Homochiral oligopeptides by chiral amplification: Interpretation of experimental data with a copolymerization model.

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We present a differential rate equation model of chiral polymerization based on a simple copolymerization scheme in which the enantiomers are added to, or removed from, the homochiral or heterochiral chains (reversible stepwise isodesmic growth or dissociation). The model is set up for closed systems and takes into account the corresponding thermodynamic constraints implied by the reversible monomer attachments, while obeying a constant mass constraint. In its simplest form, the model depends on a single variable rate constant, the maximum chain length $N$, and the initial concentrations. We have fit the model to the experimental data from the Rehovot group on lattice-controlled chiral amplification of oligopeptides. We find in all the chemical systems employed except for one, that the model fits the measured relative abundances of the oligopeptides with higher degrees of correlation than from a purely random polymerization process.

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I. INTRODUCTION

In the transition from prebiotic racemic chemistry to chiral biology one scenario suggests that homochiral peptides must have appeared before the appearance of the primeval enzymes [1, 2]. While several stochastic synthetic routes for mirror symmetry breaking that convert racemates into nonracemates have been described [3, 4], the generation of long bio-like polymers [5] made up of repeating units of the same handedness requires elaboration of new synthetic routes. Polymerization reactions of racemic mixtures of monomers in solution are typically expected to yield polymers composed of random sequences of the left- and right-handed repeat units following a binomial or Bernoulli distribution. Thus the probability for obtaining oligomers with homochiral sequence becomes negligible with increasing length [1].

Recent investigations have proposed that N-carboxyanhydride (NCA) [5, 6] and thioester derivatives [5, 7] of amino acids might have operated as relevant precursors for the formation of the early peptides [8]. Results on the polymerization of NCA monomers in organic solvents, [9–14] in water [15–17] and in the solid state [18, 19] have been published. Luisi and coworkers [20, 22] have reported the polymerization of racemic α-amino acids in solution which yields small amounts of oligopeptides of homochiral sequence whose abundances with respect to the heterochiral chains exhibit a slight departure from the binomial distribution.

This problem of the random distribution can be overcame by catalyzed polymerization of amphiphilic amino acids, in racemic and nonracemic forms, which self-assemble into two-dimensional ordered crystallites at the air-water interface [24, 25]. Based on a process involving self-assembly followed by lattice controlled polymerization, Lahav and coworkers recently proposed a general scenario for the generation of homochiral oligopeptides of a single handedness from nonracemic mixtures of activated alpha amino acids [24, 25]. Initial non-racemic mixtures undergo a phase separation by self-assembly into a 2D racemic crystalline phase and a separate enantiomorphous 2D phase of the enantiomer in excess. Each of these crystalline phases has markedly different chemical properties, thus yielding products that differ in the composition of the oligomers. So, polymerization within the enantiomorphous crystalline phase yields homochiral oligopeptides of one handedness whereas the reaction controlled by the racemic crystallites yields racemic mixtures and heterochiral products. The combination of the two routes leads to an overall chiral amplification process.

In this paper, we are interested in the lattice-controlled polymerization reactions proposed by those authors. It is important to clarify at the outset what specific aspect of the overall experimental mechanism we want model here and the way we aim to do so. The proposed experimental scheme starts from an initial excess, say $S > R$ of monomers which undergoes an initial self-assembly process into two types of two-dimensional crystallites at the air/water interface. Once formed, each one of these two crystal phases participates in the control of a subsequent type of polymerization. Thus, the racemic crystallites polymerize racemic mixtures of oligomers and the heterochiral products, whereas the other pure enantiomorphous crystallite controls the polymerization of the isotactic chains, these are formed from the monomer in excess ($S$, in this example). However, the details of the polymerizations depend in a complicated way upon the specific packing arrangements of the crystal monomers and the possible reaction pathways taken within each crystallite phase. The authors of the experiments state that the connection between the monomer packing arrangements in the crystallites and the resultant composition of the various diastereoisomeric products is “not straightforward” [25]. We therefore opt for a simple model for interpreting their data. With this objective in mind, we present a copolymerization model for the interpretation of the experimental data. The model may be termed effective in the following sense: it presupposes or takes as given the prior formation of the self-assembled 2D crystallites at the air-water interface and is concerned exclusively with the subsequent polymerization reactions. Thus the complicated microscopic details referring to the monomer packing arrangements and reaction pathways within the crystallite self-assemblies are treated implicitly with our rate constants. Our copolymerization reaction rates can satisfactorily account for the different chemical properties of the two crystalline phases (racemic 2D crystallites and pure enantiomorphous 2D crystallites) that lead to the formation of racemic mixtures, heterochiral products and isotactic oligopeptides. We contrast the fits from our model with those assuming a purely random process that obeys a binomial distribution. The final justification for considering such an effective model rests on its ability to yield good fits to the data. The goodness of the fits obtained below demonstrates that the experimental data can be fit convincingly as if the simple scheme depicted pictorially in Fig. 1 were the sole mechanism leading to the observed relative abundances. This then gives additional meaning to to term “effective”, and in the operational sense.

II. THE COPOLYMERIZATION MODEL

Our starting point is a simple model for the copolymerization of two chemically distinct monomers displaying a wide variety of product sequence compositions. The model we introduce and study here is an appropriately modified and extended version of the one considered a few years ago by Wattis and Coveney [26].

The main important differences compared to prior and related models are that we (1) consider polymerization in closed systems [27]—so that no matter flow is permitted with an external environment—and (2) we allow for reversible
monomer association steps. We also correctly include the formation (and dissociation) of the heterodimer. It turns out this must be treated on a separate basis in order to avoid double counting, which if left unchecked, would lead to a violation in the constant mass constraint. Once the heterodimer is treated correctly, this implies that the homo-trimer must also be treated separately. Beyond this, the remainder of the hetero-oligomers can be treated in a uniform way.

First, we introduce the notation to be used. Polymers are classified by three quantities: the number of A monomers of which it is composed (subscript $r$), the number of B monomers which it contains (subscript $s$) and the final or terminal monomer in the chain, denoted by a superscript. In this scheme, the monomers are denoted by $A = C_{A}^{1,0}$ and $B = C_{B}^{0,1}$; pure homopolymers are denoted by $C_{A}^{r,0}$ and $C_{B}^{0,s}$; all copolymer chains $C_{A}^{r,s}$ or $C_{B}^{r,s}$ with $r, s \geq 1$ are heteropolymers. Note also that chains of the form $C_{A}^{0,s}$ and $C_{B}^{r,0}$ are forbidden. The corresponding time-dependent concentrations are denoted by lower case variables: e.g., $c_{r,s}^{A}(t)$ and $c_{r,s}^{B}(t)$. The model is then defined by the following
homo-polymerization reactions:

\[ \text{thiophene} + \text{thiophene} \rightleftharpoons \text{thiophene-thiophene} \]

\[ \text{C}_{r,s}^A + A \xrightleftharpoons[k_{aa}^*]{k_{aa}} \text{C}_{r+1,s}^A \]

\[ \text{C}_{r,s}^A + B \xrightleftharpoons[k_{ab}^*]{k_{ab}} \text{C}_{r,s+1}^B \]

\[ \text{C}_{r,s}^B + A \xrightleftharpoons[k_{ba}^*]{k_{ba}} \text{C}_{r+1,s}^A \]

\[ \text{C}_{r,s}^B + B \xrightleftharpoons[k_{bb}^*]{k_{bb}} \text{C}_{r,s+1}^B \] (4)

This model can accommodate any two chemically distinct monomers. For the purpose of this paper, we consider the case when \( A = \text{R} \) and \( B = \text{S} \) are two enantiomers.

The overall basic scheme must be broken down into several special subcases, especially important so as to avoid undesired double counting of the heterodimer and heterotrimer reactions, see Fig. 1. Once we treat these special cases, we then pass to the corresponding set of rate equations for the concentrations.

The formation of chirally pure polymer chains denoted by \( c_{n,0}^A \) and \( c_{0,n}^B \), for \( 1 \leq n \leq N - 1 \) is described by the homo-polymerization reactions:

\[ c_{n,0}^A + c_{1,0}^A \xrightleftharpoons[k_{aa}]{k_{aa}^*} c_{n+1,0}^A \]

\[ c_{0,n}^B + c_{0,1}^B \xrightleftharpoons[k_{bb}]{k_{bb}^*} c_{0,n+1}^B \]

(5)

\( N \) is the maximum chain length permitted. In our recently reported work \cite{2}, we considered that once a monomer has been added to a homopolymer of the opposite chirality (that is, "the wrong" monomer), the polymer is inhibited and further growth is halted. This polymer could not directly react anymore and could only lose its wrong terminal monomer through the inverse reaction. In the present model, we assume such a chain can continue to grow by adding monomers of both configurations. So, for \( 2 \leq n \leq N - 1 \), the hetero-polymerization or inhibition reactions are as follows:

\[ c_{n,0}^A + c_{0,1}^B \xrightleftharpoons[k_{ab}]{k_{ab}^*} c_{n+1,1}^B \]

\[ c_{0,n}^B + c_{1,0}^A \xrightleftharpoons[k_{ba}]{k_{ba}^*} c_{1,n+1}^A \]

(6)

For both homo- and hetero-polymerization reactions, represented by Eq. 5-6, the upper limits specified for \( n \) ensure that the maximum length for all oligomers produced (or consumed) by these reaction sets, both the homo- and heterochiral ones, is never greater than \( N \). In the remainder of this paper we will consider here the natural and chiral symmetric reaction rate assignments \( k_{aa} = k_{bb} \), \( k_{ab} = k_{ba} \), and likewise for the inverse rates, \( k_{aa}^* = k_{bb}^* \) and \( k_{ab}^* = k_{ba}^* \), reducing the number of independent rate constants to four.

Even if we have the information about the composition, we can only know the chirality of the last monomer attached to the chain, we have no information regarding the specific sequence. This implies that the following two reactions are indistinguishable:

\[ c_{1,0}^A + c_{0,1}^B \xrightleftharpoons[k_{ab}]{k_{ab}^*} c_{1,1}^B \]

\[ c_{0,1}^B + c_{1,0}^A \xrightleftharpoons[k_{ba}]{k_{ba}^*} c_{1,1}^A \]

Thus for all practical purposes, \( c_{1,1}^A \equiv c_{1,1}^B \), and this suggests using the following notation: \( c_{1,1} \equiv c_{1,1}^A \equiv c_{1,1}^B \), and to define a unique direct constant rate: \( k_h = \frac{k_{aa} + k_{bb}}{2} \), and an inverse one \( k_h^* = \frac{k_{aa} + k_{bb}}{2} \). Note that if \( k_{ab} = k_{ba} \), then \( k_h = k_{ab} = k_{ba} \). Due to these characteristics, we will treat the heterodimer in a different way compared with the other hetero-polymer. The reaction of the heterodimer formation is therefore:

\[ c_{1,0}^A + c_{0,1}^B \xrightleftharpoons[k_h]{k_h^*} c_{1,1} \]

(7)
As before, the reactives and products in Eq. (7) are the same, so the differences in the free energy between initial and final states should be the same in all the reactions in these equations, implying the following thermodynamic constraint on the reaction rates:

\[
\frac{k_{ab}}{k_{ab}^*} = \frac{k_{ba}}{k_{ba}^*}.
\]  

(8)

If the heterodimer formation were not to be treated in the separate way as we have done, and were to be included, e.g. in Eq. (5) by merely changing the lower limits for $n$ ($2 \leq n \leq N - 1$) by $1 \leq n \leq N - 1$, we would be making the mistake of double counting it. The same occurs for the heteropolymers formed from the addition of a monomer to a heterodimer. The two reactions of each pair of the following equations are also indistinguishable:

\[
\begin{align*}
C_{1,1} + C^A_{1,0} & \xrightleftharpoons[k_{aa}^*]{k_{aa}} C^A_{2,1} & C_{1,1} + C^A_{1,0} & \xrightleftharpoons[k_{ba}^*]{k_{ba}} C^B_{2,1} \\
C_{1,1} + C^B_{0,1} & \xrightleftharpoons[k_{ab}^*]{k_{ab}} C^B_{1,2} & C_{1,1} + C^B_{0,1} & \xrightleftharpoons[k_{ba}^*]{k_{ba}} C^B_{1,2}.
\end{align*}
\]

Again, it is convenient to define the following direct reaction rates for these steps, $k_{ha} = \frac{k_{aa} + k_{ab}}{2}$, $k_{hb} = \frac{k_{ba} + k_{bb}}{2}$ and inverse $k_{ha}^* = \frac{k_{aa}^* + k_{ab}^*}{2}$, $k_{hb}^* = \frac{k_{ba}^* + k_{bb}^*}{2}$. Note that if $k_{aa} = k_{bb}$ and $k_{ab} = k_{ba}$, then $k_{ha} = k_{hb}$, and if $k_{aa}^* = k_{bb}^*$ and $k_{ab}^* = k_{ba}^*$, then $k_{ha}^* = k_{hb}^*$. The reactions to consider are then:

\[
\begin{align*}
C_{1,1} + C^A_{1,0} & \xrightleftharpoons[k_{ha}^*]{k_{ha}} C^A_{2,1} & C_{1,1} + C^B_{0,1} & \xrightleftharpoons[k_{hb}^*]{k_{hb}} C^B_{1,2}.
\end{align*}
\]

(9)

As we have already remarked, in our model, as in the original one for open systems \[26\], the polymeric chains that have taken up the "wrong" chirality monomer can continue to grow. Thus, we allow for the further growth of these chains by adding monomers of either chirality. This kind of polymerization reaction for $2 \leq n \leq N - 2$ is given by:

\[
\begin{align*}
C^A_{1,n} + C^A_{1,0} & \xrightleftharpoons[k_{aa}^*]{k_{aa}} C^A_{2,n} & C^B_{n,1} + C^B_{0,1} & \xrightleftharpoons[k_{bb}^*]{k_{bb}} C^B_{n,2} \\
C^A_{1,n} + C^B_{0,1} & \xrightleftharpoons[k_{ab}^*]{k_{ab}} C^B_{1,n+1} & C^A_{n+1,0} + C^A_{1,0} & \xrightleftharpoons[k_{ba}^*]{k_{ba}} C^A_{n+1,1}.
\end{align*}
\]

(10)

(11)

And for $2 \leq r \leq N - 2$, $1 \leq s \leq N - 1 - r$:

\[
\begin{align*}
C^A_{r,s} + C^A_{1,0} & \xrightleftharpoons[k_{aa}^*]{k_{aa}} C^A_{r+1,s} & C^B_{r,s} + C^B_{0,1} & \xrightleftharpoons[k_{bb}^*]{k_{bb}} C^B_{r+1,s} \\
C^A_{r,s} + C^B_{0,1} & \xrightleftharpoons[k_{ab}^*]{k_{ab}} C^B_{r,s+1} & C^A_{r,s+1} + C^A_{1,0} & \xrightleftharpoons[k_{ba}^*]{k_{ba}} C^B_{r,s+1}.
\end{align*}
\]

(12)

(13)

Note that in the elementary reaction steps, in the rate constants, and in the corresponding differential rate equations (see below), the left-right symmetry of the model is manifest, that is, possesses a discrete $Z_2$ symmetry. This symmetry can be broken spontaneously by the dynamical solutions of the differential rate equations, thus this model is apt for studying spontaneous mirror symmetry breaking.

By lifting the $Z_2$ degeneracy in the reaction rates, e.g., allowing for $k_{aa} \neq k_{bb}$ and thus leading to more independent rate constants for describing the reaction set, we could study the influence of explicit chiral bias in the model. As this is not the aim of this work, we will not consider it here.
We next write down the differential rate equations corresponding to this reaction network. We employ the rate-equation theory as in chemical kinetics. We begin with the rate equations for the chiral monomers:

\[
\frac{dc^A_{1,0}}{dt} = -k_{aa}c^A_{1,0} \left(2c^A_{1,0} + \sum_{n=2}^{N-1} N-1 \sum_{n=2}^{N-2} c^A_{1,n} + \sum_{r=2}^{N-2} \sum_{s=1}^{N-1-r} c^A_{r,s} \right) - k_{ba}c^A_{1,0} \left( \sum_{n=2}^{N-2} c^B_{n,0} + \sum_{n=2}^{N-2} c^B_{n,1} + \sum_{s=2}^{N-2} \sum_{r=1}^{N-1-s} c^B_{r,s} \right)
\]

\[
- k_{bb}c^A_{1,0}c^B_{1,1} - k_{ba}c^A_{1,0}c^B_{1,1} + k_{aa}^* \left(2c^A_{1,0} + \sum_{n=3}^{N} c^A_{n,0} + \sum_{n=2}^{N-2} c^A_{2,n} + \sum_{s=1}^{N-1} \sum_{r=1}^{N-1-r} c^A_{r,s} \right)
\]

\[
+ k_{ba}^* \left( \sum_{n=2}^{N-1} c^A_{1,n} + \sum_{n=2}^{N-2} c^A_{2,n} + \sum_{s=3}^{N-1} \sum_{r=1}^{N-1-s} c^A_{r,s} \right) + k_{ha}^* c_{1,1} + k_{ha}^* c^A_{1,1}
\]

(14)

\[
\frac{dc^B_{0,1}}{dt} = -k_{bb}c^B_{0,1} \left(2c^B_{0,1} + \sum_{n=2}^{N-1} c^B_{n,0} + \sum_{n=2}^{N-2} c^B_{n,1} + \sum_{s=2}^{N-2} \sum_{r=1}^{N-1-s} c^B_{r,s} \right) - k_{ab}c^B_{0,1} \left( \sum_{n=2}^{N-1} c^A_{n,0} + \sum_{n=2}^{N-2} c^A_{n,1} + \sum_{s=2}^{N-2} \sum_{r=1}^{N-1-s} c^A_{r,s} \right)
\]

\[
- k_{bb}c^A_{1,0}c^B_{1,1} - k_{ba}c^A_{1,0}c^B_{1,1} + k_{bb}^* \left(2c^B_{0,1} + \sum_{n=3}^{N} c^B_{n,0} + \sum_{n=2}^{N-2} c^B_{n,2} + \sum_{s=3}^{N-1} \sum_{r=1}^{N-1-s} c^B_{r,s} \right)
\]

\[
+ k_{ab}^* \left( \sum_{n=2}^{N-1} c^B_{n,1} + \sum_{n=2}^{N-2} c^B_{n,2} + \sum_{s=3}^{N-1} \sum_{r=1}^{N-1-s} c^B_{r,s} \right) + k_{ba}^* c^B_{1,1}
\]

(15)

The equations describing the concentration of the homopolymers, for \(2 \leq n \leq N - 1\):

\[
\frac{dc^A_{n,0}}{dt} = k_{aa}c^A_{n,0} \left(c^A_{n-1,0} - c^A_{n,0} \right) - k_{ab}c^A_{n,0}c^B_{0,1} + k_{aa}^* \left(c^A_{n+1,0} - c^A_{n,0} \right) + k_{ab}^* c^A_{n,1}
\]

(16)

\[
\frac{dc^B_{n,0}}{dt} = k_{bb}c^B_{0,1} \left(c^B_{n-1,0} - c^B_{0,n} \right) - k_{ba}c^B_{0,1}c^A_{1,0} + k_{bb}^* \left(c^B_{n+1,0} - c^B_{0,n} \right) + k_{ba}^* c^B_{1,1}
\]

(17)

It is necessary to treat the kinetic equations of the maximum length homopolymers \(N\) individually. Since these do not elongate further, they can not directly react, and can not be the product of an inverse reaction involving a longer chain:

\[
\frac{dc^A_{N,0}}{dt} = k_{aa}c^A_{1,0}c^A_{N-1,0} - k_{aa}^* c^A_{N,0}
\]

(18)

\[
\frac{dc^B_{N,0}}{dt} = k_{bb}c^B_{0,1}c^B_{N-1,0} - k_{bb}^* c^B_{N,0}
\]

(19)

The differential equations describing the concentration of each type of heteropolymer (included the heterodimer), for \(2 \leq n \leq N - 2\):

\[
\frac{dc^A_{1,n}}{dt} = k_{h}c^A_{1,0}c^B_{n,1} - k_{ha}c^A_{1,1}c^B_{n,0} - k_{hb}c^A_{1,1}c^B_{1,0} - k_{hh}c^A_{1,1} + k_{ha}^* c^A_{2,1} + k_{hb}^* c^B_{1,2}
\]

(20)

\[
\frac{dc^B_{1,n}}{dt} = -k_{aa}c^A_{1,0}c^B_{1,0} - k_{ab}c^A_{1,0}c^B_{1,1} + k_{ba}c^B_{0,1}c^A_{1,0} + k_{aa}^* c^A_{2,1} + k_{ab}^* c^B_{1,2}
\]

(21)

\[
\frac{dc^A_{n,1}}{dt} = -k_{bb}c^B_{n,0}c^B_{1,1} + k_{ab}c^A_{n,0}c^B_{1,1} + k_{bb}^* c^B_{n,1} + k_{ab}^* c^A_{n+1,1} - k_{ab}^* c^B_{n,1}
\]

(22)

As before, it is useful to treat individually the maximum length polymers \(N\):
\[ \frac{dc_{1,N-1}^A}{dt} = k_{ba}c_{0,N-1}^Ac_{1,0}^A - k_{ba}c_{1,N-1}^A \]
\[ \frac{dc_{N-1,1}^B}{dt} = k_{ab}c_{N-1,0}^Ac_{1,0}^B - k_{ab}c_{N-1,1}^B \]

(23)  
(24)

As was mentioned when describing the reaction network, each kind of trimer \( c_{2,1}^A \) and \( c_{1,2}^B \) must have its own differential equation in terms of \( k_{ba}, k_{bb} \):

\[ \frac{dc_{1,2}^A}{dt} = -k_{aa}c_{1,0}^Ac_{2,1}^A - k_{ba}c_{0,1}^A + k_{ba}c_{1,1}c_{0,1} + k_{aa}c_{3,1}^A + k_{bb}c_{2,2}^A - k_{ba}c_{2,1}^A \]
\[ \frac{dc_{1,2}^B}{dt} = -k_{bb}c_{0,1}^B - k_{ba}c_{1,0}^B + k_{ba}c_{0,1}c_{1,2}^B + k_{bb}c_{1,1}c_{0,1}^B + k_{bb}c_{2,2}^B - k_{bb}c_{1,2}^B \]

(25)  
(26)

For \( 2 \leq n \leq N - 3 \):

\[ \frac{dc_{2,n}^A}{dt} = k_{aa}c_{1,0}^Ac_{2,n}^A - k_{ab}c_{0,1}^A + k_{ba}c_{1,n}^A + k_{aa}c_{3,n}^A + k_{bb}c_{2,n+1}^A - k_{ba}c_{2,n}^A \]
\[ \frac{dc_{n,2}^B}{dt} = k_{bb}c_{0,1}^B + k_{ba}c_{1,0}^B + k_{ab}c_{n,1}^B + k_{bb}c_{1,n}^B + k_{bb}c_{2,n+1}^B - k_{bb}c_{n,2}^B \]

(27)  
(28)

Once again, the equations corresponding to the maximum length homopolymers \( N \) are

\[ \frac{dc_{2,N-2}^A}{dt} = k_{aa}c_{1,0}^Ac_{2,N-2}^A + k_{ba}c_{1,N-2}^A - k_{aa}c_{2,N-2}^A - k_{ba}c_{2,N-2}^A \]
\[ \frac{dc_{N-2,2}^B}{dt} = k_{bb}c_{0,1}^B + k_{ba}c_{N-2,1}^B - k_{bb}c_{N-2,2}^B - k_{ab}c_{N-2,2}^B \]

(29)  
(30)

For \( 3 \leq r \leq N - 2 \) and \( 1 \leq s \leq N - 1 - r \):

\[ \frac{dc_{r,s}^A}{dt} = k_{aa}c_{1,0}^Ac_{r,s}^A - k_{ab}c_{0,1}^A + k_{ba}c_{r-1,s}^A + k_{aa}c_{r+1,s}^A - k_{aa}c_{r,s}^A \]
\[ \frac{dc_{r,s}^B}{dt} = k_{bb}c_{0,1}^B + k_{ba}c_{r-1,s}^B + k_{bb}c_{r,s+1}^B + k_{ba}c_{r,s+1}^A - k_{bb}c_{r,s}^A \]

(31)  
(32)

(33)

For \( 3 \leq s \leq N - 2 \) and \( 1 \leq r \leq N - 1 - s \):

\[ \frac{dc_{r,s}^A}{dt} = k_{aa}c_{1,0}^Ac_{r,s}^A - k_{ab}c_{0,1}^B + k_{ba}c_{r,s-1}^A + k_{aa}c_{r,s+1}^A + k_{bb}c_{r,s}^A \]
\[ \frac{dc_{r,s}^B}{dt} = k_{bb}c_{0,1}^B + k_{ba}c_{r,s-1}^B + k_{bb}c_{r,s+1}^B + k_{ba}c_{r,s+1}^B - k_{bb}c_{r,s}^B \]

(34)  
(35)
TABLE I. Number of differential equations as a function of the maximum polymer length $N$

| $c_{1,0}^A$ | Number of eqs | $c_{0,1}^B$ | Number of eqs |
|--------------|---------------|--------------|---------------|
| $c_{1,1}$    | 1             | $c_{6,1}^B$  | 1             |
| $c_{n,0}^A$, $(2 \leq n \leq N)$ | $\sum_{n=2}^{N} n = N - 1$ | $c_{0,n}^B$, $(2 \leq n \leq N)$ | $\sum_{n=2}^{N} n = N - 1$ |
| $c_{1,n}^A$, $(2 \leq n \leq N - 1)$ | $\sum_{n=2}^{N-1} n = N - 2$ | $c_{n,1}^B$, $(2 \leq n \leq N - 1)$ | $\sum_{n=2}^{N-1} n = N - 2$ |
| $c_{2,1}^A$ | 1             | $c_{1,2}^B$  | 1             |
| $c_{n,2}^A$, $(2 \leq n \leq N - 2)$ | $\sum_{n=2}^{N-2} n = N - 3$ | $c_{n,2}^B$, $(2 \leq n \leq N - 2)$ | $\sum_{n=2}^{N-2} n = N - 3$ |
| $c_{r,s}^A$ | $(3 \leq r \leq N - 2)$ | $c_{r,s}^B$ | $(3 \leq s \leq N - 2)$ |
| $(1 \leq s \leq N - 1 - r)$ | $\sum_{r=3}^{N-2} \sum_{s=1}^{N-1-r} = \frac{1}{2}(N^2 - 7N + 12)$ | $\sum_{r=1}^{N-1-s} \sum_{s=3}^{N-2} = \frac{1}{2}(N^2 - 7N + 12)$ |
| $c_{n,N-n}^A$, $(3 \leq n \leq N - 1)$ | $\sum_{n=3}^{N-1} n = N - 3$ | $c_{r,s}^B$, $(3 \leq n \leq N - 1)$ | $\sum_{n=3}^{N-1} n = N - 3$ |

As remarked earlier, the complete reaction scheme must satisfy mass conservation in a closed system, implying that the mass variation rate must be strictly zero:

$$0 = 2c_{1,1} + 3(c_{2,1}^A + c_{1,2}^B) + \sum_{n=1}^{N} n(c_{n,0}^A + c_{0,n}^B) + \sum_{n=2}^{N-1} (n + 1)(c_{1,n}^A + c_{n,1}^B)$$

$$+ \sum_{n=2}^{N-2} (n + 2)(c_{n,2}^A + c_{n,2}^B) + \sum_{r=3}^{N-1-s} \sum_{s=1}^{N-1-r} (r + s)(c_{r,s}^A + c_{r,s}^B),$$

(36)

where the overdot stands for the time-derivative. The compliance with this constraint is an important and crucial check on the consistency of the numerical integration of the full set of differential equations Eqs. (36), which we monitor and confirm in all the simulations presented below. Analytically, this relation is satisfied by the rate equations.

As we see, there is one differential equation for each type of monomer and one for the heterodimer. The homopolymer set requires $\sum_{n=2}^{N} 2(N - 1)$ equations and the heteropolymer set a total of $2(N - 2)$ equations. The total number of kinetic differential equations describing the whole system is $N(N + 1)$, and is broken down into the separate contributions as displayed in Table I. Then, the total number of equations for describing the system as a function of maximum chain length $N$ is:

$$\#eqs = 6 + 2(N - 1) + 2(N - 2) + 2(N - 3) + (N^2 - 7N + 12) + 2(N - 3) = N(N + 1),$$

(37)

as pointed out in Ref [27]. From the computational point of view, the number of equations grows quadratically with the maximum chain length $N$.

### III. NUMERICAL RESULTS

We are interested in applying our copolymerization model to fit the experimental data measured by the Rehovot group, so our primary goal is to reproduce as closely as possible the details reported concerning the experiments on chiral amplification of oligopeptides. For this purpose, the first step is to determine the initial monomer concentrations to be employed in the simulations. The actual experiments were carried out for 0.5$mM$ solutions of monomers, thus we have employed for each case: (a) $R : S = 1 : 1$ which corresponds to an initial enantiomeric excess $ee_0 = 0\%$, so $c_{1,0}^A(0) = 0.25mM$ and $c_{0,1}^B(0) = 0.25mM$; (b) $R : S = 4 : 6$ corresponding to $ee_0 = 20\%$, so $c_{1,0}^A(0) = 0.2mM$ and $c_{0,1}^B(0) = 0.3mM$; (c) $R : S = 3 : 7$ which corresponds to $ee_0 = 40\%$, so $c_{1,0}^A(0) = 0.15mM$ and $c_{0,1}^B(0) = 0.35mM$. The remainder of the initial concentrations (the dimers and on up) are taken to be zero. Next, we systematically search for the reaction rates leading to the best fit to the given data.

Different chemical model systems were used in the experiments: namely $\gamma$-stearyl-glutamic thioethyl ester ($C_{18} - TE - Glu$), $N^\prime$-stearyl-lysine thioethyl ester ($C_{18} - TE - Lyg$), $\gamma$-stearyl-glutamic acid N-carboxyanhydride ($C_{18} - Glu - NCA$) and $\gamma$-stearyl-glutamic thioacid ($C_{18} - thio - Glu$), varying both their initial compositions and for
The composition of the oligopeptides formed was analyzed by matrix-assisted laser desorption/ionization time-of-flight mass spectroscopy (MALDI-TOF) with enantio-labeled samples. The experimental relative abundances of the oligopeptides was inferred from the ion intensity. It are these relative abundances that we aim to interpret vis-a-vis our copolymerization model.

Since only the experiments with racemic mixtures of the starting compounds required a catalyst, it is reasonable to expect that the racemic and the chiral enriched cases will follow different dynamics for a given model system. That is, the presence or absence of a specific catalyst affects the rate constants, for a given chemical system. Firstly, we will find the reaction rates for the racemic case, and afterwards, those for the enriched chiral case, allowing us to compare both. The a-priori nine free parameters we must set to run the numerical integrations are comprised by the four direct and the four inverse rate constants \( k_{aa}, k_{bb}, k_{ab}, k_{ba}, k_{aa}^*, k_{bb}^*, k_{ab}^*, k_{ba}^* \), plus the maximum polymer chain length, \( N \). We set all the inverse reaction rates to a unique value, \( k_{aa}^* = k_{bb}^* = k_{ab}^* = k_{ba}^* = 10^{-10}(s^{-1}) \), implying an almost irreversible scheme, and we determine the remainder of the parameters from fitting the copolymerization model to the relative abundance data. This required numerical integration of the set of differential equations Eqs. (14-35) which we performed using the Mathematica program package. For each independent run we verified the compliance of the numerical results with the constraint in Eq.(36), an imperative for any closed system.

Results from fitting the model to the data indicate that the maximum chain length \( N \) does not play a significant role, the Pearson product-moment correlation coefficient, \( r \), remains the same for \( N = 12, 14, 16, 18, 20 \), so we will set \( N = 12 \) for all compounds and cases treated below. Since the number of independent equations scales as \( N^2 \), this represents an important reduction on computer time and the memory used. We note that one is free to scale out the dependence of one pair of reaction constants from the rate equations by a suitable redefinition of the time variable. Thus, without loss of generality, we set the cross inhibition rates equal to unity \( k_{ab} = k_{ba} = 1(s^{-1}mol^{-1}) \) and then search for the reaction rates \( k_{aa} = k_{bb} \) leading to the best fits.

A. Racemic mixtures

In one set of experiments, the authors reported MALDI-TOF analysis of the oligopeptides formed at the air-water interface from racemic mixtures \( R : S = 1 : 1 \) of the monomers for the various model systems and catalysts. We first fit the copolymerization model to this data.

The best correlation data for the racemic \( C_{18} \text{-TE-Glu} \) system, with the \( I_2/KI \) catalyst are found for \( k_{aa} = k_{bb} = 1.7(s^{-1}mol^{-1}) \). In this case, the best fit obtains for the time scale \( t = 10^{11}(s) \). Exactly by the same process, the best correlation data for the racemic \( C_{18} \text{-TE-Lys} \) are found for \( k_{aa} = k_{bb} = 2.3(s^{-1}mol^{-1}) \) and for \( k_{aa} = k_{bb} = 1.3(s^{-1}mol^{-1}) \) when adding \( I_2/KI \) and \( AgNO_3 \) as catalyst, respectively. For the simulations here, we took the times \( t = 10^{10}(s) \) and \( t = 10^{11}(s) \) in the racemic cases with \( I_2/KI \) and \( AgNO_3 \) respectively. Finally, we fit our copolymerization model to the \( C_{18} \text{-thio-Glu} \) experimental relative abundances. The authors of the experiments affirmed that this compound undergoes a truly random polymerization, so fits from our model are expected to be slightly less satisfactory than those for the binomial distribution function. Setting the inverse reaction rates and the cross inhibition as indicated above, then the best correlation coefficients are found for \( k_{aa} = k_{bb} = 0.4(s^{-1}mol^{-1}) \). The instant or time-scale leading to these numerical values is \( t = 10^{10}(s) \).

The corresponding (experimental and numerical) relative abundances for the four compounds cited above corresponding to these values are shown in Fig. 2. The histograms show the relative abundance of each experimentally obtained oligopeptide compared to the best fit from our copolymerization model. We emphasize that we fit the model to the complete family of stereoisomer subgroups (global fit). The resulting data correlations are shown in Fig. 3 and Table IV the latter gives a detailed comparison of the best fits between individual subfamilies and the overall global fit.

In the case of the \( C_{18} \text{-Glu-NCA} \) with catalyst \( Ni(C_2H_3CO_2)_2 \), the best fit is obtained for \( k_{aa} = k_{bb} = 0.2(s^{-1}mol^{-1}) \). Results for the corresponding relative abundances are shown in Fig. 4 and the correlation from fitting is displayed in the bottom frame of Fig 3 and Table IV. Not all subfamily data sets are reported in the experimental paper [24]; here we use the fitted model to the partial data set to predict or fill in this missing subfamily data. Numerical results for the racemic case have been found for \( t = 10^{10}(s) \).

B. Chirally enriched mixtures

In a second set of experiments, the authors reported MALDI-TOF analysis of the oligopeptides formed at the air-water interface from non-racemic mixtures of the monomers for the same model systems. No catalysts were employed
FIG. 2. Relative abundance versus number of repeat units \((r, s)\) of the oligopeptides obtained from fitting the model (white) to the experimental data (black) from racemic mixtures \(R : S = 1 : 1\) of monomers. The four chemical models are indicated by the insets.

TABLE II. Comparative fits between the copolymerization model and the binomial distribution to the experimental relative abundances: racemic mixtures \(R : S = 1 : 1\) of monomers of the four model systems as indicated in the leftmost column. Only in the case of \(C_{18} − \text{thio} − \text{Glu}\) does the binomial distribution give a better global fit than the copolymerization model: this latter system provides an experimental reference system for random polymerization [25].

|          | Copolymerization model | Bin. |
|----------|------------------------|------|
|          | Fits for each subgroup |      |
|          | di | tri | tetra | penta | hexa | fit | fit |
| \(C_{18} − TE − Glu\) | 0.92 | 0.96 | 0.80 | 0.84 | -   | 0.93 | 0.75 |
| \(C_{18} − TE − Lys(I_{2}/KI)\) | 0.96 | 0.82 | 0.11 | -0.73 | 0.45 | 0.85 | 0.32 |
| \(C_{18} − TE − Lys(Ag)\) | 0.98 | 1    | 0.03 | 0.88 | 0.76 | 0.84 | 0.8  |
| \(C_{18} − \text{thio} − Glu\) | 1   | 1    | 1    | 0.98 | 0.97 | 0.95 | 0.98 |

there. We next consider fits of our model to these data sets.

The best correlations factors for both chirally enriched mixture cases (20% and 40% excesses) in the case of the \(C_{18} − TE − Glu\) system are found for the same rates, that is for \(k_{aa} = k_{bb} = 2\,(s^{-1}mol^{-1})\). The results for these
values are shown in Table IV. In Fig. 4 we display the relative abundances of the homochiral oligopeptides and in Table V both the calculated and experimental enantiomeric excesses for the 4:6 and 3:7 (R:S) mixtures. In Fig. 5 we show the data correlation. Numerical results for the non-racemic case have been found for the time scale $t = 10^{11}$ (s).

For the chiral mixtures of $C_{18} - TE - Lys$ we found the best fits for the dynamics corresponding to $k_{aa} = k_{bb} = 2.5 (s^{-1} mol^{-1})$. The results for these values are shown in Table VI. The relative abundances results for these values
FIG. 4. The $C_{18} - Glu - NCA$ system with catalyst $Ni(CH_3CO_2)_2$: relative abundance of the oligopeptides obtained from fitting the model (white) to the experimental data (black) from racemic mixtures of monomers. Compare to Fig. 4A of reference[25]

TABLE III. Comparative fits between the copolymerization model and binomial distribution to the experimental relative abundances for the racemic compositions (R:S=1:1) of the $C_{18} - Glu - NCA$ system.

|          | Copolymerization model | Bin. |
|----------|------------------------|------|
|          | Fits for each subgroup n | Global | Global |
| $C_{18} - Glu - NCA$ | di | tri | tetra | penta | hexa | hepta | octa | nona | deca | endeca | dodeca | fit | fit |
|          | 1 | - | 1 | - | 0.98 | - | 0.98 | - | 0.97 | 1 | 0.95 | 0.96 | 0.75 |

TABLE IV. Comparative fits between the copolymerization model and the binomial distribution to the experimental relative abundances measured for non-racemic mixtures of $C_{18} - TE - Glu$.

|          | Copolymerization model | Binomial |
|----------|------------------------|----------|
|          | Fits for each subgroup n | Global | Global |
| (R:S) 4:6 | di | tri | tetra | penta | hexa | fit | fit |
| 0.86 | 0.89 | 0.93 | 0.99 | - | 0.94 | 0.75 |
| (R:S) 3:7 | 0.95 | 0.94 | 0.96 | 0.99 | 0.99 | 0.95 | 0.85 |

TABLE V. Enantiomeric excesses $ee$: numerical results from the copolymerization model (experimental data) for the relative abundances of the homochiral oligopeptides for the $C_{18} - TE - Glu$ system.

|          | di | tri | tetra | penta | hexa |
|----------|----|----|------|------|-----|
| (R:S) 4:6 | 18 (26) | 24 (39) | 30 (46) | 35 (59) | - |
| (R:S) 3:7 | 37 (48) | 48 (71) | 57 (82) | 66 (92) | 73 (>99.8) |

are shown in Fig. 7 and the enantiomeric excesses obtained for 4:6 and 3:7 (R:S) mixtures are presented in Table VII. In Fig. 8, the data correlation is shown. For the simulations here, we took the instants $t = 10^{10}(s)$ and $t = 10^{11}(s)$ in the racemic cases with $I_2/KI$ and $AgNO_3$ respectively, and $t = 10^{10}(s)$ for the chirally enriched mixtures.

In the case of nonracemic $C_{18} - thio - Glu$, the best correlation coefficients are found for the same values of the reaction rates that we found in the racemic case, namely for $k_{aa} = k_{bb} = 0.4(s^{-1}mol^{-1})$. Results for the chiral cases are shown in Table VIII. As to be expected and as shown there, the correlation factors for the global fit to the binomial distribution function are slightly better than those for any simulation we could perform with the copolymerization model, so we reconfirm what was claimed by the authors of the experimental work: namely that the $C_{18} - thio - Glu$
FIG. 5. Relative abundance versus number of repeat units \((r, s)\) of the oligopeptides obtained from fitting the model (white) to the experimental data (black) from non-racemic mixtures of monomers for the \(C_{18} - TE - Glu\) system.

FIG. 6. Results from fitting the model to the experimental data: non-racemic mixtures of \(C_{18} - TE - Glu\). The solid line represents the linear correlation between experimental and numerical data obtained from fitting. The dotted line has slope equal to unity.

TABLE VI. Results for the copolymerization model and experimental data correlations for non-racemic mixtures of \(C_{18} - TE - Lys\).

| \((R:S)\) | 4:6 | Binomial |
|---|---|---|
| \(r\) | Fits for each subgroup | Global | Global |
| di | tri | tetra | penta | hexa | hepta | fit | fit |
| (R:S) 4:6 | 0.78 | 1 | 0.87 | 0.90 | 0.84 | 0.97 | 0.89 | 0.46 |
| (R:S) 3:7 | 0.93 | 1 | 0.95 | 0.97 | 0.99 | - | 0.94 | 0.65 |

system polymerizes randomly. In Figure 5 the relative abundances of the oligopeptides are shown. The data correlation is shown in Figure 6.

The best fits for both chirally enriched mixture cases (20% and 40% excesses) in the case of \(C_{18} - Glu - NCA\) are found for the same dynamics, that is \(k_{aa} = k_{bb} = 3.8(s^{-1}mol^{-1})\). The results for these values are shown in Table VI.
TABLE VII. Enantiomeric excesses: numerical results from the copolymerization model (experimental data) for the relative abundances of the homochiral oligopeptides for $C_{18} - TE - Lys$.

| ee(%) | di | tri | tetra | penta | hexa | hepta |
|-------|----|-----|-------|-------|------|-------|
| (R:S) 4:6  | 23 (34) | 30 (34) | 36 (41) | 42 (60) | 49 (62) | 54 (>99.8) |
| (R:S) 3:7  | 45 (46) | 57 (63) | 66 (73) | 75 (85) | 81 (86) | - |

FIG. 7. Relative abundance versus number of repeat units ($r, s$) of the oligopeptides obtained from fitting the model (white) to the experimental data (black) from non-racemic mixtures of monomers of $C_{18} - TE - Lys$.

FIG. 8. Results from fitting the model to the experimental data. Chiral mixtures of $C_{18} - TE - Lys$. The solid line represents the linear correlation between experimental and numerical data obtained from fitting. The dotted line has slope equal to unity.

In Fig.11 we compare the best fit against the experimentally obtained relative abundances of the oligopeptides. The corresponding data correlation is shown in Fig.12. Numerical results for the racemic case have been found for $t = 10^{11}$(s).
TABLE VIII. Results for the copolymerization model and experimental data correlations for non-racemic mixtures of $C_{18}$–thio–Glu.

| (R:S) | Fits for each subgroup n | Global fit | Global fit |
|-------|--------------------------|------------|------------|
| 4:6   | r                       |            |            |
|       | di 0.93 | tri 0.98 | tetra 0.93 | penta 0.92 | hexa 0.92 | hepta 0.91 | fit 0.91 | fit 0.93 |
| 3:7   | r       | 0.89      | 1          | 0.99      | 0.99      | 0.98      | -        | 0.96      | 0.97      |

FIG. 9. Relative abundances versus number of repeat units ($r, s$) of the oligopeptides obtained from fitting the model (white) to the experimental data (black) for the non-racemic mixtures of $C_{18}$–thio–Glu.

FIG. 10. Results from fitting the model to the experimental data for non-racemic mixtures of the $C_{18}$–thio–Glu system. The solid line represents the linear correlation between experimental and numerical data obtained from fitting. The dotted line has slope equal to unity.

IV. CONCLUSIONS

The overall scheme for the chiral amplification process leading to the experimental data investigated here involves a self-assembly step followed by a lattice-controlled polymerization [24, 25]. It is this subsequent polymerization which is the prime focus of this paper. The authors of the experimental work stress that it is not at all straightforward
TABLE IX. Results for the copolymerization model and experimental data correlations for $C_{18} - Glu - NCA$. The global fit from the binomial distribution is shown for comparison.

|                  | Copolymerization model | Binomial |                  |                  |                  |                  |                  |                  |                  |                  |
|------------------|------------------------|----------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
|                  |                      | Fits for each subgroup n |                  | Global        |                  |                  |                  |                  |                  |                  |
|                  |                      | di       | tri              | tetra             | penta          | hexa             | hepta            | octa             | nona            | deca             |
| (R:S)            |                      | fit      |                  |                  |                  |                  |                  |                  |                  |                  |
| 4:6              | -0.79                 | 0.9      | 0.63             | 0.74              | 0.95            | 0.89             | 0.77             | 0.86             | -               | 0.68             | 0.11             |
| 3:7              | -0.33                 | 0.79     | 0.81             | 0.76              | 0.86            | 0.96             | 0.75             | 0.83             | 0.89            | 0.75             | 0.38             |

FIG. 11. Relative abundances for the non racemic mixtures of $C_{18} - thio - Glu$. The experimental data set (black) is incomplete, we have used our model to fill in the missing portions of the histogram (white)

FIG. 12. Results from fitting the model to the experimental data for the $C_{18} - Glu - NCA$ system. The solid line represents the linear correlation between experimental and numerical data obtained from fitting. The dotted line has slope equal to unity.
to actually establish the correlation between the packing arrangement of the crystallites and the composition of the diastereoisomeric products that result therefrom. Therefore, our task here was to fit the outcome of these latter steps assuming an effective copolymerization scheme. The term “effective” simply means that the putative complicated correlations and interplay between the 2D crystallite phases at the air-water interface and the polymerization reaction pathways that depend on the microscopic packing arrangements within the crystals are treated here with a simple model. In this regard, our model can be regarded as a “course-grained” description of the overall process in that the microscopic details (the structures of the crystalline phases) are not resolved, but that the end-result or net effect of the pathways afforded by the crystallites can be summarized by the polymerization scheme as depicted graphically in Fig. 1.

The model as introduced is defined for fully reversible reactions and this implies that some of the reaction rates must obey a corresponding constraint as dictated by microreversibility. Thus the model is appropriate for closed systems under thermodynamic control. For the numerical fits themselves, we found that all the reverse reaction rates could be set to rather tiny values, and this in consonance with experimentally observed irreversible condensation. Thus for the present purposes, the copolymerization model is practically irreversible. The values for the forward rates of adding the same chirality monomer to the end of the growing chain are found to be greater than those for addition of a wrong chirality monomer: that is, \( k_{aa} = k_{bb} > k_{ab} = k_{ba} = 1 \) (except of course for the model system \( C_{18}^{-\text{thio}} - \text{Glu} \) serving as reference for random polymerization).

Other closed systems that lead to copolymers could be in principle be fit with our model. If for example \( k_{aa} \) and \( k_{bb} \) had different magnitudes, this would imply that a underlying chiral bias is operative either in the polymerization or in the prior formation of the two crystallites that control the polymerization. This bias could affect the packing arrangements of the crystal monomers and the reaction pathways taken within each crystallite phase. Since however our model is effective, as explained earlier, we would not be able to say whether the chiral bias is in the polymerization or in the structure of the crystallites that control the polymerization. Nevertheless, this bias in \( k_{aa} \) being different from \( k_{bb} \), would result in favoring the attachment of say, an \( S \) to an \( S \) over the attachment of an \( R \) to an \( R \), and this feature would show clearly up in the relative abundances.

Another positive feature of the model is the robustness of the fits with respect to differing initial imbalances of the enantiomers. That is, for a given chemical model (including catalyst, if any) the values of the fitted rates do not depend on the initial enantiomeric excesses of the monomers. If our rate constants are viewed as effective, that is, implicitly involving the different chemical properties of the racemic and enantiomorphous crystallite phases, then this feature suggests that the packing arrangements and reaction pathways in the solid-state do not depend (or only weakly) on the magnitude of these imbalances.

The Pearson product-moment correlation coefficient \( r \) between experimental and numerical data is greater for the copolymerization model than for the binomial distribution, except for the \( C_{18}^{-\text{thio}} - \text{Glu} \), which truly polymerizes randomly. The correlation between calculated and experimental relative abundances is also greater for the initially non-racemic situations, and the higher the initially chiral enrichment of the mixture is, the better the copolymerization model reproduces the chemical data. The results obtained here lead us to affirm that the model systems considered all undergo a non-random polymerization, as was asserted by the authors of the experiments [24, 25].

The model also qualitatively reproduces the behavior of the enantiomeric excess \( ee \), its increase with the length of the chains and the enhancement of the \( ee \) of the corresponding initial mixture of monomers. All this, in spite of the complexity of the factors that affect the reactivity within the experimental two-phase system, i.e., the microscopic crystallite packing arrangements and the possible reaction pathways within these 2D crystallites. In conclusion then, we may therefore assert that our simple scheme does provide an accurate course-grained description of the lattice-controlled polymerization reported in Ref [24, 25].

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