From a qualitative point of view, we hope that real chemistry lies somewhere between these two extremes.

To be strictly correct, every possible reaction should be included in the reaction graph. However, in practice the reaction graph must be trimmed, so that computational resources are used only for the most essential reactions. We take advantage of the fact that the vast majority of reactions are catalyzed only weakly and can be treated essentially as spontaneous reactions. The graph represents only those reactions with sufficient catalytic efficiency to make them significantly different from the corresponding spontaneous reactions. Thus, when we refer to a "catalyzed reaction," we mean a "strongly catalyzed reaction," and when we refer to a spontaneous reaction, we mean a "weakly catalyzed reaction."

3.1.1 RANDOM ASSIGNMENT OF REACTIONS In some cases changing a single monomer can have a dramatic effect on the chemical properties of a polymer, either because it causes a drastic change in the configuration of the polymer or because it alters the properties of a critical site. If this were always the case, then chemistry would be random. For a random chemistry there is no correlation between chemical formulas and chemical properties. This is unrealistic. However, it does have the advantage of being easy to implement, and lies at one extreme in the space of all possible chemistries.

Following Kauffman, we assume that out of all possible spontaneous reactions, only a fraction $p$ are catalyzed with sufficient strength to be significantly different from spontaneous reactions. The set of reactions that is catalyzed is chosen at random. To see the basic idea, imagine creating a list of all possible catalyzed reactions. For $m$ distinct monomers the number of species of length $n$ is $m^n$, the number of possible spontaneous reactions is the order of $m^n$, and the number of possible catalyzed reactions is the order of $m^p$. The reactions that are strongly catalyzed can be determined by flipping a biased coin that returns heads with probability $p$. Reactions that receive heads are assigned a non-zero value of $v$, and all others are assigned $v = 0$. The random rule generates an ensemble of possible chemistries, corresponding to all possible sequences of random choices.

Operationally the procedure described above would be very time consuming, since $m^p$ can be a very large number. It can be made much more efficient by decomposing the problem properly, focusing attention only on reactions involving species that are already present in the reaction vessel, and taking care to avoid double counting, as described by Farmer et al.

To determine catalytic efficiencies, one simple possibility is to set $v = \text{constant}$, so that all the reactions on the graph are catalyzed with the same efficiency. Another natural possibility is to choose the catalytic efficiencies at random according to a given probability distribution, for example by making the probability of a given efficiency uniform within given maximum and minimum values. We employ both of these in our simulations.

3.1.2 ASSIGNMENT OF REACTIONS BY STRING MATCHING The match rule provides an alternative artificial chemistry that is probably closer to real chemistry than the random rule discussed above. It lies at the opposite extreme—while the random rule is too disordered, the match rule is probably too ordered. For the match rule, changing a single monomer in a given polymer only causes a small change in its chemical properties. Two similar polymer strings always have similar chemical properties. The match rule assumes that the information contained in the string of a given polymer contains all the information needed to specify its chemical properties.

We roughly follow the approach used to model the immune system by Farmer et al. For convenience, in this discussion we assume a two-letter alphabet consisting of $a$ and $b$, although the rule is easily generalized to a larger alphabet. The two reactants $A$ and $B$ in Eq. (8) join together to form $C$. The character string corresponding to $C$ is matched against that of enzyme $E$. There are several possible alignments; we require that the string $E$ span the binding site between $A$ and $B$. Each allowed alignment of $E$ against $C$ is given a match score according to the number of complementary matches, i.e., the number of cases where an "a" is paired against a "b." We then compute the probability $P$ that a score as good or better would be obtained if the strings $E$ and $C$ were generated at random. We use this to define a quantity we call the specificity $s = 1/P$. The catalytic efficiency depends on the specificity through a function $v(s)$. We typically assume that high specificity corresponds to higher catalytic efficiency, and choose $v(s)$ to be linear. For a given choice of $A$, $B$, and $E$, the total catalytic efficiency is the sum of the efficiencies computed for each of the allowed alignments. For a more detailed description, see Bagley.

The match rule assigns a catalytic efficiency to every possible catalyzed reaction. The reactions with catalytic efficiencies above a given threshold $v_0$ are installed in the reaction graph. The match rule is completely deterministic, and generates a unique chemistry. The requirement that the specificity exceed a fixed threshold implies that very short polymers cannot participate in catalyzed reactions. Thus, the properties of the match rule are quite different from those of the random rule.

At this point we have studied the random rule more thoroughly than the match rule. We intend to present results using the match rule in a future paper.

3.1.3 OTHER KINETIC PARAMETERS The other relevant kinetic parameters that may vary from reaction to reaction are the forward reaction rate $k_f$, the backward reaction rate $k_b$, and the unbinding constant $k_u$. In order to use the approximation for the saturation problem described in the Appendix, it is necessary that $k_u$ be the same for all reactions. Since we can vary the catalytic efficiency for each individual reaction, this is not a serious problem.

The rate constants $k_f$ and $k_b$ play an important role. At equilibrium diverse rate constants cause the polymers of a given length to have nonuniform concentrations. However, since to first approximation the rate constants only depend on the two monomers at the binding site, the resulting nonuniformities in the concentration profile are much more regular and less pronounced than those resulting from
catalytic focusing. For convenience we typically assume that $k_f$ and $k_r$ are the same for all reactions, although in some cases we also vary them randomly.

### 3.2 METADYNAMICS

The number of possible reactions is infinite. Of course, in reality only a finite number are important. Unfortunately, it is usually impossible to state in advance which reactions can be neglected. A metadynamical simulation attempts to solve this problem by restricting attention to a variable set of reactions and chemical species, and adding or deleting as needed.

The problem of determining the essential reactions is particularly severe for polymer chemistry, where the number of possible reactions grows exponentially with the length of the polymers. For $m = 20$ and $n = 10$, for example, there are already more than Avogadro's number of possible catalyzed reactions. Even if we knew the rate constants for every reaction, we could never hope to keep track of all of them in a computer simulation.

A real chemostat only contains a finite number of molecules and hence a finite number of species, with a finite number of possible reactions between them. Continuous differential equations fail to exploit this. Suppose, for example, that at $t = 0$ all the concentration is placed in a few species, and the concentrations of the rest are set to zero. According to the laws of continuous kinetics, for $t > 0$ there are generically an infinite number of species with non-zero concentrations. This unrealistic result comes from the approximation made in treating a discrete population of molecules as though it were continuous. This is equivalent to the assumption of a well-stirred reaction vessel with infinite volume. For the problem that we address here, the fact that the reaction vessel is finite is of critical importance.

One solution to this problem is to treat the kinetics as a random process, using integer populations and simulating molecular collisions through Monte Carlo techniques. This method is efficient when the populations are low, but is computationally inefficient when the populations are high. If the population is $10^5$, for example, a differential equation solver might change the concentration by as much as one percent in a single step and still retain reasonable accuracy, while the same change with a stochastic simulator requires $10^5$ steps.

The metadynamics approach offers an alternative that is computationally more efficient when there is a wide disparity in population size. At any given time the reaction network is modeled by a finite set of continuous differential equations, representing the dominant reactions. The topological structure of this set of differential equations is represented as a graph containing only species that are either present in the reactor, or that can be produced by other species present in the reactor. As the concentrations change the dominant species and reactions may also change. The graph is changed to reflect this, which in turn changes the differential equations. The dynamics occur on two time scales, the faster time scale of the differential equations and the slower time scale for changing the graph. A typical example of a metadynamics simulation is shown in Figure 3.

![Figure 3](image.png)

**FIGURE 3** A typical metadynamics simulation. The logarithm of the concentration of each species is plotted against the logarithm of time. Initially only the four food set species $a$, $b$, $ab$, and $ba$ have non-zero concentration. Catalyzed reactions within the food set produce new species, which in turn have more catalyzed reactions. At the point where we begin the graph at $t = 100$, four new species have appeared, so there is a total of eight polymer species. Each new species can be seen appearing at the bottom of the graph as it crosses above the threshold. The system eventually approaches a steady state solution with 22 species above the threshold. The parameters are the same for all reactions, and are $k_f = 10^7$, $k_r = 10$, $\nu = 10^4$, $k_u = 10^4$, $\delta = 10^9$, and $m_0 = 3$.

To take into account the fact that the container is finite, we impose a concentration threshold corresponding to the presence of a single molecule. If the concentration of a given species is significantly below this threshold, then that species is unlikely to be present. Therefore it cannot participate in reactions that produce other new species, and these reactions can be safely ignored. We only include reactions between species that are above threshold. They may produce new species, which rise above threshold; when this happens the new species are installed in the graph and allowed to catalyze new reactions. Similarly, species that were formerly above threshold may fall below it and be removed.
Since the number of species with concentrations above threshold at any given time is finite, the graph is also finite. Adjusting the threshold makes it possible to keep the graph from becoming unmanageably large—smaller reaction vessels have higher concentration thresholds, corresponding to smaller graphs. By making the reaction vessel sufficiently small, we can ensure that the graph has less than a given number of elements. We typically strive to simulate a container that is roughly the size of a bacterium, although because of limitations in computer resources we are often forced to use smaller containers.

In our simulations the system always approaches a unique fixed point. This appears to be true independent of the way we model the reactions, i.e., whether or not we account for saturation or whether or not we include pyrophosphates. This suggests that there is a Lyapunov function for catalyzed kinetic equations in this class.\[8\]

We are able to speed up our simulations by several orders of magnitude by assuming the existence of a unique fixed point, which implies that the steady-state deterministic equations can be solved algebraically. We call the fixed point for any given set of differential equations a dynamical fixed point. The metadynamical simulation is simplified as follows: We find the dynamical fixed point of the current set of differential equations. We then examine it and check to see whether any species have moved above or below threshold in comparison to the previous dynamical fixed point. If so, we change the differential equations accordingly and find a new dynamical fixed point, and repeat the process. Eventually there are no changes compared to threshold and this procedure stops. We call the final dynamical fixed point the metadynamical fixed point. By reducing the metadynamics to a sequence of algebraic operations, on a typical workstation a simulation such as that of Figure 3 is compressed into a few seconds.

Note that although the deterministic simulation described above always reaches a metadynamical fixed point, once we reincorporate the stochastic effects of spontaneous reactions, the system may hop between many different metadynamical fixed points and the long-term dynamics become quite interesting. This is the basis for our claim that autocatalytic metabolisms can evolve, and is discussed in more detail in a companion paper.\[4\]

3.3 THE BACKGROUND OF UNCATALYZED REACTIONS

Restricting attention solely to the reaction graph is a good assumption as long as the reactions on the graph dominate over everything else. This is not always the case. For example, spontaneous reactions always dominate near equilibrium. To study the competition between catalyzed and spontaneous reactions, the spontaneous reactions must be taken into account, at least as an aggregate.

\[4\] These equations are reversible, which makes them different from many other autocatalytic equations that are known to display limit cycles, chaos, and hysteresis.

FIGURE 4 The "mandala" of polymer species. The catalytic network, shown with white letters on black ovals, includes the food set, but also consists of other species produced by catalytic reactions. There must be a continuous path in the corresponding graph of catalyzed reactions from members of the food set to each member of the catalytic network; furthermore, the concentration must be above a concentration threshold corresponding to the presence of a single molecule. The background consists of everything that is not in the catalytic network. The shadow, shown by white letters on a grey background, is a special subset of the background, consisting of species that can be produced by a spontaneous reaction involving only themselves and members of the catalytic network.
concentrations of the reaction network are high, the shadow plays a special role because it is maintained at concentrations above the rest of the background. The shadow is in this sense similar to Eigen’s quasi-species.\(^\text{15}\)

For clarity we will first discuss the problem of modeling spontaneous reactions alone, without catalyzed reactions, and then return to discuss the case when spontaneous reactions are in competition with catalyzed reactions.

The difficulty of modeling spontaneous reactions comes about because there are so many of them. To make the problem tractable, we assume that all reactions have the same forward and backward rate constants \(k_f\) and \(k_r\). The problem is then equivalent to that of modeling the spontaneous reactions in a system with only one distinct type of monomer \((m=1)\). We can lump together all polymers of a given length \(n\), adding together their concentrations to get a combined concentration \(s_n\). The allowed reactions are of the form

\[
s_i + s_j \to s_k + H.
\]

A reaction is only possible if \(i + j = k\). The contributions to the kinetics are now described in the order that the corresponding terms appear in Eq. (19): \(s_k\) receives contributions from condensation reactions between smaller species and from cleavages of longer species. It loses to condensations between polymers of length \(k\) and polymers of other lengths, as well as to cleavages of itself. If there are polymers of length \(k\) that are part of the food set, \(s_k\) will also gain from external driving.

The resulting rate equation for \(s_k\) is

\[
\frac{ds_k}{dt} = k_f \sum_{i+j=k} s_i s_j - k_r \sum_{n\neq k} (n-k+1)s_n - k_f s_k \sum_{j=1}^{m^k} s_j - k_r (k-1)s_k + n_k \delta \sigma_k - Ks_k,
\]

(19)

where \(\delta = 0\) if \(k\) is outside the food set, and \(n_k\) is the number of elements of length \(k\) in the food set. For an alphabet of \(m\) monomers, with the assumption of uniform reaction rates, the concentration computed above is divided evenly among all the species present. It is

\[
a_k = \frac{s_k}{m^k}.
\]

(20)

We now outline our approach for treating the competition between spontaneous and catalyzed reactions. Let \(s_k\) now represent the sum of the concentrations of the polymer species of length \(k\), excluding the catalytic network. The dynamics of the background due to its interactions with itself can still be taken into account by equations of the form of Eq. (19). However, the concentration for a single species is now defined to be

\[
a_k = \frac{s_k}{(m^k - l_k)},
\]

(21)

where \(l_k\) is the number of elements of the catalytic network of length \(k\). The coupling to the autocatalytic set can be taken into account by carefully counting all of the interactions. The members of the catalytic network contribute to the spontaneous background through their condensation and cleavage reactions. Similarly,

they receive concentration from the spontaneous background. The only approximation necessary comes from the assumption that all the elements of the background have the same concentration. This is not strictly true. In particular, the concentrations in the shadow are typically above those of the rest of the background. However, this is a second-order effect, and we feel that in most cases our simulations are approximately correct. The problem of coupling the catalytic network to the spontaneous background is discussed in more detail by Bagley.\(^\text{5}\)

4. EMERGENCE

In this section we present simulations of catalytic reaction networks under several different conditions. We begin by demonstrating that, under appropriate circumstances, autocatalytic metabolisms emerge at concentrations that are several orders of magnitude higher than those of the background. We explore the parameter dependence and show that there is a range of parameter values where autocatalytic metabolisms thrive.

4.1 DEPARTURE FROM EQUILIBRIUM

In Figure 5 we show a sequence of three simulations in which we drive the system further away from equilibrium by increasing the parameter \(\delta\). In each case we let the reaction vessel approach its steady-state behavior, and then plot the concentration of all the polymers in the vessel of length less than twelve. In Figure 5(a) \(\delta = 0.01\), the mean reaction number \(r = 210,000\), and the system is nearly at equilibrium. The observed behavior is in quantitative agreement with the predictions of subsection 2.2. The concentration falls off exponentially with length, all species of a given length have the same concentration, and there is no interesting structure in the concentration profile.

In Figure 5(b) \(\delta = 10^6\), which corresponds to a mean reaction number of approximately 33. The concentrations of the catalyzed reaction network are orders of magnitude above those of the background, and most of the mass of the system is concentrated in the autocatalytic metabolism.

Finally, in Figure 5(c) we show the case when \(\delta = 10^{7.5}\), corresponding to a mean reaction number of roughly 0.5. The dominance of the autocatalytic metabolism is less evident—the flow of matter through the system is so high that the reaction network has a much smaller effect on the composition of the system. This is because the flow of mass through the system is so large that there is no time for a given species to react before it is flushed from the system.

These simulations demonstrate the ambiguity of the word “non-equilibrium” in this context. On one hand, the parameter \(\delta\) is a control parameter that can be used to drive the system away from equilibrium. On the other hand, as demonstrated in Figure 5, increasing \(\delta\) does not necessarily make the physical properties of the
system deviates further from those at equilibrium. In terms of the non-uniformity of the concentration profile, \( \delta = 10^8 \) produces a larger deviation from equilibrium properties than \( \delta = 10^{7.5} \). The deviation from equilibrium properties is at a maximum when the driving away from equilibrium is finite. A flow of energy is needed to move the system away from the structureless equilibrium state, but too large a flow of energy again results in a structureless state.

We have explored several quantitative measures of the deviation of the physical properties of the system from those at equilibrium. One of these is the steady-state slope \( A \) of the concentration profile. In Figure 6 we plot the concentrations of the species in the network as a function of their length and compare them with the background, for a favorable set of parameters where the system exhibits an autocatalytic metabolism.\[6\] The logarithm of concentration as a function of length in Figure 6 gives roughly lines of slope \( A \), indicating that the concentrations decrease roughly exponentially with length. To measure \( A \) we use a least-mean-squares fit. The longer polymers of the network are at much higher concentrations than those of the background. Consequently, for the network \( A \) is much larger (less negative) than that it is for the background. \( A \) thus provides a measure of the deviation from the equilibrium profile.

In Figure 7 we plot \( A \) as a function of the mass flow \( \delta \). For small values of \( \delta \) the system is near equilibrium, and \( A \) for the network is nearly equal to \( A \) for the background. As \( \delta \) increases \( A \) increases, reaches a maximum, and then decreases. Near the maximum, \( A \) approaches zero, indicating the concentration profile is almost flat.

![FIGURE 5](image)

Concentration “landscapes” at different displacements from equilibrium.

The two horizontal axes correspond to the polymer species, arranged concentrically with the shortest polymers in the center, as in Figure 4. The vertical axis corresponds to the logarithm (base 10) of the concentration of each species. The food set consists of the polymers \( a \) and \( b \), using a variation of the network of Figure 2 with 116 catalytic links.

In (a) \( \delta = 0.1 \), and the system is near equilibrium; the concentration falls off exponentially with length but is otherwise featureless. In (b) \( \delta = 10^4 \), which gives the behavior that is most distinct from that at equilibrium; in this case the concentration of the autocatalytic set is many orders of magnitude above the corresponding equilibrium values. Finally, in (c) \( \delta = 10^{7.5} \) and the system is so far from equilibrium that it loses most of its interesting structure. (Note the change of scale in comparison with (b).) The other parameters are \( k_f = 6.49 \times 10^4 \), \( k_s = 2.50 \), \( \nu = 8.97 \times 10^3 \), \( k_u = 5.00 \times 10^4 \), and \( m_0 = 2.0 \).

![FIGURE 6](image)

Mean concentration as a function of polymer length at steady state.

The solid line (N) corresponds to the network, the dotted line (B) corresponds to the background, and the dashed line (E) corresponds to the equilibrium concentrations. The simulation is performed for the network of Figure 2 with parameters \( k_f = 6.49 \times 10^4 \), \( k_s = 2.50 \), \( \nu = 8.97 \times 10^3 \), \( k_u = 5.00 \times 10^4 \), \( \delta = 1.79 \times 10^4 \), and \( m_0 = 2.0 \).

\[6\] Note that this network has different parameters than those of Figure 5.
The mass concentrated in the metabolism provides another natural measure of the deviation of its properties relative to those at equilibrium. For example, in Figure 8 we plot the fraction of the mass in the background, the food set, and the catalytic network (subtracting the food set), as a function of $\delta$. There is a central regime where the majority of the mass of the system is concentrated in the autocatalytic metabolism. Note that this regime overlaps with the regime where $\Lambda$ is large. However, the two are somewhat skewed; $\Lambda$ peaks at roughly $\delta = 10^5$, while the mass peaks when $\delta > 10^3$.

We feel that the need for a balanced energy flow for “interesting behavior” reflects a general principle. Another possible example is the fact that life evolved on Earth and not on Mercury or Pluto.

4.2 DEPENDENCE ON PARAMETERS

How special are the parameters for which autocatalytic metabolisms occur? To answer this question as quantitatively as possible, we have systematically explored the parameter space, testing for the presence of autocatalytic metabolisms. We used two measures of the dominance of the autocatalytic set. One of these is $\Lambda$, and the other is the mass ratio $R = N/(B + F)$, where $N$ is the mass of the network (neglecting the food set), $B$ is the mass of the spontaneous background, and $F$ is the mass of the food set.

In Figure 9 we show the behavior of $\Lambda$ under variations of $\delta$ and $\nu$. Note that $\Lambda$ remains near zero for a broad range of parameter values. For comparison, in Figure 10 we plot the mass ratio $R$ as a function of the same parameters, but for a network with fewer catalytic links. The behavior is more sharply peaked, but there is a broad range in which the autocatalytic set contains the majority of mass in the reaction vessel.

In Figure 11 we show the behavior of $\Lambda$ under variations of $\nu$ and the unbinding rate constant $k_u$. This figure illustrates how distinct the behavior of the autocatalytic metabolism is from that of the background. In one regime, roughly corresponding to lower values of $\nu$ and lower values of $k_u$, $\Lambda$ behaves just as it does for the spontaneous background. It is more negative and forms a relatively flat surface as a function of parameters. The other regime, which corresponds to larger values of the two parameters, has higher values of $\Lambda$. The transition from one regime to the other is quite sharp.

Finally, to illustrate the effect of varying the forward rate constant $k_f$, in Figure 12 we show the effect of varying $k_f$ and $\delta$. Once again we see a broad parameter

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**FIGURE 7** The slope of the concentration profile vs. the mass flux $\delta$. (See Figure 6 for an example of a concentration profile.) The network (N) corresponds to the solid curve, the background (B) to the dotted curve, and equilibrium (E) to the dashed curve. Parameters are the same as those for Figure 6, except that $\delta$ is varied.

**FIGURE 8** The fraction of the total mass for the network (N), the background (B), and the food set (F). (The fraction for the network excludes the food set.) The simulation is the same as that of Figure 7.
**FIGURE 9** $\Lambda$ vs. $\log_{10} \delta$ and $\log_{10} \nu$ for the variation of the network of Figure 2 with 118 catalytic links. The parameters are $k_f = 3.02 \times 10^4$, $k_r = 2.70$, $k_u = 7.11 \times 10^4$, and $m_0 = 2.0$.

**FIGURE 10** The mass ratio $R = N/(B + P)$ vs. $\log_{10} \delta$ and $\log_{10} \nu$ for the network of Figure 2. $k_f = 6.49 \times 10^2$, $k_r = 2.50$, $k_u = 5.00 \times 10^4$, $m_0 = 2.0$.

**FIGURE 11** $\Lambda$ vs. $\log_{10} \nu$ and $\log_{10} k_u$ for the variation of the network of Figure 2 with 118 catalytic links. $k_f = 3.02 \times 10^4$, $k_r = 2.70$, $\delta = 1.41 \times 10^2$, and $m_0 = 2.0$.

**FIGURE 12** $\Lambda$ vs. $\log_{10} \delta$ and $\log_{10} k_f$ for the variation of the network of Figure 2 with 118 catalytic links. $k_r = 2.70$, $\nu = 5.26 \times 10^5$, $k_u = 7.11 \times 10^4$, and $m_0 = 2.0$. 

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*Spontaneous Emergence of a Metabolism*
regime in which the values of the slope are quite high. Interestingly, for small values of $\delta$ there is a regime in which increasing $k_f$ beyond roughly $10^5$ causes a decrease in $A$.

It is evident from Figures 6–12 that for reaction networks with fixed topologies and uniform kinetic parameters, the dependence on parameters is fairly smooth. The parameters that approximately give the maximum value of $A$ are shown in Table 2. To measure the size of the regime where autocatalytic sets are dominant, we swept the parameters of the system one at a time, holding the other parameters constant at values corresponding to the maximum. We somewhat arbitrarily say that the system supports an autocatalytic metabolism if the mass ratio is greater than one. For convenience we held the reaction graph fixed. In one case we used the graph of Figure 2 which has a minimal number of catalytic links; in the other case we used a graph with the same reactions, but 118 catalytic links. The results are summarized in Table 2.

| Table 2 | Parameter regime that supports autocatalytic metabolisms. The kinetic parameter values roughly corresponding to the maximum value of $A$ are shown in the third column. To measure the size of the parameter regime that supports autocatalytic metabolisms, we varied each parameter one at a time, holding the others constant at their optimum values. We say that the system supports an autocatalytic metabolism when the mass ratio $R > 1$. For the experiment shown in (a) we keep the reaction graph fixed at that of Figure 2, which has 15 reactions and 15 catalytic links. In (b) we used a variation with the same reactions but 118 catalytic links. |
|-----------------|-----------------|-----------------|-----------------|
| **Global parameters** | **Description** | **Optimum** | **Range** |
| (a) | | | |
| $k_f$ | Forward reaction rate | $6.49 \times 10^2$ | $< 1, 10^{1.8}$ |
| $k_r$ | Reverse reaction rate | $2.50$ | Not computed |
| $\nu$ | Catalytic efficiency | $8.97 \times 10^6$ | $[10^{6.1}, > 10^{10}]$ |
| $k_u$ | Unbinding rate | $5.00 \times 10^4$ | $[10^3, 10^9]$ |
| $\delta$ | Mass flux | $1.79 \times 10^1$ | $[10^1, > 10^7]$ |
| (b) | | | |
| $k_f$ | Forward reaction rate | $3.02 \times 10^4$ | $< 1, 10^5$ |
| $k_r$ | Reverse reaction rate | $2.70$ | $[10^2, 10^3]$ |
| $\nu$ | Catalytic efficiency | $5.26 \times 10^5$ | $< 1, 10^9$ |
| $k_u$ | Unbinding rate | $7.11 \times 10^4$ | $[10^7, > 10^9]$ |
| $\delta$ | Mass flux | $1.41 \times 10^2$ | $[10^2, > 10^9]$ |

The overall conclusion that we draw from these studies is that in our artificial chemistry there is a broad regime in which autocatalytic metabolisms dominate. At this point, since the relevant kinetic parameters for real polymers are unknown, we cannot say whether they lie within the favorable regime. In particular, it seems that real catalytic efficiencies and forward rate constants are certainly on the low end of the favorable regime. Determining whether or not real systems lie within the favorable regime requires further investigation.

5. SENSITIVITY

The experiments of the previous section demonstrate that as long as the topology is fixed and the parameters are uniform, the behavior of autocatalytic metabolisms depends smoothly on parameters. In this section we investigate the sensitivity of catalytic networks to variations in topology and to non-uniform variations of the parameters of each individual reaction.

5.1 SENSITIVITY TO TOPOLOGY

The topology of a reaction network is clearly important. In Table 2, for example, we have already seen that a reaction network with more catalytic links produces autocatalytic metabolisms across a wider parameter regime. To test the dependence on topology more directly, we simply generate many different reaction networks at random and simulate their dynamics. For example, in Figure 13 we fix the reactions and the kinetic parameters, but randomly vary the catalytic links. When there is a small number of catalytic links, most reaction networks have highly negative values of $A$ near the equilibrium value. When there is an intermediate number of catalytic links, $A$ is highly sensitive to the topology—some networks depart significantly from equilibrium, while others do not. Finally, when there is a large number of catalytic links, almost all networks have roughly the same value of $A$, making a significant departure from equilibrium.

This result is intuitively reasonable. When there are only a small number of catalytic links, the configuration has to be rather special to generate an autocatalytic set. As the number of links increases the existence of an autocatalytic set becomes more and more likely. Note, however, that the existence of an autocatalytic set is by no means a sufficient condition for the system to generate a metabolism; there are many graphs that have autocatalytic sets in them, yet show no significant departure from equilibrium. Thus, there is also an additional dynamic effect—adding more and more catalytic links enhances the nonlinear feedback necessary to deviate from equilibrium properties.
5.2 SENSITIVITY TO PARAMETERS

For simplicity, in the results presented so far we assumed the kinetic parameters were the same for all the reactions in the network. This is obviously unrealistic; in any real chemical network, the parameters of each reaction are different. To test this sensitivity, in Figure 14 we vary the catalytic efficiency of each individual reaction while we hold the efficiency of the other reactions constant. Varying the efficiency of some reactions has a large effect on the steady-state slope, while varying others does not.

The autocatalytic set of Figure 14 has only 15 links, one per reaction, so perhaps it is not surprising that it is highly sensitive to the value of each catalytic efficiency. If a similar experiment is performed using the same reaction network, but with 118 catalytic links, the opposite occurs. The properties are relatively insensitive to the catalytic efficiency, as shown by Bagley.\(^5\) This robustness comes about because there are typically many catalysts for each reaction. This result is consistent with that of Figure 13.

Figure 15 shows the results of varying the the forward rate constant \(k_f\) for the catalytic network of Figure 2. As we see, there is also considerable sensitivity to variations of \(k_f\).

The experiments of this section demonstrate that the sensitivity of autocatalytic metabolisms to variations of parameters depends on the density of the reaction graph. In some parameter regimes the properties of autocatalytic metabolisms are quite sensitive to parameters. This is not surprising—the kinetic equations are highly nonlinear, and such sensitive behavior is common in highly nonlinear systems. This enhances the perception of autocatalytic metabolisms as “emergent phenomena.” Without actually performing a simulation, in these parameter regimes the behavior of an autocatalytic metabolism is difficult to predict in advance. In other parameter regimes, however, almost all parameter settings yield similar behavior.
5.3 ROLE OF DIVERSITY

In order for catalysis to be able to concentrate the material of the system into only a few species, it is necessary for some catalytic efficiencies to be high, while others are low. Thus, it is clear that diversity of parameters is a necessary condition of catalytic focusing. When we set $\nu > 0$ for some reactions and $\nu = 0$ for the remainder, we are automatically introducing diversity. It is our impression that additional diversity in the kinetic parameters, such as a distribution of values for $\nu$, $k_f$, and $k_r$, tends to increase the ability of the system to focus, but at present we have not demonstrated this quantitatively.

6. METABOLIC ROBUSTNESS

In this section we make more precise the sense in which the autocatalytic activity we have observed is similar to that of a metabolism, and investigate robustness to changes in the food set.

6.1 WHAT IS A METABOLISM?

A metabolism takes in material from its environment and reassembles it in order to propagate the form of the host organism. Autocatalytic metabolisms take material from the food set, and through a chain of reactions whose rates are accelerated by catalysis, boost the concentration of the members of the autocatalytic set. They differ from contemporary metabolisms in that they do not require a host organism.

Another function of a metabolism is to extract free energy from the environment and make it available for the functions of the organism. Through the pyrophosphate mechanism discussed in subsection 2.4.2, autocatalytic sets can extract energy from light and use it to energize polymers. This in turn alters the effective forward and backward rate constants, enhancing the tendency for polymerization. As we see in Figure 15, for an autocatalytic set to take over the medium in which it resides, it is critical that the rate constants be in the proper regime. For real chemistry the unenergized values of $k_f$ are probably in the unfavorable regime. This mechanism for extracting energy is critical to the existence and therefore the “functioning” of an autocatalytic metabolism.

In conclusion, although autocatalytic metabolisms are certainly much less sophisticated than the metabolisms of contemporary organisms, they exhibit the same rudimentary properties.

6.2 ROBUSTNESS

The fact that their underlying deterministic dynamical equations appear to have a unique stable fixed point endows autocatalytic metabolism with a considerable degree of robustness to trauma. A perturbation in the concentration of one of its elements, for example, quickly dies out. Thus, “self-repair” of autocatalytic metabolisms is built into their chemical kinetics.

Another notion of robustness concerns variations in diet. Some metabolisms can digest only one kind of food, and are, therefore fragile to changes in their environment. An example is the Panda, which can only digest bamboo. Others can digest many different types of material, and are, therefore, quite robust to changes in their environment. An example is the cockroach, which to the average urban dweller seems capable of digesting anything.

In this section we investigate the robustness of autocatalytic metabolisms by simply varying their food set. For example, in Figure 16 we simulate the steady-state concentration profile for five different food sets in conditions where everything else is held constant. In the first case, called the “default,” the food set consists of the four species $\{a, b, ab, bb\}$, which are all input at the same rate. This food set gives rise to clear dominance of the autocatalytic metabolism over the background. In the other cases the supply of at least one of these four species is eliminated, and

[1] However, a perturbation that adds new elements to the autocatalytic metabolism may be strongly amplified. This possibility is discussed in detail in the companion paper.
the influx of the other species is adjusted so that the total rate of influx of mass remains constant. The results are shown in Figure 16. The concentration profiles cluster into two groups: In cases I and III the concentration profile is essentially unaltered, giving the same dominance over the background observed with the default food set. Apparently the network is able to resupply the missing elements with little alteration. For the other two alterations, in contrast, the autocatalytic set appears to “die”: the concentrations change radically, dropping below the level corresponding to that with no catalysis. The clustering evident in Figure 16 demonstrates a qualitative difference between an autocatalytic metabolism that is “alive” and one that is “dead,” and reinforces our use of the word “metabolism” for this behavior.

Note that we originally thought that survival under changes of the food set might be easily explained by simply examining the reaction graph. However, there are several cases where the autocatalytic set dies even though a metabolic pathway is available that could potentially replenish the missing element. The persistence of the autocatalytic metabolism depends on the topology, but it also depends on complicated nonlinear cooperative effects associated with the kinetics.

7. DISCUSSION
7.1 RELEVANCE TO EXPERIMENTS

One of our ultimate goals is the discovery of autocatalytic metabolisms in the laboratory. We hope that the parameters quoted in Table 2 will at least provide a crude starting point for laboratory investigations. In any case, we have demonstrated some qualitative principles that may serve as a guide in real experiments, such as:

- For an autocatalytic metabolism to dominate over its background, polymerization must be favored even in the absence of catalysis. In other words, the primary role of catalysis is to focus the concentration into a few species, rather than to effect an overall tendency for everything to polymerize. Energetic chemical species such as pyrophosphate may play an important role in shifting the effective equilibrium constant to favor overall polymerization.
- It is important to drive the system the proper distance from equilibrium. For a significant effect the mean reaction number (as defined in subsection 2.4.1) should be in the vicinity of 100, although the optimal value also depends on the other parameters.
- The largest possible deviation from equilibrium occurs when all the parameters have finite values.[2] This is particularly surprising for the catalytic efficiency $\nu$ and the forward rate constant $k^+$, which one might have thought should be as large as possible. However, since $\nu$ and $k^+$ are almost certainly on the low side of the favorable regime, from a practical point of view in experiments, an effort should be made to find systems for which they are as large as possible.
In order for an autocatalytic set to be robust, the steady-state concentrations of its components should be within a few orders of magnitude of each other.

We have posed several questions whose experimental resolution could prove or disprove the assumptions of this model, or more likely, could guide revisions to make it more realistic. The number of individual kinetic parameters involved is enormous, and it is unrealistic to expect that they could be measured in detail. However, experiments that measure the distribution of parameters could be very useful. A measurement of the distribution of catalytic efficiencies as a function of length for randomly chosen polypeptides or nucleic acids would be particularly helpful.

By heating and cooling appropriate mixtures of amino acids, Sidney Fox has observed the formation of protein-like polymers that form cell-like enclosures, called proteinoid microspheres. It is possible that autocatalytic behavior played an important role in the formation of proteinoids, through a mechanism analogous to that discussed here. The problem of determining experimentally whether or not an autocatalytic metabolism is present in the laboratory has not been solved, and deserves more attention.

The formation of an enclosure is a desirable property for an autocatalytic metabolism. An autocatalytic metabolism might form its own enclosure, or it might inhabit a pre-existing enclosure. Morowitz et al. have proposed that the spontaneous formation of self-replicating lipid vesicles may have played a critical role in the formation of proteinoids, through a mechanism analogous to that discussed here. Furthermore, they argue that they might be capable of spontaneously producing amino acids, as well as a membrane potential. Such vesicles might provide a natural home for autocatalytic metabolisms. In the spirit of Oparin's original proposal, it is conceivable that an autocatalytic metabolism could establish a cooperative relationship with such a vesicle and evolve so that it was able to regulate its functions.

7.2 POSSIBLE SCENARIO FOR THE ORIGIN OF LIFE

Our results lend support to the hypothesis that the emergence of metabolisms preceded the emergence of template-based self-replication. The formation of autocatalytic metabolisms might play a role in several possible scenarios for the origin of life. The following gives one possible scenario:

The only possible exception is the mass concentration $m_0$, which we have not studied in detail.

7.3 CRITIQUE

We feel that there are at least two major problems with the model for autocatalytic metabolisms that we have presented here, at least as it pertains to experiments:

1. **Reliance on a mass flux to drive the system away from equilibrium.** It seems implausible that a flux of appropriate material would exist for a sustained period of time in a natural setting. Although we have also explored the possibility of light as an energy source, using pyrophosphate as an energy transducer, in its present form this mechanism essentially only serves to shift the equilibrium point, and cannot focus the material of the system to create a metabolism all by itself. Another mechanism for focusing the energy of the system, perhaps based on oxidation-reduction reactions, might solve this problem in a more realistic manner.

2. **Catalytic properties of short polymers.** The random rule generates reactions whose catalysts may be short polymers. This is unrealistic. The match rule we have formulated takes this to the opposite extreme, and only allows polymers longer than a certain critical size to participate in any catalytic reactions, either as reactants or as catalysts. Because of this the problem of maintaining a flux from the food set to the catalytic network becomes difficult. Of course, there...
are always spontaneous reactions between short polymers, which automatically expand the food set to a certain critical size, as described by Eq. (6). This might be sufficient, but in any case it complicates the problem.

We need to explore this question more carefully. One possible line of investigation is to use an artificial chemistry that is intermediate between the two proposed here, in which short polymers cannot be catalysts, but can be reactants. Still another solution might follow from finding an alternative energy source, as suggested in Eq. (1) above.

From a broader perspective of studying self-organization, this model suffers from a problem that is generic for many artificial life models. The process of making abstractions and simplifications builds a straitjacket that limits its emergent properties. The model presented here is essentially a “connectionist” model for chemistry, and suffers from the limitations inherent to this level of description. In order to make this model tractable, we have reduced real polymers, which have complex spatial structure, to an interaction rule and a set of rate equations that measures only one aspect of their aggregate behavior. To see further emergence of functional properties, such as the formation of enclosures, it may be necessary to study an artificial chemistry that is richer than that we have studied here. Unfortunately, such a model will inevitably require more computational power. This seems to be an unavoidable trade-off that occurs in all artificial life models. Even with a Connection Machine, one must be very clever to compete with Nature, who has more than Avogadro’s number of parallel processors.

7.4 ARE THEY ALIVE?

The artificial metabolism that we have studied here represents an organizational state that is between that of living and non-living systems. Given the appropriate conditions, within its own artificial universe an autocatalytic metabolism emerges spontaneously. It metabolizes energy, capturing the material of its universe and incorporating it into its own form. It also reproduces itself, although it does so continuously rather than discretely.

A good point of comparison is the class of oxidation reactions that produce “rust.” Rust can be an autocatalytic reaction, in the sense that the presence of rust engenders the formation of more rust. For example, collectors of bronze coins are very careful to avoid “bronze disease,” a form of rust that degrades a bronze coin, and spreads rapidly from coin to coin; if a coin is discovered to have bronze disease, it is immediately quarantined, to avoid the infection of other coins.

The key difference between autocatalytic metabolisms and simple autocatalytic reactions has to do with diversity and specificity. For a given metal, such as bronze, there is typically only one variety of rust. For a given polymeric medium, such as amino acids, there may be an enormous number of different autocatalytic metabolisms, each of which contains a different mixture of polymeric species. As we discuss in a companion paper, when the stochastic dynamics of spontaneous reactions are taken into account, which autocatalytic metabolism dominates at any given time depends on the history of the medium. Autocatalytic metabolisms can act like “seeds”—the presence of a minimal set of the elements of an autocatalytic metabolism, even at low concentrations, may allow that metabolism to take over its medium, and block the formation of other autocatalytic metabolisms. Autocatalytic metabolisms are both diverse and specific. Finally, as discussed in the companion paper, they also have the necessary feedback mechanisms to propagate themselves and amplify variations.

Insofar as an autocatalytic set is alive, it is a much cruder and less specific form of life than an organism with reproductive machinery based on templating. There is no genetic code for autocatalytic metabolisms. The message is simply the set of chemical species present in the soup.

There has been some debate concerning whether or not Gaia, i.e., the Earth, can be considered to be a living organism in and of itself. The relevant issues bearing on the question of whether or not autocatalytic metabolisms are alive are closely related to those for Gaia. The properties of autocatalytic metabolisms and Gaia are strikingly similar. First, an autocatalytic metabolism causes a substantial deviation from the equilibrium properties of the medium it occupies. Second, an autocatalytic metabolism may be the sole inhabitant of its medium. In this case, like Gaia, the autocatalytic metabolism evolves only through the richness of its own dynamics. One type of autocatalytic metabolism spontaneously evolves into another. There is internal competition and cooperation.

An objection to the consideration of Gaia as a living organism has been raised by Dawkins, who points out that since it is the sole inhabitant of its environment, there is no mechanism for selection. While we argue that autocatalytic metabolisms might “evolve” on their own, they need not remain isolated, and in general it is unlikely that they would. It is quite natural to imagine diverse autocatalytic metabolisms in separate enclosures, geographically isolated from each other, occasionally coming in contact, and competing and/or cooperating with each other. Some preliminary simulations involving spatially isolated autocatalytic metabolisms have been made by Bagley. Thus, Dawkin’s objections do not apply to autocatalytic metabolisms. A significant difference between autocatalytic metabolisms and Gaia is that the constituents of Gaia are themselves alive, whereas the constituents of autocatalytic metabolisms are not. Insofar as they are alive, Gaia is a simple organism made out of very complex organisms, whereas autocatalytic metabolisms are simple organisms made out of inanimate matter.

7.5 SUMMARY

We have demonstrated that under appropriate conditions an autocatalytic set can concentrate much of the mass of its environment into a focused set, with concentrations orders of magnitude above equilibrium. When this occurs we call the network an autocatalytic metabolism. The use of the term metabolism is supported
by the arguments of Section 6, and by the demonstration that some autocatalytic metabolisms are robust, and are capable of digesting a variety of different food sets.

Autocatalytic metabolisms can be highly sensitive to both the topology of the reaction network and the kinetic parameters of individual reactions. Because of this sensitivity, without performing a simulation it is difficult to state in advance whether or not a reaction network supports an autocatalytic metabolism. Nonetheless, there are ranges of parameters where most parameter values or most topologies support an autocatalytic metabolism. An indication of the range of parameter values where autocatalytic metabolisms are common is given in Table 2, and Figures 6-12.

These simulations demonstrate there is a favorable regime in which the flow of energy through the system is “just right” to make its properties radically different from those at equilibrium. There is an intriguing correspondence with behavior observed in other natural and artificial systems. For example, Langton has performed experiments constructing cellular automata rules at random. The rules are constrained to have a fixed probability of producing a non-zero state, which is regulated by a parameter \( \lambda \). In an analogy to thermodynamics \( \lambda \) corresponds to the Boltzmann factor \( \Delta E/kT \), and thus, in a generalized sense, measures how far the system is driven from equilibrium. Langton finds that the most interesting behavior occurs at a finite value of \( \lambda \), intermediate between the equilibrium value and the largest possible deviation from equilibrium. It is also interesting to note the roughly sigmoidal shape of the transition in Figure 9, which is similar to the phase transitions observed by Langton. This correspondence may be more than accidental, since both phenomena involve measurements of quantities that go through a phase transition when graph closure is obtained.

This model adds support to the idea that the emergence of a metabolism may have preceded the emergence of a self-replicator based on templating machinery. Perhaps more important, it provides an example of a “self-organizing structure” that is intermediate in complexity between life and non-life. There is an old idea, clearly articulated by Herbert Spencer, that in some general sense the evolution of form and organization depends on diversity arising from physical laws. Autocatalytic metabolisms illustrate how diversity in nonlinear feedback loops can couple to a flow of energy and radically alter the properties of a system from those at equilibrium, giving rise to information-carrying, propagating structures. This follows naturally from the rules of chemical kinetics. Although autocatalytic metabolisms are quite simple and lack the sophistication of modern organisms, they nonetheless fulfill the same functions as a contemporary metabolism, and challenge our notion of what it means to be alive.

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APPENDIX

KINETIC EQUATIONS

The effect of saturation can be taken into account exactly (within the limit of validity of mass action) by breaking down a catalysed reaction into each of its parts. This involves keeping track of several different intermediate products: A bound to E, B bound to E, C bound to E, and AB bound to E. Each reaction thus has four intermediaries. The autocatalytic networks that we simulate may have many species, and typically have many more reactions than species. Thus, treating each catalysed reaction in full detail would enormously increase the complexity of the simulation.

Another difficulty of the simulation comes about because the reactions are all reversible; if we allowed them to be irreversible, we would have effectively assumed the solution. This means that traditional schemes such as the Michaelis-Menten approximation are not applicable.

We simplify the saturation problem by assuming that A and B bind to each other as soon as they are bound to the enzyme E. We then break the reversible reaction of Eq. (8) into two irreversible reactions of the form

\[ A + B + E \xrightarrow{\nu_E} CE + H, \]
\[ C + H + E \xrightarrow{\nu_E} ABE. \]

(22)

ABE and CE are complexes formed in the reaction. The further dissociation of the bound complexes into free elements is treated through the irreversible reactions

\[ \frac{\text{CE}}{k_u} \rightarrow C + E, \]
\[ \frac{\text{ABE}}{k_u} \rightarrow A + B + E. \]

(23)

\( k_u \) is a constant that characterizes the rate at which the reactants unbind from the enzyme. Taking these together, and adding on the effect of the spontaneous reactions gives the rate equations

\[ \frac{dCE}{dt} = -k_u CE + k_f (1 + \nu_E) AB, \]
\[ \frac{dABE}{dt} = -k_u ABE + k_r (1 + \nu_E) CH, \]
\[ \frac{dC}{dt} = k_u CE - k_r (1 + \nu_E) CH, \]
\[ \frac{dA}{dt} = k_u ABE - k_f (1 + \nu_E) AB, \]
\[ \frac{dE}{dt} = k_u CE + k_u ABE - (1 + \nu_E)(k_f AB - k_r CH). \]

(24)

Note that we are using the same symbol for the bound complex \( ABE \) and its concentration. The reason the last equation differs from the others is because the enzyme E is involved in two bound complexes, and so has two terms involving \( k_u \).

In comparison with the exact reaction scheme, the approximate reaction scheme of Eq. (24) reduces the number of intermediates by a factor of two. However, as described in the remainder of this section, by making a simple substitution of variables, we can make a simplification that goes considerably beyond this, so that the number of equations that must be solved including saturation is only twice the number required without saturation.

Suppose that a given species \( i \) participates in many different reactions, and is bound up in many different complexes \( \xi_m \). For convenience, assume that each has the same dissociation constant \( k_u \). Using the approximation scheme above, the kinetic equation for \( dx_i/dt \) is of the form

\[ \frac{dx_i}{dt} = k_u \sum_m \xi_m + \sum \text{(reaction terms)}. \]

(25)

The first term takes into account all of the bound complexes that contribute to species \( i \) when they dissociate. (reaction terms) consists of a sum of terms, one for each reaction that species \( i \) participates in. Each term is of the form of Eq. (24), according to whether \( i \) plays the role of A, C, or E. Note that these only involve free species; none of the bound complexes \( \xi_m \) are involved. Furthermore, the dissociation terms that involve \( \xi_m \) are all linear. Thus, if we define the new variable

\[ \bar{x}_i = \sum_m \xi_m, \]

we can rewrite Eq. (25) as

\[ \frac{dx_i}{dt} = k_u \bar{x}_i + \sum \text{(reaction terms)}. \]

(27)

Similarly, each bound complex has a kinetic equation of the form

\[ \frac{d\xi_m}{dt} = -k_u \xi_m + \sum \text{(reaction terms)}. \]

(28)

There is only one reaction term, corresponding to the particular reaction associated with the complex. If we add up the equations associated with all the reactions that \( x_i \) participates in, we obtain

\[ \frac{d\bar{x}_i}{dt} = -k_u \bar{x}_i + \sum \text{(reaction terms)}. \]

(29)

Thus, we can solve for \( \bar{x}_i \) without having to solve for each intermediate complex from which it is derived. Since none of the reaction terms involve the bound complexes, we never have to solve for them. The number of variables is just twice the number of polymer species, rather than more than twice the number of reactions. We have taken saturation into account, at the small cost of doubling the effective number of species.
REFERENCES


