# On the chemical freezing of phase separation in immiscible binary mixtures

Daniele Carati and René Lefever Faculté des Sciences, CP 231, Université Libre de Bruxelles 1050 Bruxelles, Belgium. (Dated: Submitted to PRE, February 1996, Revised February 1997)

We discuss the thermodynamic and kinetic conditions under which chemical reactions may prevent the coarsening terminating spinodal decomposition and freeze the unmixing of binary mixtures at some early, pattern forming, stage of evolution. Under very general conditions, we establish that: (i) this pattern freezing phenomenon can only occur in nonequilibrium systems the level of dissipation of which exceeds a finite, non zero threshold value; (ii) at least two independent chemical processes must take place; (iii) chemistry must be destabilizing, which requires that at least one of these processes must be autocatalytic; (iv) pattern formation is possible even outside of the spinodal region, i. e., without involving a phase separation phenomenon, in unsymmetrical mixtures where the potential energies between pairs of identical particles are sufficiently different. This latter condition replaces for non-ideal chemically reacting binary mixtures the *unequal diffusion coefficients condition* which governs the appearance of Turing patterns in the classical reaction-diffusion theory.

### I. INTRODUCTION

The phase ordering dynamics of a binary mixture, such as a binary alloy, quenched from a high-temperature, homogeneous state to a point below the spinodal line is wellknown [1]. In this domain, the mixture is unstable with respect to long wave-length, small amplitude concentration fluctuations and, at first, decomposes into two coexisting phases exhibiting an interconnected, labyrinthine morphology. Subsequently, on a long time scale, this pattern slowly coarsens and finally transforms into two homogeneous macroscopic phases separated by a minimal interfacial boundary region. This coarsening, also called Ostwald ripening, is due to the fact that in the spinodal domain, all nonzero Fourier modes with a wave number k smaller than some upper cutoff value  $k_u$  are unstable while the zero-mode, k = 0, is only marginally stable. Under those conditions, the dynamics selects a spatial organization which originates from the amplification of large wavelength modes, i. e., corresponds to vanishingly small wave numbers  $(k \to 0)$ . Indeed, though the growth of these large scale modes is extremely slow because it requires the diffusion of matter over large distances, in the end, it governs the spatial organization because it minimizes free energy.

Over the last twenty years, several theoretical studies have investigated the idea that coupling spinodal decomposition with chemical reactions may, in the absence of hydrodynamic effects, freeze this coarsening process and stabilize a stationary inhomogeneous state characterized by an intrinsic wavelength, i. e., a wavelength determined by molecular and kinetic parameters rather than by externally imposed boundary conditions and/or geometrical constraints [2–6, 8].

In general, the starting point of these theoretical studies is a nonlinear diffusion equation, deriving from a square-gradient free energy functional [9], in which source terms are added to model the chemical reactions occurring in the mixture. So far, the choice of these chemical

terms has relied either on intuitive considerations [2, 4– 6, 8], or on a master equation derivation which exploits the kinetic similarities of some chemical reactions with spin-exchange processes [3]. This basically mean field type of approach was pioneered by Huberman [2], who predicted that the influence of chemistry on spinodal decomposition may, not only narrow the band of unstable modes, but more importantly, introduce a lower cutoff,  $k_{\ell}$ , below which all modes, and in particular the mode k = 0, are stable. Considering a two step autocatalytic reaction, this author further showed, within the framework of this autocatalytic model, that chemistry restricts the size of inhomogeneities which arise during spinodal decomposition under conditions where the system is out of equilibrium with respect to both chemical equilibrium and phase composition.

The theoretical results reported in more recent works support this prediction that chemistry may suppress the coarsening which normally terminates the isothermal phase separation of immiscible, incompressible fluids. More than that even: in these studies, this suppression is obtained thanks to chemical processes which are considerably simpler than the one considered by Huberman, suggesting hence, that the phenomenon in question is ubiquitous in nature [5] and that the class of chemical reactions which qualify as candidates for a possible experimental demonstration, is extremely broad. A major simplification in this respect, is the fact that in none of the models studied recently, is the chemistry involved autocatalytic: typically, the reactions considered are simple isomerisation processes like  $A \rightleftharpoons B$ , [3, 4, 8], associationdissociations reactions like  $A + B \rightleftharpoons C$  [5], or standard adsorption-desorption surface reactions [6].

This apparently extremely general effect of chemistry on spinodal decomposition is however obtained through a procedure which we find thermodynamically unsatisfactory [10] because it consists in adopting chemical rate laws which, contrary to those adopted for diffusion, do not take into account the non-ideality of the systems considered. More precisely, it amounts to assuming that, somehow, far from thermodynamic equilibrium, the relationship binding chemical affinities to activity coefficients [11, 12] may be neglected and that, as a result, it then becomes possible to model chemical and diffusional processes as being independent from each other [13].

The conditions for which this far from equilibrium independency assumption (FFEIA) applies are unclear. Our objective here is therefore to investigate the behavior of mixtures undergoing phase separation in the presence of chemical reactions along a different line of approach which (i) does not rely on the FFEIA, (ii) allows us to encompass equilibrium as well as far from equilibrium situations and (iii) is based on thermodynamic considerations which are model independent.

We shall study the case of chemically reactive binary mixtures which is also the case investigated in previous works. We shall limit ourselves to situations which involve no hydrodynamic effects. From a chemical point of view, this means that we restrict ourselves to reaction schemes which *a priori* rule out the occurrence of pressure or density variations of chemical origin. No other assumption will be made concerning the kinetics of the chemical processes considered.

It is characteristic of chemical reactions and diffusion that the thermodynamic forces governing these processes are related, being function, in the first case, of the chemical potentials and, in the second case, of the chemical potentials spatial derivatives. The thermodynamic fluxes conjugate to these forces, i. e., the chemical reaction rates and diffusion rates, are thus also function of the chemical potentials; this relationship is furthermore well-known if one admits the validity of the usual phenomenological laws relating thermodynamic forces and fluxes. Under those conditions, choosing the chemical potentials determines the coupling between diffusional and chemical processes. In particular, it determines the equilibrium state towards which reaction-diffusion systems evolve in the absence of external constraints. As embodied in the classical Duhem-Jouget theorem [11], it results therefrom that in equilibrium systems, diffusional stability automatically insures the stability of chemical equilibrium. In order to determine what happens to this classical stability relationship under nonequilibrium conditions, question which underlies the present study, it is essential in modeling chemical and diffusional fluxes to respect this property that both kinds of fluxes are given functions of the same chemical potentials, which themselves are thermodynamic quantities subjected to well-defined, general requirements.

We undertake the investigation of the behavior of binary mixtures undergoing simultaneously spinodal decomposition and chemical reactions by addressing this problem in detail in section II. Taking as usual in mean field approaches, the square-gradient free energy as starting point, we discuss the form that chemical potentials must have within this framework to be thermodynamically consistent. Subsequently, we express in terms

of these chemical potentials the generalized reactiondiffusion equation governing the spatio-temporal behavior of the system. We analyze the linear stability properties of the uniform steady state solutions of this equation in section III. We derive from this analysis in section IV A and IV B, the thermodynamic and kinetic conditions under which chemical reactions may introduce a lower cutoff  $k_l$  which (i) excludes the zeroth mode from the band of unstable modes, (ii) permits to avoid the phenomenon of coarse graining and (iii) allows instead for the freezing of spinodal decomposition at an intermediary stage. This analysis allows us to predict also that the difference in potential energies between pairs of identical particles plays an essential role in the stability properties of the systems considered here. Notably, the value of this parameter determines whether or not there can already exist a finite band of unstable modes outside of the spinodal domain; in other words, whether or not a symmetry breaking instability may be observed which leads to pattern formation without involving the unmixing phase transition. The results are illustrated on an example in section IVC.

## II. MODELING OF CHEMICALLY REACTIVE BINARY MIXTURES

### A. Free Energy and chemical potentials

We consider a binary mixture subjected to isothermal, isobaric conditions which below (above) some critical temperature  $T_c$ , exhibits a miscibility gap. As usual, we suppose that the coarse grained concentrations of the components 1 and 2 forming this mixture, respectively  $c_1(\mathbf{r})$  and  $c_2(\mathbf{r})$  (expressed in mass per unit volume), can be defined at each space point  $\mathbf{r}$ . In functional form, the Gibbs free energy of this system can be written as:

$$G = \int d\mathbf{r} \ g(\mathbf{r}),\tag{1}$$

where  $g(\mathbf{r})$  is a free energy density,  $d\mathbf{r}$  denotes a volume element and integration extends over the entire system. Working in the spirit of the treatments currently adopted to study the problem at hand, we assume that composition gradients are small compared to the reciprocal of intermolecular distances and we expand  $g(\mathbf{r})$  about its value  $g^b$  for a bulk phase of uniform composition. Assuming further that no external field or other source of spatial anisotropy is present, we only retain in this expansion the terms compatible with the tensoral invariance of  $g(\mathbf{r})$  with respect to symmetry operations of rotation [9]. To the leading order, this yields:

$$g(\mathbf{r}) = g^{b}(c_{1}, c_{2}) + \frac{1}{2} \sum_{i,j=1}^{2} \kappa_{ij}(c_{1}, c_{2}) \nabla c_{i} \cdot \nabla c_{j}.$$
 (2)

The scalar term  $g^b$  takes into account the non-ideal, energetic and/or entropic destabilizing effects responsible for the spinodal decomposition of uniform bulk phases. The  $\kappa_{ij}$ 's describe the interactions between ij-pairs of particles and are the elements of a symmetric matrix when the medium is isotropic. Contrary to  $g^b$ , these gradient terms should always be stabilizing: in the spinodal domain, their effect should insure the minimization of interfacial boundary regions and the existence of an upper cut-off,  $k_u$ , on the band of unstable Fourier modes.

As explained in the introduction, it is important in order to study the coupling of chemical reactions and spinodal decomposition, to take into account that these two processes are controlled by thermodynamic forces which are linked to the components chemical potentials  $\mu_1$  and  $\mu_2$ . The remaining part of this section serves the purpose of setting up the reaction-diffusion equation obeyed by the system dynamics in agreement with this requirement and in as general form form as possible,.

Let us first explicitate that in (2),  $g(\mathbf{r})$  is a first order homogeneous function of the concentrations  $c_i$ . For any number  $\alpha$ , one has:

$$g^b(\alpha c_1, \alpha c_2) = \alpha \ g^b(c_1, c_2) \tag{3}$$

$$\kappa_{ij}(\alpha c_1, \alpha c_2) = \alpha^{-1} \kappa_{ij}(c_1, c_2). \tag{4}$$

Choosing  $\alpha$  to be equal to the density  $\rho = c_1 + c_2$ , we reexpress the functional dependence of  $g^b$  and of the  $\kappa_{ij}$ in terms only of the coarse grained massic fraction  $x \equiv x_1(\mathbf{r}) = c_1/\rho$  so that the free energy density (2) thus rewrites as  $(x_2(\mathbf{r}) = 1 - x)$ :

$$g(\mathbf{r}) = \rho \ g^b(x) + \frac{1}{2} \sum_{i,j=1}^2 \frac{\kappa_{ij}(x)}{\rho} \nabla c_i \cdot \nabla c_j.$$
 (5)

Assuming that the density  $\rho = c_1 + c_2$  is constant throughout the system, immediately transforms this expression into the familiar Cahn-Hilliard form

$$g(\mathbf{r}) = \rho \left[ g^b(x) + \frac{\kappa(x)}{2} (\nabla x)^2 \right], \tag{6}$$

where  $\kappa(x) = \kappa_{11}(x) + \kappa_{22}(x) - 2 \kappa_{12}(x)$  measures that contribution to non-ideality which arises from the existence of an unbalance between the mean attractive forces between like molecules and the attractive force between unlike molecules. This parameter must be positive so that *per se* the occurrence of inhomogeneities results in an increased free energy. Surface tension effects then insure that concentration gradients cannot grow *ad infinitum* and that the spinodal instability is bounded by an upper cutoff; as a consequence, inhomogeneities corresponding to large wave numbers (small wavelengths) are always damped.

Given that  $\rho = c_1 + c_2$  is constant, it could seem, at first sight, that introducing this factor in (5) complicates

notations unnecessarily. This is the case when spinodal decomposition takes place in the absence of chemical reactions: to write down the diffusion equation describing the system evolution, it is then sufficient to know the chemical potential difference

$$\Delta \mu \equiv \mu_1 - \mu_2 = \frac{\delta G(x, 1-x)}{\delta x},$$

so that the factor  $\rho = c_1 + c_2$  may be forgotten. Here however, we need to keep track of the complete dependence of G with respect to  $c_1$  and  $c_2$ . Indeed, later on, to incorporate chemical reactions and write down the reactiondiffusion equation giving the system evolution, we shall need to know the chemical potentials of each component separately. In deriving the latter from the classical defining relations

$$\mu_1 = \frac{\delta G(c_1, c_2)}{\delta c_1},\tag{7}$$

$$u_2 = \frac{\delta G(c_1, c_2)}{\delta c_2}, \tag{8}$$

it is essential to remember that  $c_1$  and  $c_2$  must be treated as independent quantities. Replacing expression (5) for g in (1) and using (7,8), we obtain that

ŀ

$$\mu_1 = \mu_1^b(x) - \frac{1}{2} [\kappa(x) + \Delta(x)] \nabla^2 x$$
$$- \frac{1}{2} \partial_x [x \ \kappa(x) + \Delta(x)] (\nabla x)^2, \qquad (9)$$

$$\mu_{2} = \mu_{2}^{b}(x) + \frac{1}{2} \big[ \kappa(x) - \Delta(x) \big] \nabla^{2} x 
 + \frac{1}{2} \partial_{x} \big[ (1-x) \kappa(x) - \Delta(x) \big] (\nabla x)^{2}, \quad (10)$$

where  $\mu_1^b$  and  $\mu_2^b$  are the bulk chemical potentials of the components, and  $\Delta(x) \equiv \kappa_{11}(x) - \kappa_{22}(x)$  measures deviations from ideality which arise when the attractive forces pulling together identical molecules are different; clearly, this contribution to non-ideality is distinct from the one described by  $\kappa(x)$ . Depending upon whether  $\Delta(x)$  is equal to zero or not, we shall call the mixture symmetrical or unsymmetrical[19]. In deriving (9,10) we have used the classical relation

$$x \ \frac{\partial \mu_1^b(x)}{\partial x} + (1-x) \ \frac{\partial \mu_2^b(x)}{\partial x} = 0, \tag{11}$$

which the bulk chemical potentials  $\mu_1^b(x)$ , and  $\mu_2^b(x)$  must satisfy in view of the extensivity properties of G. Eq. (11) is of particular interest for the following because it allows us to specify the bulk chemical potentials  $\mu_1^b$  and  $\mu_2^b$  in terms of a single unknown function Z(x). One has:

$$\mu_1^b(x) = \mu_1^0 + \int_1^x dx' \ (1 - x') \ Z(x'), \qquad (12)$$

$$\mu_2^b(x) = \mu_2^0 - \int_0^x dx' \ x' \ Z(x'), \tag{13}$$

where the chemical potentials  $\mu_1^0$  and  $\mu_2^0$  of components 1 and 2 in pure form are functions of temperature and pressure only. The integrations extend from the situation in which the components are pure to the one in which they form a uniform mixture the composition of which is given by  $c_1 = \rho x$ ,  $c_2 = \rho (1 - x)$ . In terms of the chemical potentials (9,10), the free energy density writes as  $g(\mathbf{r}) = c_1 \ \mu_1 + c_2 \ \mu_2$ . Integrating the Laplacian terms of this expression over a volume of mixture, enclosed by a surface on which boundary terms vanish [9], we recover the Cahn-Hilliard form (6) where  $g_b$  is given by

$$g^{b}(x) = \rho \left[ x \ \mu_{1}^{b}(x) + (1-x) \ \mu_{2}^{b}(x) \right]. \tag{14}$$

Noteworthy, while the chemical potentials (9,10) depend on the self-interactions difference  $\Delta(x)$ , the usual Cahn-Hilliard free energy (6) does not. As a result, classically  $\Delta(x)$  plays no role in the spinodal decomposition problem of purely diffusive binary mixtures. We shall see below that this situation changes in the presence of chemistry. Chemical rates, are in general more complicated functions of chemical potentials than diffusion and as a rule they depend upon  $\Delta(x)$  explicitly. The latter quantity becomes then a essential parameter for the stability properties of the mixture.

#### B. Examples: perfect and regular solutions

With the derivation of expressions (6) and (9,10) the relations existing between the square-gradient free energy density of a binary mixture and the chemical potentials of its components have been stated in general form. This formalism involves essentially three unknown functions, Z(x),  $\kappa(x)$  and  $\Delta(x)$ , which have to be determined on the basis of the properties particular to the system considered.

The first two functions govern the behavior of purely diffusive systems. For the sake of concreteness, before we pursue our general treatment, let us specify these functions for two classical types of mixtures: perfect and regular solutions. Our purpose here is purely illustrative. The results established in the next sections will not be restricted to the choice of a particular mathematical form for Z(x) and  $\kappa(x)$ .

For simplicity and without loss of generality, we set  $\rho = 1$  in the following. We first consider the case of two components forming a perfect solutions in a uniform bulk phase. This corresponds to the choice:

$$\begin{aligned} \kappa(x) &= 0\\ Z(x) &= \frac{RT}{x \ (1-x)} \end{aligned}$$

Replacing these expressions and (12-14) in (6), it immediately follows that  $g \equiv g(\mathbf{r})$  is constant in space and represents the free energy density of a perfect solution,

$$= g^{0} + RT [x \ln(x) + (1-x) \ln(1-x)]. \quad (15)$$

 $g^0 = \mu_1^0 x + \mu_2^0 (1 - x)$ , is the free energy density before mixing; the second term is the entropy of mixing which is of ideal form.

As a second example, we consider a symmetrical nonuniform regular solution with a constant, stabilizing surface tension term. This amounts to putting:

$$\begin{aligned} \kappa(x) &= \kappa > 0 \quad (\kappa = \text{const.}) \\ Z(x) &= \frac{RT}{x \ (1-x)} - 2\Omega \end{aligned}$$

and yields:

g

$$g(\mathbf{r}) = g^{0} + RT \left[ x \ln(x) + (1-x) \ln(1-x) \right] + \Omega x (1-x) + \frac{\kappa}{2} (\nabla x)^{2}, \qquad (16)$$

where  $\Omega$  measures the bulk non-ideality; when  $\Omega > 0$ , this system exhibits a critical point for  $x = x_c = 1/2$  and  $T = T_c = \Omega/2R$  [11, 14].

#### C. Diffusive flux and spinodal instability

The rate of diffusion is proportional to the divergence of the flux  $\mathbf{J}_x$  of particles 1 with respect to particles 2. The thermodynamic force,  $\mathbf{F}$ , driving this process is the gradient of the chemical potential difference between molecules 1 and 2 ( $\mathbf{F} = -\nabla[\mu_1 - \mu_2]$ ). To relate  $\mathbf{J}_x$  and  $\mathbf{F}$ , we adopt the usual phenomenological law:

$$\mathbf{J}_x = -L(x)\nabla(\mu_1 - \mu_2) \tag{17}$$

where L(x) is a function of composition (Onsager's coefficient), the positivity of which is required in order that entropy production be positive. In the absence of chemistry, the diffusion equation for x thus reads:

$$\partial_t x = \nabla \left[ L(x) \nabla (\mu_1 - \mu_2) \right]. \tag{18}$$

The stability of a homogeneous states  $x = x_0$  is easily determined by considering the evolution of a small perturbation,  $\delta x(\mathbf{r}, t) = x(\mathbf{r}, t) - x_0$ , that moves the system away from this state:

$$\partial_t \delta x(\mathbf{r}, t) = L_0 \nabla^2 \left[ \delta \mu_1(\mathbf{r}, t) - \delta \mu_2(\mathbf{r}, t) \right].$$
(19)

Here,  $L_0$  stands for  $L(x_0)$ . We also introduce the simplified notations  $\kappa_0 = \kappa(x_0)$ ,  $Z_0 = Z(x_0)$  and  $\Delta_0 = \Delta(x_0)$ . The expressions for  $\delta\mu_1$  and  $\delta\mu_2$  are calculated from relations (9-10) after substituting in them expressions (12-13) for the bulk chemical potentials. This yields:

$$\delta\mu_1(\mathbf{r},t) = \left[ (1-x_0)Z_0 - \frac{\kappa_0 + \Delta_0}{2} \nabla^2 \right] \delta x(\mathbf{r},t), (20)$$

$$\delta\mu_2(\mathbf{r},t) = \left[-x_0 Z_0 + \frac{\kappa_0 - \Delta_0}{2} \nabla^2\right] \delta x(\mathbf{r},t).$$
(21)

Eq. (19) may now be written in Fourier space as

$$\partial_t \delta x(k,t) = L_0 k^2 \big[ \mu_1'(x_0,k) - \mu_2'(x_0,k) \big] \, \delta x(k,t), \quad (22)$$

where,  $\mu'_1$  and  $\mu'_2$  are obtained from (20-21):

$$\mu_1'(x_0,k) = (1-x_0)Z_0 + \frac{1}{2} [\kappa_0 + \Delta_0]k^2 \qquad (23)$$

$$\mu_2'(x_0,k) = -x_0 Z_0 - \frac{1}{2} [\kappa_0 - \Delta_0] k^2.$$
 (24)

Replacing (23,24) in (22), yields that the linear growth coefficient of mode k is given by:

$$\Gamma(k) = -L_0 k^2 \left[ Z_0 + \kappa_0 k^2 \right].$$
 (25)

Since  $\kappa_0$  must be positive, the sign of  $Z_0 = \partial^2 g(x)/\partial x^2$ at  $x = x_0$  determines the nature of the free energy extremum; the homogeneous states is stable for positive  $Z_0$ . If  $Z_0$  is negative, the homogeneous state  $x_0$  is unstable with respect to perturbations the wave numbers k of which lie in the range:

$$0 < k < \sqrt{-\frac{Z_0}{\kappa_0}} \equiv k_u. \tag{26}$$

In that case, the system leaves the homogeneous state and tends to develop two domains of different concentration separated by an interface. The condition  $Z_0 = 0$ thus defines the so-called spinodal domain. The growth factor of the fastest growing mode during the early evolution of spinodal decomposition, i. e.,

$$k_f = \sqrt{-\frac{Z_0}{2\kappa_0}},\tag{27}$$

is then given simply by:

$$\Gamma(k_f) = L_0 \frac{Z_0^2}{4\kappa_0}.$$
(28)

#### D. Coupling of diffusion and chemical reactions

Suppose now that in addition to diffusion, the composition of the binary mixture may also vary due to chemical reactions and/or relaxation phenomena (excitation processes, conformational changes, photochemical processes, etc.). Suppose further that there are R linearly independent processes of this sort, and that the mixture is an open system in contact with external reservoirs of constant composition which may maintain it out of thermodynamic equilibrium.

The molecular mechanisms underlying these processes need not be detailed. It suffices for our general purpose to know that they amount to transformations which interconvert the mixture components into each other. Phenomenologically, these processes can be represented in chemical reaction form as:

$$a_{1r}X_1 + a_{2r}X_2 + \sum_i a_{ir}B_i \rightleftharpoons$$
$$b_{1r}X_1 + b_{2r}X_2 + \sum_i b_{ir}B_i (29)$$

where the index  $r = 1, \ldots, R$  labels the chemical transformations, the  $B_i$ 's represent the composition variables describing the state of the external reservoirs and the coefficients  $a_{jr}, b_{jr}, (j = 1, 2, i)$  represent the molecularities of components 1, 2 and  $B_i$ , respectively in the forward and backward direction of chemical reaction r.

As the external reservoirs composition and the mixture density are constant[20], the mass balance equation giving the spatio-temporal evolution of the system reads:

$$\rho \ \partial_t x = \sum_{r=1}^R \nu_{1r} M_1 w_r + \nabla \big[ L(x) \nabla (\mu_1 - \mu_2) \big], \qquad (30)$$

where  $w_r$  is the rate of reaction r. Since mass is conserved in chemical reactions, the stoichiometric coefficients,  $\nu_{kr} = b_{kr} - a_{kr}$ , and molecular masses,  $M_k$  of components 1 and 2 (k = 1, 2) are linked by the relation

$$\sum_{k=1}^{2} \nu_{kr} M_k = 0 \qquad (\forall r). \tag{31}$$

It is therefore convenient for the following to define  $\nu_r \equiv \nu_{1r}M_1 = -\nu_{2r}M_2$ , and putting  $\rho = 1$ , to rewrite (30) more simply as:

$$\partial_t x = \sum_{r=1}^R \nu_r \ w_r + \nabla [L(x)\nabla(\mu_1 - \mu_2)].$$
(32)

Two qualitatively different kinds of transformations are encompassed by (29) according to whether or not the reservoirs in contact with the mixture are involved. In the latter case, one has  $a_{ir} = b_{ir} = 0$  ( $\forall i$ ) so that only the components 1 and 2 of the mixture participate in the reaction. In the former case, at least some of the  $a_{ir}$  and  $b_{ir}$  coefficients are different from zero and the reaction is a reservoir "driven" reaction or exchange process. In driven reactions some of the  $B_i$ 's may act as catalysts, e. g., when  $a_{ir} = b_{ir}$ . To each reaction rate  $w_r$  is associated a thermodynamic force, or chemical affinity  $\mathcal{A}_r$ . With the notations

$$\vec{\mu}_r = a_{1r}M_1\mu_1 + a_{2r}M_2\mu_2 + \vec{\mu}_{Br} \tag{33}$$

$$\overleftarrow{\mu}_r = b_{1r}M_1\mu_1 + b_{2r}M_2\mu_2 + \overleftarrow{\mu}_{Br},$$
 (34)

where  $\vec{\mu}_{rB} = \sum a_{ir} M_{B_i} \mu_{B_i}$  and  $\overleftarrow{\mu}_{rB} = \sum b_{ri} M_{B_i} \mu_{B_i}$ , the affinity  $\mathcal{A}_r$  can be expressed in terms of chemical potentials as

$$\mathcal{A}_r = \overrightarrow{\mu}_r - \overleftarrow{\mu}_r, \tag{35}$$

while the reaction rates  $w_r = \vec{v}_r(\vec{\mu}_r) - \vec{v}_r(\vec{\mu}_r)$  are the difference of two terms corresponding respectively to the direct rate of reaction  $\vec{v}_r$ , which depends upon  $\vec{\mu}_r$ , and to the reverse rate of reaction  $\vec{v}_r$  which depends upon  $\vec{\mu}_r$ .

By definition, thermodynamic equilibrium is the state for which the affinity and rate of each chemical reaction simultaneously vanish:

$$\mathcal{A}_r = 0, \qquad \qquad w_r = 0. \tag{36}$$

We conclude therefrom that, when  $\overleftarrow{\mu}_r = \overrightarrow{\mu}_r \equiv \widetilde{\mu}_r$ , the equality  $\overrightarrow{v}_r(\widetilde{\mu}_r) = \overleftarrow{v}_r(\widetilde{\mu}_r)$  must hold. Since, this equality has to be obeyed independently from the value of  $\widetilde{\mu}_r$ , the functions  $\overrightarrow{v}_r(\cdot)$  and  $\overleftarrow{v}_r(\cdot)$  must be identical so that we may set  $\overrightarrow{v}_r(\cdot) = \overleftarrow{v}_r(\cdot) = v_r(\cdot)$ . The most general expression for the rate of a chemical reaction is thus[21]:

$$w_r = v_r(\vec{\mu}_r) - v_r(\vec{\mu}_r). \tag{37}$$

In addition, by the thermodynamic postulate that the entropy production of linearly independent chemical processes must be positive,  $w_r$  and  $\mathcal{A}_r$  must always have the same sign. This implies that  $v_r$  must be a monotonically increasing (nonlinear) function.

When the affinity is small, Eq.(37) reduces to the linear relation  $w_r = \gamma \mathcal{A}_r$ , where  $\gamma$  is a positive constant. For the general nonlinear regime, it is usual to write the reaction rate as

$$w_r = v_r(\vec{\mu}_r) \left( 1 - \exp[-\mathcal{A}_r/RT] \right), \tag{38}$$

which is compatible with (37) if  $v_r(\mu) \propto \exp(\mu/RT)$ .

At this stage, it is important to stress that a wide variety of reaction-diffusion systems can be described by Eq. (32). Nevertheless, in the next section, we shall see that the stability properties of their uniform stationary states can be classified quite simply into four general categories once the chemical potentials are known and the thermodynamic requirements mentioned above concerning the rates  $w_r$  are taken into account.

#### III. LINEAR STABILITY OF HOMOGENEOUS STATIONARY STATES

### A. Linear growth coefficients $\tilde{\Gamma}(k)$

Contrary to what happens with purely diffusive systems (cf. section II C), in the case of chemically reacting mixtures, the homogeneous stationary state concentrations  $x_0$  can no longer be chosen at will. For a given temperature and pressure, their value is fixed by the chemical reactions and the state of the external reservoirs with which the system exchanges matter and energy, i. e., by the solutions  $x_0$  of the conservation relation:

$$\sum_{r=1}^{R} \nu_r \left[ v_r(\vec{\mu}_r(x_0)) - v_r(\overleftarrow{\mu}_r(x_0)) \right] = 0.$$
 (39)

Using the same notations as in section II C, the linear stability of these states w. r. t. to a small perturbation  $\delta x(\mathbf{r}, t)$  is given by

$$\partial_t \delta x(\mathbf{r}, t) = \left[ \sum_{r=1}^R \nu_r \left( \delta v_r(\vec{\mu_r}) - \delta v_r(\vec{\mu_r}) \right) + L_0 \nabla^2 (\delta \mu_1 - \delta \mu_2) \right].$$
(40)

The quantities  $\delta v_r(\vec{\mu_r})$  and  $\delta v_r(\vec{\mu_r})$  given by

$$\begin{split} \delta v_r(\vec{\mu}_r) &= \left. \frac{dv_r}{d\vec{\mu}_r} \right|_{\vec{\mu}_r(x) = \vec{\mu}_r(x_0)} \delta \vec{\mu}_r(\mathbf{r}, t) = \vec{v}'_r \delta \vec{\mu}_r(\mathbf{r}, t) \\ \delta v_r(\vec{\mu}_r) &= \left. \frac{dv_r}{d\vec{\mu}_r} \right|_{\vec{\mu}_r(x) = \vec{\mu}_r(x_0)} \delta \vec{\mu}_r(\mathbf{r}, t) = \vec{v}'_r \delta \vec{\mu}_r(\mathbf{r}, t) \end{split}$$

define  $\vec{v}'_r$  and  $\overleftarrow{v}'_r$ . Introducing the dimensionless parameters

$$z_{0} = \left(\frac{L_{0}}{\kappa_{0}\vec{v}_{1}'}\right)^{\frac{1}{2}} Z_{0}, \qquad \rho_{r} = \frac{\overleftarrow{v}_{r}'}{\overrightarrow{v}_{r}'},$$
$$\tau_{r} = \left(\frac{\kappa_{0}}{L_{0}\vec{v}_{1}'}\right)^{\frac{1}{2}} \vec{v}_{r}', \qquad \delta = \frac{\Delta_{0}}{\kappa_{0}}, \qquad (43)$$

and proceeding as in section II C, the linear growth coefficient  $\tilde{\Gamma}(k)$  corresponding to the rescaled time and space variables

$$\tilde{t} = \vec{v}_1' t, \quad \tilde{\mathbf{r}} = \left[\frac{L_0 \kappa_0}{\vec{v}_1'}\right]^{-\frac{1}{4}} \mathbf{r},$$
(44)

can be written in compact form as

$$\tilde{\Gamma}(k) = -k^4 + [M_0(\delta) - z_0] k^2 + N_0 z_0.$$
(45)

The explicit expressions for  $M_0(\delta)$  and  $N_0$  are:

$$M_0(\delta) = C_1(x_0)\delta + C_2(x_0), \qquad (46)$$

$$N_0 = C_1(x_0) \left(1 - 2x_0\right) + C_2(x_0) \tag{47}$$

with

$$C_1(x_0) = \sum_{r=1}^R \frac{\nu_r \tau_r}{2} \quad \left[ a_{1r} M_1 + a_{2r} M_2 + (b_{1r} M_1 + b_{2r} M_2) \rho_r \right], \quad (48)$$

$$C_2(x_0) = \sum_{r=1}^R \frac{\nu_r \tau_r}{2} \quad \left[ a_{1r} M_1 - a_{2r} M_2 - (b_{1r} M_1 - b_{2r} M_2) \rho_r \right].$$
(49)

The  $\rho'_r$ s and  $\tau'_r$ s are positive functions whatever the value of  $x_0 \in [0, 1]$ . Furthermore, it should be kept in mind that the value of  $x_0$ , as given by Eq. (39), does not depend on the coefficients  $\kappa_{ij}$  multiplying the gradient terms of g, and thus does not depend on  $\delta$ . This is noteworthy because  $\delta$  largely controls the system stability. Indeed, one may already observe that in the expression for  $\tilde{\Gamma}(k)$ (cf. Eq. (45)),  $\delta$  only appears in the term of order  $k^2$ ; by varying  $\delta$ , the sign and magnitude of this term, and hence the stability of  $x_0$ , can be modified at will.

#### B. Classification of instabilities

For non-reactive systems, it has been shown in section II C that the change of sign of  $Z_0$ , or equivalently here of  $z_0$ , defines the boundary of the spinodal domain:  $Z_0 < 0$  is the condition under which (25) admits a finite band of unstable modes which includes the mode k = 0. Here, in order to classify the wider class of behaviors which become possible in the presence of chemistry, we remark the following:

(i) Since the linear stability of the homogeneous stationary state solutions of (39) with respect to the mode k = 0 is given by

$$\tilde{\Gamma}(0) = N_0 \ z_0,\tag{50}$$

and hence, only depends on the sign of the functions  $N_0$ and  $z_0$ , the stability diagrams reporting the behavior of  $\tilde{\Gamma}(k)$  can be divided into the four basic cases described in III B 1-4.

(ii) In the large wavenumbers domain  $(k \to \infty)$ , the eigenvalues  $\tilde{\Gamma}(k)$  are always negative: surface tension, which prevents too sharp interfaces to develop, and diffusion, which is all the more important that the wavelength of spatial heterogeneities is small, cooperate to

damp short wavelength fluctuations efficiently. As a consequence, if at k = 0,  $d\tilde{\Gamma}(0)/dk^2 = M_0(\delta) - z_0$  is positive,  $\tilde{\Gamma}(k)$  must pass through a maximum for

$$k = \tilde{k}_f = \sqrt{\frac{M_0(\delta) - z_0}{2}}.$$
 (51)

(iii) Replacing this expression of  $k_f$  in (45), one finds that there exists a finite band of unstable modes, from which the mode k = 0 is excluded, if the inequalities

$$\tilde{\Gamma}(0) = N_0 \ z_0 < 0,$$
(52)

and

$$\tilde{\Gamma}(\tilde{k}_f) = \left[\frac{M_0(\delta) - z_0}{2}\right]^2 + N_0 z_0 > 0$$

$$= \frac{1}{4} C_1(x_0)^2 \delta_0^2 + \frac{C_1(x_0)}{2} [C_2(x_0) - z_0] \delta_0$$

$$+ \frac{1}{4} [C_2(x_0) - z_0]^2 + N_0 z_0 > 0$$
(53)

are fulfilled.

(iv) In strongly unsymmetrical mixtures, i. e., in the limit  $|\delta| \to \infty$ , inequality (53) is always satisfied. Looking for the transition point where  $\tilde{\Gamma}(\tilde{k}_f)$  passes from the negative to the positive values, so that the finite band of unstable modes appears, we consider (53) as an equation in  $\delta$  and solve for its roots. This yields the values

$$\delta_{\pm} = \frac{1}{C_1(x_0)} \bigg[ -C_2(x_0) + z_0 \pm 2\sqrt{-N_0 z_0} \bigg], \quad (54)$$

which replaced in (51) permit to calculate the wavenumber  $k_c$  corresponding to this point of marginal stability. One has:

$$k_c = (-z_0 N_0)^{\frac{1}{4}} . (55)$$

In brief, the main outcome of the above analysis, is that if the mode k = 0 is stable (cf. inequality (52)), and if the self-interaction difference parameter  $\delta$  does not belong to the interval  $(\delta_{-}, \delta_{+})$ ,

$$\delta \ni (\delta_{-}, \delta_{+}), \tag{56}$$

then the uniform stationary state  $x_0$  is unstable with respect to a finite band of wavenumbers,

$$0 < k_{-} \le k \le k_{+},\tag{57}$$

the boundaries of which can be written in terms of the values of  $\tilde{k}_f$  and  $k_c$ , given by (51,55), as [22]:

$$k_{\pm} = \tilde{k}_f \sqrt{1 \pm \sqrt{1 - \left(\frac{k_c}{\tilde{k}_f}\right)^4}}.$$
 (58)

Let us now analyze more in detail the four cases which may be encountered depending on the sign of  $z_0$  and  $N_0$ .

#### 1. Turing kind of instability

If  $z_0 > 0$  and  $N_0 < 0$ , the uniform stationary state  $x_0$  lies above the spinodal line and  $\Gamma(0) < 0$ . The components of the mixture are thus miscible. Nevertheless, pattern formation could take place if the value of  $\delta$  fulfills condition (56). No phase separation being involved, the symmetry breaking instability appearing in that case is reminiscent of the Turing instability well-known in classical reaction-diffusion theory. The originality of the instability mechanism found here however is that it does not require the diffusion coefficients of two reactants, the so called "activator" and "inhibitor", to be unequal. Instead, it is the difference of self-interactions between identical particles which controls the formation of patterns with an intrinsic wavelength. If condition (56) is not satisfied, i. e., if  $\delta(x_0) \in (\delta_-, \delta_+)$ ,  $\Gamma(k) < 0$  for all values of k and the stationary state  $x_0$  is stable.

#### 2. Spinodal instability

If  $z_0 < 0$  and  $N_0 < 0$ , the stationary state  $x_0$  lies below the spinodal line. There exists a band of unstable modes which includes the uniform perturbation mode k = 0. We conclude that spinodal decomposition should proceed in this case essentially as in the absence of chemistry. The fastest growing mode is either (51) or the mode k = 0, depending upon whether  $M_0(\delta) - z_0 > 0$  or not. When the system is strongly unsymmetrical, the early growth of  $\tilde{k}_f$  is faster than that of the fastest mode of the purely diffusive case. Indeed, under the transformation (43,44) expressions (27,28) become in dimensionless form  $k_f = \sqrt{-z(x_0)/2}$  and  $\Gamma(k_f) = z(x_0)^2/4$ . By comparing with (51) and (53), it is clear that for  $|\delta(x_0)| \to \infty$ , the inequality  $\tilde{\Gamma}(\tilde{k}_f) - \Gamma(k_f) > 0$  holds.

#### 3. Chemical instability

If  $z_0 > 0$  and  $N_0 > 0$ , the stationary state  $x_0$  lies above the spinodal line. Nevertheless, there exists a band of unstable modes which includes the uniform perturbation mode k = 0. This instability originates from nonlinearities associated with the reaction scheme considered rather than from the non-ideality of molecular interactions in the mixture. It should in general involve a multistationary state phenomenon and is in this respect distinct from spinodal decomposition or from the Turing kind of instability described in case 1. As in the preceding case, depending upon whether  $M_0(\delta) - z_0 > 0$  or not, the fastest growing mode is either (51) or the mode k = 0; again also in strongly asymmetric mixtures,  $\tilde{k}_f$ grows more rapidly than the fastest mode of the purely diffusive system.

## 4. Ostwald ripening freezing instability

If  $z_0 < 0$  and  $N_0 > 0$ , the stationary state  $x_0$  lies below the spinodal line, but if  $\delta(x_0) \in (\delta_-, \delta_+)$ , one has  $\tilde{\Gamma}(k) < 0$  for all values of k: the chemical reactions completely inhibit the phase separation. If on the contrary,  $\delta(x_0) \ni (\delta_-, \delta_+)$ , there exists a finite band of unstable modes,

$$0 < k_{\ell} \equiv k_{-} < k < k_{+} \equiv \tilde{k}_{u}, \tag{59}$$

which excludes the mode k = 0. This is the interesting case where patterns with an intrinsic wave length may appear while at the same time the mixture is immiscible. As explained in the introduction, such a behavior amounts in the terminology introduced by [6, 7] to a freezing of the Ostwald ripening stage of phase separation. We shall therefore refer to it as the Ostwald ripening freezing instability (ORFI). The following properties are noteworthy:

(i) It is clear from their definitions that the purely diffusive upper cut-off,  $k_u = \sqrt{z_0}$  (cf.(26) rewritten using (44)), and the lower and upper boundaries of the unstable modes band, respectively  $k_-$  and  $k_+$  (cf. (51,55,58)), depend on different parameters and hence, can be varied independently from each other. For example, only  $k_{\pm}$  depends on  $\delta$  and on the concentrations  $B_i$  maintained constant in the external reservoirs. It is thus to be expected that by changing these concentrations, one may vary the position of the unstable band  $(k_{\ell}, \tilde{k}_u)$  with respect to that of the purely diffusive system. In fact, these two bands could even become disconnected, so that  $k_u < k_{\ell} < \tilde{k}_u$ , if the inequality

$$-C_1(x_0) \left[ \delta + x_0 \right] z_0 > 0 \tag{60}$$

holds, in which case, chemistry hampers the damping by diffusion of large k modes.

We shall not, in the present work, attempt to determine and to classify the great variety of patterns which appears once the homogeneous stationary state  $x_0$  becomes unstable. This requires a nonlinear analysis which we plan to report elsewhere [16]. The objective of our discussion in the next section is to precise more explicitly the thermodynamic and kinetic properties which condition the appearance of patterns and of the ORFI just defined. The results of this discussion are illustrated by simulating numerically the behavior of an example in section IV C

### IV. THERMODYNAMIC AND KINETIC CONDITIONS ON THE CHEMICAL FREEZING OF PHASE SEPARATION

#### A. Dissipation threshold

Let us first consider what kind of instability may occur if the homogeneous state is an equilibrium state ( $x_0 =$   $x_e$ ). At equilibrium, all the affinities vanish  $(\vec{\mu}(x_e) = \vec{\mu}(x_e))$  and consequently  $\vec{v}'_r = \vec{v}'_r \equiv v'_r$ , or equivalently,  $\rho_r(x_e) = 1$ . In that case, the parameters  $M_0, N_0$  reduce to:

$$M_e = N_e = -\sum_{r=1}^{R} \nu_r^2 \ \tau_r, \tag{61}$$

so that the eigenvalue equation simply reads:

$$\tilde{\Gamma}_e(k) = -\left[k^2 - N_e\right]\left[k^2 + z_e\right].$$
(62)

We have mentioned in section II D that the  $v_r$  are monotonically growing functions; their derivatives are thus always positive and so are the  $\tau_r$ . Eq. (61) shows that the parameters  $M_e$  and  $N_e$  are always negative at equilibrium independently of the functions  $\tau_r$  in the reaction rates and independently of the functions  $\kappa_e$  and  $z_e$  appearing in the free energy. As (62) shows and in agreement with Duhem-Jouget theorem, the stability properties of homogeneous equilibrium state are thus entirely determined by the sign of  $z_e$  which controls diffusional stability.

The spinodal instability can then be observed at equilibrium in the domain corresponding to the spinodal region of non-reactive systems  $(z(x_e) < 0)$ . On the contrary, the ORFI can never be observed under equilibrium conditions since  $N_e$  is always negative in that case.

Furthermore, since  $N(x_e)$  is strictly negative at equilibrium, the ORFI can only appear if the external baths drive and maintain the system at a finite distance from its chemical equilibrium state. To see this, let us suppose that  $x_0$  belongs to the thermodynamic branch of steady states[15] and lies close to the equilibrium state:  $x_0 = x_e + \delta x_0$  with  $|\delta x_0|/x_e \ll 1$ . The first terms in the expansion of  $N_0$  are

$$N_0 = N_e + \frac{dN}{dx_0} \delta x_0 + \dots, \tag{63}$$

and the minimal distance from equilibrium at which the condition  $N_0 > 0$  may be realized, is:

$$\delta x_0 > |\frac{N_e}{\frac{dN}{dx_0}}|. \tag{64}$$

The amplitude of the distance from equilibrium  $\delta x_0$  has thus a lower bound below which the ORFI cannot be observed. It is equivalent to conclude that the ORFI is a dissipative instability, or that the patterns are *dissipative* structures [15].

#### B. Properties of reaction schemes

In order that the ORFI be possible, the chemical scheme must include at least two reactions  $(R \ge 2)$ . Indeed, if R = 1, the stationary condition for homogeneous

states is equivalent to the equilibrium condition and we have shown in the previous section that the ORFI cannot be observed at equilibrium. Furthermore, the scheme must include at least one autocatalytic reaction, i. e., a reaction in which the stoichiometric coefficients of at least one component of the mixture  $(X_1 \text{ or } X_2)$  are non zero for both the direct and the reverse transformations  $(a_{r1} \neq 0, b_{r1} \neq 0 \text{ or } a_{r2} \neq 0, b_{r2} \neq 0)$ . Indeed, if all the reactions in the scheme are non autocatalytic, they can be rewritten as:

$$\nu_r X + \sum_i a_{ri} B_i \rightleftharpoons \nu_r Y + \sum_i b_{ri} B_i, \qquad (65)$$

where  $\nu_r$  stands now for  $a_{r1}$  and  $b_{r2}$ . In that case, the parameter  $N(x_0)$  reduces to:

$$N(x_0) = -\frac{1}{2} \sum_{r=1}^{R} \nu_r^2 \ \tau_r(x_0) \left[ 1 - x_0 + \rho(x_0)(1 + x_0) \right] \ (66)$$

which is negative and the ORFI is impossible. Hence, it is interesting to note that simple reaction schemes cannot give rise to the ORFI. In particular, binary systems undergoing simple reactions like  $B_1 + X \rightleftharpoons Y + B_2$  will not induce pattern formation, even under far from thermodynamic equilibrium conditions.

### C. Example

To illustrate by an example the ORFI described in section III, we consider the following two step autocatalytic reaction scheme:

$$B_1 + 2X_1 + X_2 \rightleftharpoons 3X_1 + B_2 \tag{67}$$

$$B_1 + X_1 + X_2 \rightleftharpoons 2X_2 + B_2. \tag{68}$$

 $B_1$  and  $B_2$  are initial and final products the concentrations of which are kept constant by external reservoirs. The mixture free energy is defined by choosing for z(x)the regular solution form:

$$z(x) = \frac{RT}{x(1-x)} - 2\Omega \tag{69}$$

In the reservoir, for simplicity, we suppose that the mixture of  $B_1, B_2$  is ideal and define accordingly their chemical potentials as

$$\mu_{B_1} = \Psi + RT \,\ln(x_{B_1}) \tag{70}$$

$$\mu_{B_2} = \Psi + RT \, \ln(1 - x_{B_1}). \tag{71}$$

Adopting for the reaction rates the exponential form (38), the kinetic equation for x reads:

$$\partial_t x = \tau_1 \left[ \exp(\frac{\mu_{B_1} + 2\mu_1 + \mu_2}{RT}) \exp(\frac{3\mu_1 + \mu_{B_2}}{RT}) \right] - \tau_2 \left[ \exp(\frac{\mu_{B_1} + \mu_1 + \mu_2}{RT}) + \exp(\frac{2\mu_2 + \mu_{B_2}}{RT}) \right] + L\nabla^2(\mu_1 - \mu_2), \quad (72)$$

1 9.0

where

Г

$$\mu_{1} = \mu_{1}^{0} + RT \ln(x) + \Omega (1-x)^{2} -\frac{\kappa + \Delta}{2} \nabla^{2} x - \frac{\kappa}{2} (\nabla x)^{2}, \qquad (73)$$

$$\mu_{2} = \mu_{2}^{0} + RT \ln(1-x) + \Omega x^{2} + \frac{\kappa - \Delta}{2} \nabla^{2} x - \frac{\kappa}{2} (\nabla x)^{2}.$$
(74)

We have seen that the affinity  $A_T/RT = 2 \ln(x_{B_1}/x_{B_2})$  of the overall reaction

$$2 B_1 \rightleftharpoons 2 B_2 \tag{75}$$

cannot be taken equal to zero, which implies that  $x_{B_1}$ cannot be equal to 1/2. Here, we set  $x_{B_1} = 0.87$ and, for simplicity, we take the parameters depending on the intermolecular interactions as being constant:  $\kappa(x) \equiv \kappa = 1.0, \ \Delta(x) \equiv \Delta = -0.3, \ \Omega = 2.2.$  Letting further the chemical relaxation times  $\tau_1, \tau_2$  as well as the Onsager coefficient L and RT be equal to one, and choosing for to the reference chemical potentials the values  $\mu_1^0 = 0.122, \mu_2^0 = 0.183, \Psi = -0.288$  one finds that (72) admits three homogeneous stationary solutions  $x_0$ , namely:  $x_0^a = 1/2, x_0^b = 0.024279$  and  $x_0 = x_0^c =$ 0.26959. The stationary state  $x = x_0^b$  is stable for all k's, while  $x = x_0^c$  is unstable for a range of k values which includes the mode k = 0. The stationary state  $x = x_0^a$ , which lies in the spinodal region, is on the contrary, is unstable w. r. t. the finite band of wavenumber  $k_{-} \approx 0.3453 < k < k_{+} \approx 0.5295$  from which the mode k = 0 is excluded. Taking this state as average initial condition and integrating (72) numerically, one finds (see Fig. 1) that instead the Ostwald ripening process which would be observed in the absence of chemistry, a stable pattern with hexagonal symmetry finally establishes itself in the course of time.

A complete discussion of the various nonlinear behaviors which may arise in models such as (67,68) will be presented elsewhere.

# To investigate the idea that chemical reactions may freeze the unmixing of immiscible mixtures at an early stage of spinodal decomposition, we have developed an approach which is based on general thermodynamic arguments, encompasses equilibrium as well as nonequilib-

rium situations and does not rely on the FFEIA.

CONCLUSION

V.

Within the framework of this approach, we have established that the linear stability properties of chemically reacting binary mixtures can be classified into four categories, one of which (case 4, section III) corresponds to the soft mode stabilization effect leading to phase separation freezing. To encounter this case, it is indispensable that the system considered be dissipative, that it undergoes a chemical process comprising at least two independent reaction steps and that at least one of these steps be autocatalytic. It is a remarkable fact that those properties are exactly those of the model studied by Huberman in his twenty years old pioneering paper[2].

¿From a kinetic point of view, our results do not support the recent claims that the chemical freezing of phase separation is an ubiquitous phenomenon[5] susceptible to be observed with very simple chemistry and hence, with a broad class of chemical processes. In this respect, it may also be pointed out that the recent molecular dynamic studies which have been devoted to the ORFI [17], and which constitute a way of investigating this problem completely independent from our thermodynamic approach, do not confirm the predictions based on the FFEIA.

Finally, our results draw attention to the enlarged variety of symmetry breaking instabilities which may be encountered in non-ideal systems, even in domains of parameter space where the occurrence of phase transitions can be excluded. This feature which we have reported earlier in the case of anisotropic ternary mixtures [18] is further illustrated by the chemically reacting binary systems considered here, notably by their behavior in response to variations of the self-interaction parameter  $\delta$ .

#### Acknowledgments

D. C. acknowledges the financial support of the Fonds National de la Recherche Scientifique (F. N. R. S., Belgium). This work is part of a Research Program of the Centre for Nonlinear Studies and Complex Systems of the University of Brussels (U. L. B.). The simulations have been performed on the Cray J-19 of the Computation Center of the University of Brussels.

 For reviews, see e. g., J. D. Gunton, M. San Miguel and P. S. Sahni, in: Phase Transitions and Critical Phenomena, vol. 8, edited by C. Domb and J. L. Lebowitz (Academic Press 1983). K. Binder, in: Phase Transformations of Materials (Material Science and Technology 5), edited by P. Haasen (Weinheim: VCH 1991).

[2] B. A. Huberman, J. Chem. Phys. 65 (1976) 2013; see also, D. J. Scalapino and B. A. Huberman, Phys. Rev. Lett. **39** (1977) 1365; E. Coutsias and B. A. Huberman, Phys. Rev. B **24** (1981) 2592.

- [3] S. Puri and H. L. Frisch, J. Phys. A: Math. Gen. 27 (1994) 6027.
- [4] S. C. Glotzer, D. Stauffer and N. Jan, Phys. Rev. Lett. 72 (1994) 4109.
- [5] S. C. Glotzer, E. A. Di Marzio and M. Muthukumar, Phys. Rev. Lett. **74** (1995) 2034.
- [6] J. Verdasca, P. Borckmans and G. Dewel, Phys. Rev. E 52 (1995) 4616.
- [7] Q. Tran-Cong, T. Nagaki, O. Yano and T. Soen, Macromolecules 24 (1991) 1505. For recent experimental results, see A. Imagawa and Q. Tran-Cong, Macromolecules 28 (1995) 8388; Q. Tran-Cong and A. Harada (1996) 76 1162, and the references cited therein.
- [8] J. J. Christensen, K. Elder and H. C. Fogedby, Phys. Rev. E54 (1996) 2212.
- [9] J. W. Cahn and J.E. Hilliard, J. Chem. Phys. 28 (1958) 258.
- [10] R. Lefever, D. Carati and N. Hassani, Phys. Rev. Lett. 75 (1995) 1674.
- [11] I. Prigogine and R. Defay, *Chemical Thermodynamics* (Longmans, Green, New York, 1962).
- [12] M. Gitterman, J. Stat. Phys. 58 (1990) 707.
- [13] S. C. Glotzer, D. Stauffer and N. Jan, Phys. Rev. Lett. 75 (1995) 1675.
- [14] G. N. Lewis and M. Randall, *Thermodynamics* (Longmans, Green, New York, 1961); E. A. Guggenheim, *Thermodynamics* (North-Holland, Amsterdam, 1967)
- [15] P. Glansdorff and I. Prigogine, Thermodynamics of Struc-

ture, Stability and Fluctuations (Wiley-Interscience, New York, 1971); G. Nicolis and I. Prigogine, *Selforganiza*tion in Nonequilibrium Systems (Wiley-Interscience, New York, 1977).

- [16] D. Carati and R. Lefever, in preparation
- [17] S. Toxvaerd, Phys. Rev. E 53 1996, 1.
- [18] R. Lefever and D. Carati, Physica A 213 1995, 90.
- [19] The deviations of  $\Delta(x)$  from zero may be due to a difference in the range over which the interactions between pairs of identical particles operate as well as to a difference in the intensities of the attractive forces. In this sense, the notion of symmetry used here is more general than the one associated with symmetrical mixtures classically [14]
- [20] We are primarily interested in the influence of chemical reactions on unmixing under isobaric conditions; we want to avoid the additional complications which pressure and density variations, leading to the need of introducing hydrodynamic equation into the description, would entail.
- [21] It is well-known that thermodynamics does not determine chemical reaction rates; it only imposes that at equilibrium the relations (36) hold. The fact that the functions  $\vec{v}_r$  and  $\vec{v}_r$  have to be equal, is not in contradiction with this statement. Indeed, the reaction rate itself,  $w_r$ , is not prescribed since it still contains the unknown function  $v_r$ . Our analysis does not require that a particular form for this function  $v_r$  be postulated
- [22] it is easy to verify that when (56) holds, the value of  $\tilde{k}_f$  is always larger than that of  $k_c$

FIG. 1: Snapshots of the density profile for example (70) for a) t=0, b) t=200, c) t=450 and d) t=30 000. The domaine size is 90  $\times$  90 and the simulation is made with 128  $\times$  128 points. The values of the parameters are given in the text.