Derivation of the Matalon-Packter law for Liesegang patterns

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Theoretical models of the Liesegang phenomena are studied and simple expressions for the spacing coefficients characterizing the patterns are derived. The emphasis is on displaying the explicit dependences on the concentrations of the inner and the outer electrolytes. Competing theories (ion-product supersaturation, nucleation and droplet growth, induced sol-coagulation) are treated with the aim of finding the distinguishing features of the theories. The predictions are compared with experiments and the results suggest that the induced sol-coagulation theory is the best candidate for describing the experimental observations embodied in the Matalon-Packter law.

I. INTRODUCTION

Systems that exhibit pattern formation are common in nature, and patterns often emerge in the wake of a moving front.² A well known example that will be studied below is the Liesegang phenomenon,³ the formation of precipitation patterns by a moving chemical reaction front. In a typical experimental setup, one has a chemical reactant (inner electrolyte) dissolved in a gel matrix and a second reactant (outer electrolyte) poured onto the gel. The concentration of the outer electrolyte is much higher than that of the inner one and so it diffuses into the gel and reacts with the inner reactant. The reaction product then precipitates and, frequently, the observed precipitation patterns are a family of bands or rings (depending on the geometry of the system) clearly separated in the direction perpendicular to the diffusion. This process is believed to be responsible for many precipitation patterns such as the structure of agate rocks⁴ or, to cite a somewhat speculative example, the pattern of human gallstones.

Although the Liesegang patterns have been known for more than 100 years,⁵ there is still disagreement as to the mechanisms underlying this phenomenon. The main problem is the lack of a theory that could be compared with experiments quantitatively and, to a lesser degree, the lack of experiments that produce results amenable to quantitative theoretical analysis. At first the task appears to be simple. Theoretical analysis. At first the task appears to be simple. The main problem is the lack of a theory that could be compared with experiments quantitatively and, to a lesser degree, the lack of experiments that produce results amenable to quantitative theoretical analysis. At first the task appears to be simple. The theoretical analysis.

(i) After a transient time, the position $x_n$ of the $n$th band is related to the time $t_n$ of its formation through the so-called time law, $x_n \sim \sqrt{t_n}$, first discussed in Ref. 4. The time law is a rather obvious consequence of the diffusive motion of a reaction front in a gel and thus it is considered to be understood.

(ii) For large $n$, the ratio of the positions of two consecutive bands approaches a constant value

$$
\frac{x_n}{x_{n-1}} \rightarrow 1 + p,
$$

with $0.05 < p < 0.4$ typically. This last property is known as the Jablczynski law,⁶ or the spacing law, and $1 + p$ is called the spacing coefficient. Most of the detailed experimental observations concern this law. It has been found that the spacing coefficient is a nonuniversal quantity. It depends, for example, on the concentrations $a_0$ and $b_0$ of the outer and inner electrolytes which can be controlled experimentally.

Based on several different experiments, Matalon and Packter⁷ concluded that

$$
p=F(b_0) + \frac{G(b_0)}{a_0},
$$

where $F$ and $G$ are decreasing functions of their arguments. For various reagents, they found $F(b_0) \sim b_0^{-\gamma}$ with $0.2 \leq \gamma \leq 2.7$. The function $G(b_0)$ is less known but it is generally observed that it decreases with increasing $b_0$. The observation expressed in Eq. (2) is usually called the Matalon-Packter law.

A problem with the Matalon-Packter law is that, experimentally, $a_0$ and $b_0$ can usually be changed only by a factor 5-10 and, furthermore, $p$ itself has a significant error bar.
since it is determined from a finite number of \( x_n \)'s (\( n \sim 10 \to 20 \)). Thus the functional form (2) is far from unique; it should be considered as a power law fit to \( p^a \).

(iii) Finally, the width law states that the width \( w_n \) of the \( n \)-th band is an increasing function of \( n \) and typically \( w_n \sim x_n^\alpha, \alpha \geq 0 \). Since the definition of \( w_n \) is open to debate and the error bars in the measurements of \( w_n \) are rather large, this quantity has largely been ignored in the quantitative discussions of Liesegang phenomena.

Although there are a large number of interesting and often puzzling observations about Liesegang phenomena in particular systems, the above three points appear to be the only ones which describe quantifiable results and carry some generality.

One would expect that so few experimental facts can be explained easily. Indeed, there are several competing theories,\(^{10\sim 21}\) many of which\(^{10\sim 17}\) fare as well as far as the derivation of the time and spacing laws are concerned.\(^{22}\) All these theories follow how the diffusive reagents (ions in the outer and inner electrolytes) \( A \) and \( B \) turn into immobile precipitate \( D \),

\[
A + B \to \ldots C \to \ldots \to D, 
\]

by taking into account various scenarios for the intermediate steps denoted as \( \ldots C \ldots \). Since the precipitate appears through some kind of supersaturation, further differences in theories arise from the details of treating the nucleation thresholds and the growth kinetics of the precipitate.

The simplest (and first developed) theory is based on the concept of supersaturation of ion-product\(^{10}\) and has been developed by many researchers.\(^{11\sim 13,23}\) In this theory, there is no intermediate step in between \( A, B \) and \( D \). When the local product of the concentrations of the reactants, \( ab \), reaches some critical value, \( q^* \), nucleation of the precipitate \( D \) occurs. The nucleated particles grow and deplete \( A \) and \( B \) in their surroundings. As a consequence, the local level of \( ab \) drops and no new nucleation takes place. This continues until the reaction zone (where \( ab \) is maximum) moves far enough that the depletion effect of the precipitate becomes weak. Then \( ab = q^* \) is reached again and nucleation can occur. The repetition of this process leads to the formation of bands.

In the next level of complexity, theories contain a single intermediate step in \( \ldots C \ldots \) with the mechanism of band formation based on the supersaturation of the intermediate compound \( C \).\(^{16,17}\) It is assumed that \( A \) and \( B \) react to produce a new species \( C \) which also diffuses in the gel. The nature of \( C \) is not really specified; it may be a molecule as well as a colloid particle. When the local concentration of \( C \) reaches some threshold value, nucleation occurs and the nucleated particles, \( D \), act as aggregation seeds. The \( C \) particles near to \( D \) aggregate to the existing droplet (hence become \( D \)) provided their local concentration is larger than a given aggregation threshold. These models are characterized by two thresholds, one for nucleation and one for droplet growth. The depletion mechanism is similar to the one described for the ion-product theory and it leads to band formation. We shall refer to this theory as the theory of nucleation and droplet growth.

A variation of the single intermediate compound theories is based on the idea of an induced sol-coagulation process.\(^{14,15}\) The compound \( C \) is assumed to be the sol and this sol coagulates if the following two conditions are satisfied. First, the concentration of \( C \) exceeds a supersaturation threshold \( c > c^* \) and, second, the local concentration of the outer electrolyte is above a threshold \( a > a^* \) that is often referred to as the critical coagulation concentration threshold. The band formation is a consequence of the nucleation and growth of the precipitate combined with the motion of the front where \( a = a^* \).

The time- and the spacing laws follow from the above theories. Thus, to select the correct theory (provided there is a single theory for all Liesegang phenomena), one would have to calculate the functional form of \( p(a_0, b_0) \) in order to find agreement or disagreement with the Matalon-Packter law. The spacing coefficient, however, is obtained from the numerical solution of complicated coupled nonlinear differential equations and, as a consequence, the results are in an implicit form that is not particularly useful for deducing \( p(a_0, b_0) \). A notable exception is the simplest version of the ion-product theory which was used by Wagner\(^{11}\) to derive a result, \( p \sim a_0^{-3/2}, \) that is in variance with the Matalon-Packter law. Wagner’s result raises the question whether the present theories contain the Matalon-Packter law at all and our aim here is to discuss this question and make an attempt at answering it.

In attempting to derive the Matalon-Packter law, our basic aim is to keep the calculations and formulas simple so that the explicit dependence on \( a_0 \) and \( b_0 \) could be displayed. Accordingly, we have to make simplifying assumptions. The first two of these assumptions are about the experimental setup and the majority of the experiments satisfy the constraints described there. The remaining assumptions are based on observations of the time evolution of the concentrations of the reactants and reaction products obtained from numerical solution of the appropriate equations, from simulations of the process,\(^{17}\) and also from experience with the analytical solution of the problem of chemical reaction zone.\(^{24}\) We believe that the important features of the original problem are not lost after making the assumptions listed below.

1. The experimental setup can be described by one-dimensional reaction-diffusion equations (the gel is in a test tube whose length is much larger than its diameter).
2. The concentration of the outer electrolyte \( (a_0) \) is assumed to be much larger than that of the inner electrolyte \( (b_0) \) and it is also assumed that \( a(0,t) \) is kept fixed at \( x = 0 \), \( a(0,t) = a_0 \). In experiments one typically has \( 0.005 \leq b_0/a_0 \leq 0.1 \).
3. Reaction zones and the precipitation bands are much narrower than the diffusion length.
4. All concentration profiles can be approximated by straight lines in the neighborhood of the reaction zones and precipitation bands.
5. The slopes of the straight lines are determined by the diffusion lengths and by the asymptotic concentrations of the reacting species.
(6) An existing precipitation band acts as a sink [with $b(x_n,t>t_n)=0$ or $c(x_n,t>t_n)=0$ boundary condition] for the inner electrolyte and for the intermediate compound (this is an assumption about the reaction and aggregation rates being sufficiently large).

Using the above assumptions, we can derive relatively simple expressions for $p(a_0,b_0)$ for all three classes of theories (in Sec. II for supersaturation of ion-product, in Sec. III for nucleation and droplet growth, and in Sec. IV for induced sol-coagulation theories). The results are compared both with the corresponding numerical solutions of the nonsimplified models and with experimental data. Section V contains our final remarks.

In closing the Introduction, we list the symbols frequently used in the text:

- $a(x,t)$ - concentration of outer electrolyte, $A$;
- $b(x,t)$ - concentration of inner electrolyte, $B$;
- $a_0, b_0$ - initial concentrations of $A$ and $B$;
- $\kappa=b_0/a_0$ - dimensionless ratio of initial concentrations;
- $c(x,t)$ - concentration of sol (or intermediate compound);
- $D_A, D_B, D_C$ - diffusion coefficients of $A, B$, and $C$;
- $D_f$ - effective diffusion coefficient of reaction front ($A + B \rightarrow C$);
- $q^\ast$ - ion-product threshold;
- $c^\ast$ - nucleation (coagulation) threshold;
- $a^\ast$ - threshold for induced coagulation;
- $x_n$ - position of the $n$th precipitation band;
- $t_n$ - time of formation of the $n$th band;
- $1+p=x_{n+1}/x_n$ - spacing coefficient.

II. ION-PRODUCT SUPERSATURATION

Let us begin with the simplest case of the reaction $A + B \rightarrow D$. The ion-product supersaturation theory$^{10-13}$ is based on the assumption that precipitation of $D$ at time $t$ and at a site $x$ occurs if the product of ion concentrations $a(x,t)b(x,t)$ exceeds a threshold value

$$ a(x,t)b(x,t) \geq q^\ast. \tag{4} $$

In the mean-field theory, the equations describing this reaction-diffusion process are given by

$$ \frac{\partial a}{\partial t} = D_a \frac{\partial^2 a}{\partial x^2} - k \theta(ab-q^\ast) - \lambda abd, \tag{5} $$

$$ \frac{\partial b}{\partial t} = D_b \frac{\partial^2 b}{\partial x^2} - k \theta(ab-q^\ast) - \lambda abd, \tag{6} $$

$$ \frac{\partial d}{\partial t} = k \theta(ab-q^\ast) + \lambda abd, \tag{7} $$

where $D_A$ and $D_B$ are the diffusion constants of $A$ and $B$, and $\theta(x)$ is the step function describing an infinitely sharp threshold for precipitation with $k$ being the rate constant. The last terms on the right-hand sides represent the aggregation onto the existing precipitate of concentration $d(x,t)$. The initial conditions to the above equations are given by $a(x,0) = a_0 \theta(-x)$, $b(x,0) = b_0 \theta(x)$, and $d(x,0) = 0$.

These equations can be solved numerically, and our analytical expression found for $p(a_0,b_0)$ in the above process is in accord with the numerical results. Before deriving the results, however, let us show how the structure of the result may emerge from a simple dimensional analysis.

If we assume that both the precipitation ($k$) and aggregation ($\lambda$) rates go to infinity, then it follows from dimensional considerations that the dimensionless spacing coefficient can be expressed through the available dimensionless combinations of the parameters, and so we have

$$ p = P \left( \frac{b_0}{a_0} ; \frac{D_b}{D_a} ; \frac{q^\ast}{a_0b_0} \right). \tag{8} $$

Further simplification can be made by assuming that $P$ is not singular in its first argument and by setting this argument to zero since in experiments $b_0/a_0 \ll 1$. Then the Matelon-Packter law can emerge only if $P$ is linear in its third argument, thus giving

$$ p = P_1 \left( \frac{D_b}{D_a} \right) + P_2 \left( \frac{D_b}{D_a} \right) \frac{q^\ast}{a_0b_0}. \tag{9} $$

We emphasize that this is not a derivation of the desired result. This is, however, what we shall show to be valid in the experimentally investigated range ($0.05<p<0.4$) of possible $p$-s. Our calculation is based on a simple analytical approach combined with numerical evaluation of $P_1$ and $P_2$.

Figure 1 gives a characteristic picture showing the concentration profiles as well as the ion-product profile past the formation of the precipitation band $x_n$ and just before the appearance of the $(n+1)$th band. It is clear from the figure that the linearization of the concentration profiles is a good assumption, and, accordingly, in the neighborhood of $x_n$, we can approximate the concentrations as

$$ a(x,t) = a_0 \left( 1 - \frac{x}{\sqrt{2D_A t}} \right). \tag{10} $$
The condition for the \( n+1 \)-st band to appear is that the maximum of the \( ab \) product reaches the value \( q^* \). This leads to the following two equations:

\[
a(x_{n+1},t_{n+1})b(x_{n+1},t_{n+1}) = q^*,
\]

\[
\frac{\partial}{\partial t} [a(x,t)b(x,t)]_{t_{n+1},t_{n+1}} = 0.
\]  

The second equation gives a simple relationship between \( t_{n+1} \), \( x_n \) and \( x_{n+1} \):

\[
\sqrt{2D_a t_{n+1}} = 2x_{n+1} - x_n,
\]

which allows us to eliminate \( t_{n+1} \) from Eq. (12) and to obtain an equation for \( x_{n+1}/x_n \) and \( x_{n+1}/x_{n-1} \).

Assuming now that \( n \) is large enough that we can use the approximation

\[
\frac{x_{n+1}}{x_n} = 1 + p,
\]

and one obtains the following equation for \( p \),

\[
\Phi(p) = \frac{p^{3/2}(1+p)}{(1+2p)^2\sqrt{2+p}} = \frac{D_b}{D_a a_0 b_0} q^*.
\]

This equation clearly has a \( p > 0 \) solution for small values of \( \zeta = \sqrt{D_b/D_a} q^* (a_0 b_0) \). \( p \) increases with \( \zeta \) and goes to infinity as \( \zeta \to 1/4 \), accounting for the fact that no bands can be formed if \( a_0 b_0 \) is too small for \( ab \) to reach the critical value \( q^* \).

At first sight, the Matalon-Packter law does not follow from this theory, since \( p \) depends on \( a_0 \) and \( b_0 \) only through \( a_0 b_0 \). Furthermore, another problem is that, for small \( p \), we obtain Wagner’s result, \(^{11} \) namely the dependence of \( p \) on \( a_0 \) is of the form

\[
p \sim a_0^{-2/3},
\]

in contrast to the Matalon-Packter prediction \( p \sim 1/a_0 \). However, one should notice that the range of \( p \) where \( p \sim a_0^{-2/3} \) holds is certainly much smaller than the experimentally observed values \( 0.05 < p < 0.4 \). In this latter interval, \( \Phi(p) \) is very well approximated as a straight line (see Fig. 2) with parameters \( \Phi(p) = -0.0035 + 0.2p \). Thus, to a good approximation, we can write (16) as a Matalon-Packter law with \( F(b_0) = 0.02 \) and \( G(b_0) \sim 1/b_0 \),

\[
p = 0.02 + 5 \sqrt{\frac{D_b}{D_a a_0 b_0}} q^*.
\]

In this approach we obtain \( P_1 = \text{const.} \) and \( P_2 \approx \sqrt{D_b/D_a} \). Numerical solution of Eqs. (5) and (6) indicates that, in reality, the \( p \) is given to an excellent accuracy by

\[
p \approx 0.18 - 0.052 \left[ \frac{D_b}{D_a} \right]^{1/3} + \left[ 7.5 - 2.57 \left( \frac{D_b}{D_a} \right)^{1/6} \right] a_0 b_0 q^*.
\]

For \( D_a = D_b \), the order of magnitude for \( F(b_0) = 0.12 \) agrees with the order of magnitude values of \( F \) in experiments\(^6 \) and the \( 1/a_0 \) dependence is also the experimentally observed behavior. However, in experiments, \( F(b_0) \) is not a universal number. It changes not only as \( D_b/D_a \) is varied, but it also depends on \( b_0 \). Furthermore, the experimentally observed \( G(b_0) \) is also more varied than being just \( G(b_0) \sim 1/b_0 \). Thus we must conclude that although it is possible to derive the Matalon-Packter law from the ion-product supersaturation theory, the resulting functions \( F(b_0) \sim \text{const} \) and \( G(b_0) \sim 1/b_0 \) are too simple to account for experimental observations.

### III. NUCLEATION AND GROWTH

The theory at the next level of complexity assumes that the \( A + B \) reaction yields the precipitate \( D \) through an intermediate compound \( C \). This compound may, in principle, be a molecule \( AB \) but it may as well be a sol particle formed by \( A \) and \( B \) and, possibly, by some other background ions. All of this rather complicated situation is described by assuming that the \( C \)-s can be treated as diffusing species which precipitate if their concentration exceeds a threshold \( c^* \). The precipitate, \( D \), grows by collecting the neighboring \( C \)-s and the various theories differ in the sophistication of the description of this nucleation and growth process.\(^{16,17} \) In our case the nucleation threshold is sharp at \( c^* \) and precipitate will be assumed to be a perfect sink for the \( C \)-s.

In order to understand the formation of precipitate, we observe that the reaction zone (where the \( C \)-s are produced) is narrow\(^{24} \) and that \( c(x,t) \) reaches its maximum there (see Fig. 3 for characteristic concentration profiles just before the \( n+1 \)-st band forms). The \( n \)-th band acts as a sink and thus the \( C \)-s (or about half of them) formed at the front, \( x_f \), end up in this band.

With a good approximation, we can assume that at time \( t_{n+1} \), when \( c \) reaches the threshold value \( c^* \) and the \( (n+1) \)-th band is about to form, the concentration of \( C \)-s exhibits a triangularlike shape and it varies linearly between \( c(x_n) = 0 \) and \( c(x_{n+1}) = c^* \). As a consequence, we can estimate the current flowing to the \( n \)-th band as
Next we note that the front becomes quasistationary in the large-time limit (its velocity goes as \( \dot{x}_f \sim 1/\sqrt{t} \)) and so we can assume that the above expression for the current is valid in the whole time interval \([t_n, t_{n+1}]\) (numerical solutions of the reaction-diffusion equations support this assumption). Then the amount of \( C \) which disappears into the \( n \)-th band during the time interval \([t_n, t_{n+1}]\) can be computed as

\[
N_C \approx |j|(t_{n+1} - t_n) = D_c c^* \frac{t_{n+1} - t_n}{x_{n+1} - x_n}.
\]

On the other hand, \( N_C \) can also be estimated as the amount of \( C \)-s produced by the front \( \{c_0(x_{n+1} - x_n)\} \), where \( c_0 \) is a constant\(^{24} \) minus the \( C \)-s which are in the triangle shape, \( c^* (x_{n+1} - x_n) \),

\[
N_C \approx (c_0 - c^*) \cdot (x_{n+1} - x_n).
\]

Equating the two estimates of \( N_C \) then yields

\[
\frac{D_c c^*}{c_0 - c^*} = \frac{(x_{n+1} - x_n)^2}{t_{n+1} - t_n}.
\]

We can use now the fact that the band is formed at the front whose position, \( x_f \), is determined by an effective diffusion coefficient \( D_f \):\(^{24} \)

\[
x_{n+1} = x_f(t_{n+1}) = \sqrt{2D_f t_{n+1}},
\]

and find

\[
\frac{p}{1 + p/2} \approx p = \frac{D_c c^*}{D_f (c_0 - c^*)}.
\]

The parameters \( D_c \) and \( c^* \) in the above expression are material parameters. The dependence of \( p \) on \( a_0 \) and \( b_0 \) enters through the effective diffusion coefficient \( D_f \) of the front and through the concentration, \( c_0 \), of \( C \)-s left behind by the moving front. The remaining task is thus to determine \( D_f \) and \( c_0 \). We start with the case of \( D_a = D_b \), which can be discussed with relative ease.\(^{24} \) It has been shown for this case that \( D_f / D_a \) is determined from the following equation:

\[
\text{erf} \left( \frac{D_f}{2D_a} \right) = \frac{1 - \kappa}{1 + \kappa},
\]

where \( \text{erf}(x) \) is the error function\(^{26} \) and \( \kappa = b_0/a_0 \). The remarkable feature of the solution of this equation is that, for the experimentally relevant values \( 0.005 \leq \kappa \leq 0.1 \), the inverse of \( D_f \) is very well approximated by a linear function of \( \kappa \) (see Fig. 4),

\[
\left( \frac{D_a}{D_f} \right) \approx \eta_1 + \eta_2 \cdot \kappa = 0.158 + 4.03 \cdot \kappa.
\]

This is an important observation since it follows from (25) that \( p \sim 1/D_f \) for small \( p \) and the Matalon-Packer law will emerge as a consequence of the above equation.

The general case \( D_a \neq D_b \) is more complicated but the results are similar. The effective diffusion coefficient, \( D_f \), is obtained from solution of the following equation.\(^{25} \)

\[
H \left( \sqrt{\frac{1}{2} \frac{D_f}{D_a}} \right) = a_0 \frac{D_f}{D_a} H \left( \sqrt{\frac{D_a}{2D_b}} \right),
\]

where \( H(x) = \left[ 1 - \text{erf}(x) \right] \exp(x^2) \). The behavior of \( D_a / D_f \) as a function of \( \kappa \) for \( D_a \neq D_b \) is similar to that seen in Fig. 4. One can observe that \( D_a / D_f \) is linear in \( \kappa \) in the experimentally relevant region and thus Eq. (27) emerges again but the constants \( \eta_1 \) and \( \eta_2 \) are now dependent on \( D_b / D_a \).

We turn now to the calculation of \( c_0 \). The starting point is the observation that the width of the reaction zone is much smaller than the diffusion length\(^{24} \) and thus the reaction zone can be approximated as a point at \( x_f(t) = \sqrt{2D_f t} \) where both concentrations \( a \) and \( b \) approach zero. This means that the field \( b(x,t) \) satisfies the diffusion equation with the reaction term replaced by the boundary condition \( b(x=x_f(t),t)=0 \) and the other boundary condition being \( b(x\to\infty,t) = b_0 \). Once this moving boundary problem is solved, the number of \( C \)-s, \( N_C \), produced up to a given time is obtained by the time integral of the current \( j_b \equiv D_b \partial b/\partial x \) evaluated at \( x_f \). Finally, \( c_0 \) is found by dividing \( N_C \) by the advance of the front in time \( t \), \( c_0 = N_C / x_f \) (here we use the fact that the density of \( C \)-s produced by the front is constant in space\(^{24} \)). The result of this calculation is given by
For $D_a = D_b$ and $\kappa \ll 1$ we can use Eq. (26) in conjunction with the asymptotics of the error function $^26$ to obtain $c_0 = b_0$. The physical meaning of this result is clear. For $\kappa \ll 1$, we have $D_b \ll D_f$ and thus the front moves fast into the region of $B$-s and, consequently, the $B$-s can be treated as immobile particles yielding $c_0 = b_0$. Corrections to the $c_0 = b_0$ result can also be calculated using again the asymptotics of the error function. $^26$ To first order in $D_b/D_f$, we find

$$c_0 \approx b_0 \left( 1 + \frac{D_b}{D_f} \right) = b_0 \left( 1 + \frac{D_a}{D_f} - \frac{D_b}{D_f} \right). \tag{30}$$

We can use now the linearity of $D_a/D_f$ in $\kappa = b_0/a_0$ to obtain the following parametrization for $c_0$:

$$c_0 = b_0 (\sigma_1 + \sigma_2 \kappa), \tag{31}$$

where $\sigma_1 = 1 + \eta_1 D_b/D_a$ and $\sigma_2 = \eta_2 D_b/D_a$ are numbers of the order of 1. Since $\sigma_1$ and $\sigma_2$ are of the same order of magnitude and since $\kappa \ll 1$, in the following we shall neglect the $\kappa$ term in (31),

$$c_0 \approx \sigma_1 b_0. \tag{32}$$

It is important to note here that a similar omission of the $b_0/a_0$ term would not be justified in the linearized form of $D_a/D_f$. $^27$ There we have a constant term, $\eta_1 \approx 0.2$, that is an order of magnitude smaller than the coefficient, $\eta_2 \approx 4$, in front of $b_0/a_0$. Thus, for the relevant values of $b_0/a_0$, the two terms contribute equally.

The expressions (30), (31), and (32) remain good approximations up to the point where $D_b/D_f \approx 1$. This happens, however, only at rather large values of $D_b/D_a$ for $\kappa \ll 1$. Since the diffusion coefficients of usual electrolytes in aqueous solutions are usually within a factor 2 of each other$^29$ and their ratios rarely exceed 5, $^28$ we believe that the approximation (31) can be used for realistic experimental setups.

Having determined $D_f$ and $c_0$, we can now return to Eq. (25) for $p$ and find again a version of the Matalon-Packter law,

$$p = \frac{D_c e^* \eta_1}{D_a (\sigma_1 b_0 - e^*)} + \frac{D_c e^* \eta_2 b_0}{D_a (\sigma_1 b_0 - e^*) a_0}$$
$$= F(b_0) + \frac{G(b_0)}{a_0}. \tag{33}$$

In order to gain confidence in the above result, we have calculated $p$ for $\kappa$ in the experimental range $0.005 < \kappa < 0.1$ by using the appropriate reaction-diffusion equations $^3\text{Eqs. (5), (6), and (7)}$ must be modified and supplemented by another equation in order to take into account the nucleation and growth processes described at the start of the section (for details see Ref. 17). The results are displayed in Fig. 5 where the Matalon-Packter law derived above (solid line on the figure) is shown to perform very well considering the simplicity of the derivation. It should be noted that nonlinear dependence on $\kappa$ sets in for $\kappa \approx 0.06$ and that taking into account the $\kappa$-dependence of $c_0$ (dashed line) modifies the straight line with the right curvature.

One can observe from Eq. (33) that the nucleation and growth theory gives more complicated $b_0$ dependences for $F(b_0)$ and $G(b_0)$ than the ion-production theory. Power law forms for $F$ and $G$ are found in the limit of $c^* \ll b_0$ (which might be the experimental limit) where one obtains $F(b_0) \sim 1/b_0$ and $G(b_0) \sim 1/b_0$ constant. It is remarkable that these are similar to the ion-product results but with the roles of $F$ and $G$ interchanged. Comparing the two theories, we see that the nucleation and growth theory performs better in the sense that it can produce power law behavior for $F(b_0)$ as observed in some experiments and, at the same time, has a $G(b_0)$ which depends on $b_0$ more weakly but it is a decreasing function of $b_0$ in agreement with experiments.

In conclusion, the nucleation and growth theory provides us with a Matalon-Packter law that is closer to experiments and, perhaps, describes some of them. Nevertheless, the functions $F$ and $G$ appear to be too rigid to accommodate all experimental findings. It should also be noted that this theory has a prediction $G(b_0)/F(b_0) \sim b_0$ that is experimentally easily distinguishable from the prediction of the ion-product theory, $G(b_0)/F(b_0) \sim 1/b_0$.

IV. INDUCED SOL-COAGULATION

The induced sol-coagulation theory $^{14,15}$ is a generalization of the nucleation and growth theory. The sol, C, is produced at the reaction front in the $A + B \rightarrow C$ reaction and it flocculates if the following two conditions are satisfied. First, $c$ must exceed a supersaturation threshold $c^*$, and, second, the concentration of the outer electrolyte, $a$, must be above the critical coagulation concentration threshold, $a^*$. The second condition arises in systems where the $A$ ions screen the repulsive electrostatic interaction among the sol particles.

Figure 6 shows characteristic concentration profiles in this process just as the $n + 1$-st band is appearing. The first
The remarkable feature of this picture is that the reaction front is way ahead of the precipitation zone (this does seem to happen in some experiments\(^{(39)}\)). This is understandable, since \(a(x_f, t) \to 0\) at the front, while at the place where precipitation occurs one must have \(a(x_{n+1}, t_{n+1}) = a^*\). The second important feature is that the reaction front leaves \(c^*\) behind at a fixed concentration \(c_0\), as already discussed in the previous section. In order that precipitation could occur, the parameters must be assumed such that \(c_0 > c^*\). Further assuming that the sol does not diffuse too fast \((D_s \ll D_a)\), one of the conditions for precipitation \((c \geq c^*)\) is always satisfied behind the front and far away from the last existing band \((x_n)\).

Consequently, the position, \(x_{n+1}\), where the next band appears will be determined by the arrival of the concentration “front” \(a = a^*\) to a position where \(c = c^*\). The \(a\) and \(c\) profiles near \(x_n\) can be written as \(a = a_0(1 - x/x_f)\) and \(c = c_0(x - x_n)/\sqrt{2D_c(t - t_n)}\), the above conditions yield

\[
\frac{c_0(x_{n+1} - x_n)}{\sqrt{2D_c(t_{n+1} - t_n)}} = c^* ,
\]

\[
a_0 \left(1 - \frac{x_{n+1}}{x_f(t_{n+1})}\right) = a^* ,
\]

where \(x_f(t) = \sqrt{2D_f t}\) is the position of the reaction front at time \(t\).

In order to calculate \(p\), we note now that Eq. (35) yields

\[
\frac{x_{n+1}}{x_f(t_{n+1})} = 1 - \frac{a^*}{a_0} = \frac{x_n}{x_f(t_n)} .
\]

The above equation can then be used in conjunction with \(x_f(t_{n+1}) = \sqrt{2D_f t_n}\) to obtain \(p\) from Eq. (34),

\[
p \approx \frac{2D_c a^*}{D_f c_0^2 (1 - a^*/a_0)^2} .
\]

Since the values of \(D_f\) and \(c_0\) come from reaction \(A + B \to C\) which is decoupled from the next stages of nucleation and growth processes, the calculations and approximations of these quantities can be taken over from Sec. III. Moreover, we shall assume that \(a^*\) is much smaller than \(a_0\) so that the right-hand side of Eq. (37) can be expanded in \(a^*/a_0\). There is no experimental information on \(a^*\) and, although \(a^*/a_0\ll 1\) seems to be a natural assumption, we must be cautious with the \(a^*/a_0 \to 0\) limit. The problem is that the picture we are working with for the induced sol-coagulation theory (Fig. 6) is not valid for arbitrary small \(a^*\). Indeed, the fact that the position of the \(A + B \to C\) reaction front, \(x_f\), is far ahead of the point of formation of the \(n+1\)-st band means that diffusion length of the \(C\) particles is smaller than \(x_f(t_{n+1}) - x_n\).

\[
\sqrt{2D_c(x_{n+1} - x_n)} < x_f(t_{n+1}) - x_n .
\]

Using Eqs. (36) and (37) and neglecting higher order terms in \(p\) and \(a^*/a_0\), the above equation yields

\[
\frac{D_c a^*}{D_f c_0} < 1 .
\]

The meaning of this result is the following. The place where the new band nucleates moves to the \(A + B \to C\) reaction zone and thus the inequality (38) gets violated upon increasing \(D_c\) or \(c_0\) and decreasing \(a^*\). It is clear thus that, at fixed values of the other parameters, \(a^*/a_0\) cannot be taken to zero.

The lack of the limit \(a^*/a_0 \to 0\) has the consequence that the result for the nucleation and growth theory (33) cannot be obtained from (37). The matter of finding an expression for \(p\) valid in the \(a^*/a_0 \to 0\) limit and reducing to (33) appears to be a rather difficult task since various length scales become comparable approaching this limit. We have not been able to obtain such an interpolating formula.

Returning to (39), we see that the material parameters can be such (e.g., if \(c^*/c_0 \ll 1\)) that \(a^*/a_0\) can be small without the inequality (39) being violated. For such a range of parameters we can expand (37) in \(a^*/a_0\) and obtain again a version of the Matalon-Packeter law,

\[
p = \frac{2D_c a^* a_0}{D_f c_0^2 b_0^2} + \frac{2D_c a^*}{D_f c_0^2 b_0^2} (2 \gamma_1 a^* + \gamma_2 b_0) .
\]

The spacing coefficient (40) can again be compared with that obtained from the appropriate reaction diffusion equations (the equations for the nucleation and growth theory must be augmented with the condition that nucleation can occur only if \(a > a^*\)). An exhaustive numerical study is practically impossible due to the number of parameters in the problem \((\kappa, a^*/a_0, c^*/a_0, D_b/D_a, D_c/D_a)\), plus the rates of reaction, nucleation, and aggregation. One can easily miss regimes of nontrivial behavior in this high dimensional parameter space and we claim with this numerical study only that the Matalon-Packeter law as given by Eq. (40) is indeed observed for a reasonable range of parameters (Fig. 7).

As one can see from (40), the induced sol-coagulation theory provides us with

\[
F(b_0) = 1 - b_0^2 , \quad G(b_0) = C + \alpha b_0^2 .
\]

The power law form \(F(b_0) \sim b_0^{-2}\) is close to what has been observed in some experiments and the fact that \(G(b_0)\) is a decreasing function of \(b_0\) is also in accord with the observations. It should also be clear that a more precise
theory of the induced sol-coagulation process should reproduce, in the $a^2/\alpha_0 \to 0$ limit, the result $F \sim b_{0,1}^{-1}$ obtained in the nucleation and growth theory. Thus, assuming that the crossover between the $b_{0,2}^{-1}$ and the $b_{0,1}^{-1}$ behaviors is smooth, one should be able to find regimes where $F(b_0) \sim b_{0,1}^{-1}$ with $1 \leq \gamma \leq 2$. Since this covers a large portion of the experimentally observed range $0.2 \leq \gamma \leq 2.7$, we conclude that the induced sol-coagulation theory provides the best description of the pattern formation in Liesegang phenomena.

V. FINAL REMARKS

Our main results are summarized in Eqs. (19), (33), and (40) giving the Matalon-Packter law for the three main theories, respectively. Comparing these formulas, we arrived at our main conclusion, namely that the induced sol-coagulation theory is the best in describing the experimental observations on the spacing coefficient of Liesegang patterns.

There is no doubt that the arguments used in the derivation of $p$ can be refined and made more precise. The main aim of our work, however, was the demonstration that the Matalon-Packter law can be understood in terms of simple pictures for which it is possible to develop analytical arguments. We hope that one can build on these results and achieve a better understanding of Liesegang phenomena. In particular, it would be important to find a description of the induced sol-coagulation process which contained the crossover to results for the nucleation and growth model. Furthermore, we have just considered the simplest cases associated with the reaction $A + B \to C$. In experiments, equally important are the $A_2 + B \to C$ and $A + B_2 \to C$ cases and, obviously, one should also explore the possibilities associated with these and, possibly, more complicated reaction schemes. Finally, an interesting and important test of the conclusion that the induced sol-coagulation theory is preferable in the studies of Liesegang phenomena would be the application of this theory for the quantitative description of revert patterns.

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FIG. 7. Spacing coefficient for the induced sol-coagulation theory. The dependence of $p$ on $k = b_0/\alpha_0$ is displayed for the following choice of parameters: $D_x/D_\infty = 1, D_y/D_\infty = 0.3, c^2\alpha_0 = 0.86, a^2\beta_0 = 7.8$, and with all the reaction rates taken to be large. The dashed line is the "linear" Matalon-Packter law (40) while the solid line is the "nonlinear" version of (40) where the $k$-dependence of $c_0$ (31) is kept when going from Eq. (37) to Eq. (40).