Convective Instabilities Derived from Dissipation of Chemical Energy

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ABSTRACT
Oxidation reactions of a series of organosulfur compounds by chlorite are excitable, autocatalytic, and exothermic and generate a lateral instability upon being triggered by the autocatalyst. This article reports on the convective instabilities derived from the reaction of chlorite and thiourea in a Hele-Shaw cell. Reagent concentrations used for the development of convective instabilities delivered a temperature jump at the wave front of 2.1 K. The reaction zone was 2 mm and due to normal cooling after the wave front, this induced a spike rather than the standard well-studied front propagation. Localized spatiotemporal patterns develop around the wave front. This exothermic autocatalytic reaction has solutal and thermal contributions to density changes that act in opposite directions due to the existence of a positive isothermal density change in the reaction. The competition between these effects generates thermal plumes. The fascinating feature of this system is the coexistence of plumes and fingering in the same solution as the front propagates through the Hele-Shaw cell. Wave velocities of descending and ascending fronts are oscillatory. Fingers and plumes are generated in alternating frequency as the front propagates. This generates hot and cold spots within the Hele-Shaw cell, and subsequently spatiotemporal inhomogeneities. The small ΔT at the wave front generated thermocapillary convection which competed effectively with thermogravitational forces at low Eötvös numbers. A simplified reaction-diffusion-convection model was derived for the system. Plume formation is heavily dependent on boundary effects from the cell dimensions.

This article reports of self-organization derived from the dissipation of chemical energy. The unique aspect of this article is as follows: generally, addition of convection to reaction-diffusion mechanisms destroys most coherence and self-organization. It is not so, as reported in this article. The delicate balance between thermogravitational, thermocapillary diffusion effects can deliver remarkable self-organization in ungelled environments.

INTRODUCTION
Pure reaction-diffusion structures and propagation have been extensively studied and are well-known from the basis of a Turing instability.1,2 The presence of a reaction front is also a well-known feature in many physical, biological, and chemical processes. For example, a simple chemical reaction type process of \( A + B \rightarrow C \) will exhibit a front, a spatially-localized region where production of \( C \) can be achieved, provided diffusing reagents are initially separated in space.3 Even if \( A \) and \( B \) are initially not separated, a lateral instability can still emerge if the prevailing reaction is excitable, e.g., autocatalytic.4 When a spatially extended system becomes unstable, the resulting dynamics have sensitive dependence on whether the system is convectively unstable, in which case perturbations grow in time but convected away fast enough that they die at each fixed position in the lab frame considered, or absolutely unstable, in which there arises a perturbation and a location where this perturbation does not decay. With an autocatalytic reaction system, a perturbation using an autocatalyst does not decay and will grow until diffusion or convection-limited. The first stage in understanding pattern formation in such reactive flows is to understand the description and calculation of the properties of the reaction zone. This involves answering the question of where and at what rate does the reaction product appear. This problem has been studied with respect to Liesegang ring band formation in which simplified theories have predicted band formation solely from properties of the front and phenomenological theories of diffusion-limited aggregation.5,6 The width of the reaction zone...
This fascinating cooperativity has been observed in real-life phenomena in engineering systems, geological formations, supernovae dynamics, CO$_2$ sequestration, earth mantle convection, and atmospheric and ocean layering convective motions. 

This article assesses the active role that chemical reactions can have on parameters of flow dynamics in chemical reactive flows. We have, previously, been able to detail the mechanism of the "drive" reaction that drives the hydrodynamic instabilities reported here.\textsuperscript{11-13} The reaction used is the autocatalytic chlorite-thiourea reaction.\textsuperscript{11,14} Autocatalytic reactions coupled to diffusion can generate traveling fronts of constant speed whereby products of the reaction invade and progressively consume reactants in space and time.\textsuperscript{15} In the absence of a gelled environment, convection can deform such fronts due to density differences between reactants and products. Density differences can result in differences in molar volumes of products and reactants. This is especially so in the chlorite-thiourea reaction where products such as sulfate and chloride are dense ionic products while reactants are lighter organic reagents. The exothermicity of the reaction, however, can temporarily "flip" this density relationship until thermal equilibrium is attained.\textsuperscript{16} Thus, convective motions will appear around descending fronts since we have a dense solution overlying a lighter one (reactants). This is an unstable front since any deformity of the surface will tend to grow. A Rayleigh-Taylor instability (RTI) in which denser solution sinks into the rising less dense reactant solution will deform the interface into density fingering since the density gradient points opposite to the body force. This is due to the buoyancy-induced convective deformation of a reaction-diffusion front. The combination of convection to a reaction-diffusion front results in reaction-diffusion-convection dynamics. Our preliminary experiments have shown that addition of heat and, subsequently, convection (both Marangoni and Rayleigh-type) have not produced more cooperativity, but more complexity due to, among others, interfacial effects which can introduce stochasticities.\textsuperscript{17} Universally, after an induction period, all thiocarbamide-oxyhalogen systems produce a large pH change (as H$^+$ ions are rapidly produced) as well as large heat generation. The initial oxidation of the S(-II) species to S(0) (e.g., sulfenic acids) has a negative acid feedback loop, but subsequent oxidations all the way to S(VI) produce a positive acid feedback loop.\textsuperscript{11} The only prerequisite for the observations of the full range of exotic dynamics is that the oxidant (chlorite) should be in stoichiometric excess over the reductant.\textsuperscript{11} The advantage, then, is that since [H$^+$] is allowed to vary and is determined by the reaction's stoichiometry, then an acid propagation wave can be observed and used to follow the lateral instability through an acid-base indicator. This approach has also been utilized in following waves generated in the chlorite-tetrathionate (CT) reaction.\textsuperscript{24-26} The stoichiometry of the ClO$_2^-$-thiourea reaction is\textsuperscript{11,12,27}

\begin{equation}
2\text{ClO}_2^- + \text{SC(NH}_2)_2 + \text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + \text{OC(NH}_2)_2 + 2\text{Cl}^- + 2\text{H}^+.
\end{equation}

\textbf{Propagating species: ClO}_2(aq), SO$_4^{2-}$, H$^+$, Q (heat)

We have derived a general mechanism to account for most chlorite-based oxidations of sulfur compounds, and it appears the chlorite-thiourea reaction can be explained, in general terms by this model. Reaction (R2) is a general stoichiometric equation for the oxidation of thiocarbamide compounds by chlorite.\textsuperscript{17} The oxidation products are sulfate and a urea-type residue, $R_1R_2C=O$,

\begin{equation}
R_1R_2C=S + 2\text{ClO}_2^- + \text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + R_1R_2C=O + \text{pH}^+ + 2\text{Cl}^-.
\end{equation}

These reactions show common clock reaction characteristics in which initially there is a quiescent period with no activity in the reaction indicators. At the end of the induction period, there is a sudden production of sulfate, SO$_4^{2-}$, chlorine dioxide, ClO$_2$, and acid, H$^+$. The acid and sulfate are formed simultaneously from the oxidation of the sulfonic acid intermediate, $R_1R_2\text{CSO}_3\text{H}$,

\begin{equation}
R_1R_2\text{CSO}_3\text{H} + 2\text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + R_1R_2C=O + 4\text{H}^+ + 2\text{e}^-.
\end{equation}

Chlorine dioxide is formed by an extraneous reaction between the excess chlorite, ClO$_2^-$, and left-over autocatalyst: hypochlorous acid, HOCl,\textsuperscript{23}

\begin{equation}
2\text{ClO}_2^- + \text{HOCl} + \text{H}^+ \rightarrow 2\text{ClO}_2(aq. \text{Yellow}) + \text{Cl}^- + \text{H}_2\text{O}.
\end{equation}

HOCl, being the autocatalytic species, controls the rates of reaction and wave propagation under isothermal conditions. The mechanism for autocatalysis involves the reactive intermediate Cl$_2$O$_2$ leading to quadratic autocatalysis\textsuperscript{24}

\begin{equation}
\text{ClO}_2^- + \text{HOCl} + \text{H}^+ \rightarrow \text{Cl}_2\text{O}_2 + \text{H}_2\text{O} \quad \text{(R5)}
\end{equation}

followed by

\begin{equation}
\text{Cl}_2\text{O}_2 + 2\text{e}^- + 2\text{H}^+ \rightarrow 2\text{HOCl}.
\end{equation}

So, for almost all our modeling studies, the rate of reaction can be simplified into $\text{Rate} = k[A][B]$ for a standard A + B reaction where B represents the autocatalyst.

Apart from H$^+$, SO$_4^{2-}$, and ClO$_2^-$, the most important product parameter in terms of wave propagation dynamics is a large....
and almost instantaneous heat evolution. The heat generated is very high, such that the reaction of thiourea and chlorite has a reaction enthalpy in excess of \(-1170\) kJ mol\(^{-1}\).\(^{14,29}\) The transformation from purely organic reagents to mostly inorganic products; sulfate, H\(^+\), Cl\(^-\) results in high energies of solvation and an increased solution density. By adjusting initial reagent concentrations, temperature jump for our reaction system could be maintained between \(\Delta T = 2.2–3.8\) K. This is the range that produced the most exotic wave and front dynamics. The unreacted solution far ahead of the wave front maintained the room temperature while the leading edge of the wave front could be 2.2–3.8\(^\circ\) above this temperature. The reaction zone is a thin sliver of approximately 2 mm.\(^{10}\) The experimentally-determined reaction zone and temperature profile are shown in Figs. 1 and 2. This was obtained using thermocouples lined along the wave front.

**INDICATORS**

There are several physical parameters that can be utilized to follow the development of patterns and/or waves in this reaction system. Apart from neutral density polystyrene beads, there are four possible indicators that can be used to follow the traveling wave and the subsequent spatiotemporal patterns: acid-base, BaCl\(_2\), ClO\(_2\) color, and freshly-prepared starch. Each has a specific role that the others cannot duplicate. BaCl\(_2\) gives a white precipitate of BaSO\(_4\), which is produced in reaction (R3).\(^{31}\) This article dwells only on the use of acid-base indicators. This method is profiled below: in unbuffered solutions, protons produced by the reaction can lead to large changes in pH of the solution. Typical pH values are in the range of approximately 5.5 in reactant solutions and 1.5–3.5 in product solutions [initial reactant concentrations determine the amount of H\(^+\) ions formed, see stoichiometry (R2)]. Acid-base indicators are best suited for the study of front propagation and spatiotemporal patterns due to their vivid color at low micromolar concentrations. Acid-base indicators are also very good at being able to characterize the reaction zone of the wave front: intensity and degree of color change with indicators methyl red and bromophenol blue is indicative of the extent of reaction. This gives a very good estimate of the reaction front (which we have so far estimated at about 2–3 mm;\(^{18}\) exact values for conditions utilized for our experiments were obtained in this study, see Fig. 1). Acid-base indicators are not effective in aiding in the observation of hydrodynamic patterns since they form a homogeneous solution with a reaction solution.

**EXPERIMENTAL SECTION**

**Materials**

The following analytical grade reagents were used without further purification: thiourea (Aldrich); perchloric acid 70\%–72\%, bromophenol blue, methyl red, and methyl orange (Fisher). The structures of these three dyes used are shown below. Sodium chlorite is sold in its technical grade form (about 80\%). A single recrystallization (ethanol-water mixture) brought the assay to >98\%. The analysis of sodium chlorite was performed iodometrically by adding excess acidified iodide and titrating the liberated iodine against standard thiosulfate with freshly-prepared starch as indicator.\(^{32}\)
can be integrated as an initial value problem, giving

Previously, we had established that, in shallow layers, 1.7 mm < d < 2.3 mm, wave front propagation is proportional to the temperature jump at the wave front, ΔT. The temperature jump is proportional to the heat production by the reaction Q. Q, in turn, is a function of the extent of the reaction, \( \xi \),

\[
Q = f(\Delta H_R, \xi, c_\infty).
\]

\( \Delta H_R \) is the enthalpy change of the reaction and \( c_\infty \) represents initial concentrations subject to stoichiometric constraints that the ratio of oxidant to reductant is greater than 2. Thus, at constant initial concentrations, the variable which controls amount of heat generated within the reaction zone is the extent of reaction, \( \xi \). \( \xi \) varies from zero just in front of the wave to 1.0 at the end of the reaction zone. The maximum value of ΔT can be derived from the heat capacity of the reaction solution in the limit of extent of reaction approaching unity. The highest temperature is recorded just behind the wave front, and, subsequently, a drop in temperature is recorded based on Newton’s law of cooling. In shallow layers, the wave accelerates until it attains a constant lateral velocity. Since the Marangoni effect dominates in shallow layers, the wave velocity is, thus, determined by the rate of reaction and thus reaction kinetics. This bistable autocatalytic chlorite-thiourea reaction has the extent of reaction that is a nonlinear function of the instantaneous chemical reactant concentrations. The autocatalysis term will dominate the kinetics of the reaction; this is the R5 + R6 sequence. The reaction that delivers acid that determines the position of the front is the oxidation of the sulfonic acid in reaction (R3),

\[
R_1R_2^\prime CSO\cdot H + HOCl + H_2O \rightarrow SO_4^{2-} + R_1R_2^\prime C=O + 3H^+ + Cl^-. \tag{R7}
\]

Thus, the rate of formation of acid is dependent on the rate of formation of HOCl. The R5 + R6 can be summarized as

\[
A + B \rightarrow 2B, \text{ with } B = HOCl. \tag{R8}
\]

The rate of formation of HOCl is

\[
\text{Rate} = k[A][B], \text{ initial concentrations } a_0, b_0. \tag{2}
\]

Equation (2) can be integrated as an initial value problem, giving the concentration of HOCl at any time as

\[
[HOCI](t) = \frac{(a_0 + b_0)}{1 + \left(\frac{b_0}{a_0}\right) e^{-k(a_0 + b_0)t}}. \tag{3}
\]

Equation (3) is a sigmoidal trace which is limited by the diffusion of reactants in the reaction zone and the depletion of the substrate. Thus, we expect an ever-increasing rate of reaction which tails off when thiourea is depleted. These kinetics are much more complex than those expected from the standard \( A + B \rightarrow \text{ products} \), which has been the staple of most previous studies of chemical reactive flows.
WAVE FRONT DYNAMICS

Our studies have three different waves in one wave front. The leading front is the acid wave, determined by the change in pH indicator. This is derived from reaction (R3). The second wave can be visually observed by barium sulfate precipitation in reaction solutions spiked with barium chloride. This is also derived from reaction (R3), but the proton’s diffusion coefficient means that the BaSO4 wave trails behind the reaction front. The third wave is of yellow chlorine dioxide, derived from reaction (R4). The chlorine dioxide wave could be enhanced by the addition of freshly-prepared starch indicator preserved with HgI2. The acid front, determined by the acid-base indicator, was then annihilated by bleaching from chlorine dioxide. Thus, a colorless solution emerges way behind the leading wave front.

EXPERIMENTAL MEASUREMENTS

There are several parameters that are important in determining observed spatial inhomogeneities, most important being the initial reagent concentrations as they determine the amount of heat produced which will fuel wave propagation. Our initial studies of wave propagation had involved the following initial concentrations: [NaClO3]0 = 4.00 × 10−3 M, [CS(NH2)2]0 = 2.00 × 10−4 M. Upon triggering the reaction, the temperature increases, with ΔT = 3.3 K. This was also collaborated through calculations from the expected heat of reaction, heat capacity of the vessel, and assuming reaction solution has the same specific heat as water. The solution depth was varied from 1.7 mm to 3.5 mm. Depths shallower than 1.1 mm could not deliver a completely flat surface due to the surface tension between the plexiglass and the reaction solution. Depths of 1.3 mm < d < 2.1 mm delivered a lateral instability of an acid with no structure. Wave velocity commenced with a rapid acceleration followed by a steady wave velocity as would be predicted by the rate of formation of HOCl in Eq. (3). The isothermal density change was measured as Δρ = 2.8 × 10−3 g cm−3. The change in surface tension with temperature, as measured by the capillary rise method, ∂σ/∂T = −9.5 × 10−5 N m−1 K−1. A negative value for ∂σ/∂T is expected for most liquids. ∂σ/∂T was measured only for the product solution. The expansion coefficient for the product solution, ∂ρ/∂T, was determined experimentally as −0.4 kg m−3 K−1.

VIDEO IMAGING TECHNIQUES

In previous work performed in our laboratory, the pH value of solution varied from 5.5 to approximately 2.6. Such a range was amenable to the observation of acid front by the use of methyl orange and methyl red indicators. In the reported experiments in this article, the final pH varied between 2.6 and 3.8. While methyl orange/red could catch the lowest parts of the pH range, they were not as effective in the higher ranges. Thus, a mixture of the indicators, methyl orange and bromophenol blue, was used to determine not only the wave front but also the extent of the reaction in the reaction zone before bleaching by chlorine dioxide occurred. Based on the mixture of pH indicators used, the unreacted solution had a bluish-green color, the wave front is red, and it varies in reddish intensity based on the extent of reaction. Chlorine dioxide formation in the wake of the acid front, as mentioned previously, bleaches the indicators to a white color.

EXPERIMENTAL RESULTS

All experiments reported here were run with the following initial reagent concentrations: [CS(NH2)2]0 = 1.68 × 10−3 M, [ClO2]0 = 4.00 × 10−3 M for an oxidant to reductant ratio of 24:1. This ensured the total consumption of the thiourea. Product solutions contained chloride (excess), chloride, chlorine dioxide, sulfate, and acid. These experiments were not buffered so that acid variations could be used to determine reaction front and rate. Initiation was solely through the addition of autocatalyst at the edge of the Hele-Shaw cell. Same wave dynamics were observed from electrochemical initiation when a platinum electrode is placed on one end of the vessel and a voltage of 3.0 V is passed through. The autocatalyst, HOCl, was added in the form of a basic solution of chlorine water through a micropipette.

Added at the edge of the Hele-Shaw cell, the autocatalyst immediately commenced the reaction, and this was observed through the immediate acid formation [reaction (R3)] and yellow chlorine dioxide [reaction (R4)]. Due to the positive isothermal density change, the acid wave descended down the cell to the bottom of the cell. The proximity to the edge of the vessel ensured efficient heat exchange of the warm front with the ambient temperature. The efficient heat exchange with the unreacted cold solution ensured its rapid finger down the cell. This was the initiation of the rest of the wave front dynamics reported in this article. Figure 4 shows a sketch of all the relevant and expected forces in this reacting-diffusing-convecting medium. Our previous studies had shown a dominant Marangoni effect at the surface in shallow layers. The surface tension gradient on the free surface will induce a top layer which has to be materially balanced by a return back flow at the bottom of the fluid. Villiers and Platten1,2 modeled this flow for high Prandtl number fluids, Pr ≥ 4, and evaluated a horizontal velocity profile of

\[ V_x = h(h - 2/3); 0 \leq h \leq 1, \quad (4) \]

where \( V_x \) is the horizontal velocity and \( h \) is the normalized solution depth. The sketch of this flow is shown in Fig. 3 for depths between

![FIG. 3. Calculated 2-axis velocity profile for a Benard-Marangoni instability with lateral heating but with no reaction. It is a single-component system (acetone), and calculated for Pr = 4.24. The lower back flow is necessary for material balance.](image-url)
1.7 mm and 3.3 mm. These shallow layers had a dominance of thermocapillary effects. Figure 4 shows fluid flow with a reaction solution with reagent concentrations that gave a front temperature jump of 3.3 K. The thin sliver of product solution pulled forward due to thermocapillary effects is brought to thermal equilibrium with unreacted bulk solution resulting in fingering regime of double-diffusive convection. As the finger proceeds down the bulk unreacted solution, it is swept backward in accordance to expected fluid flow depiction in Fig. 3. Depths and values of $\Delta T$ could be adjusted to deliver various patterns and fluid flow dynamics. These patterns include thermal plumes and convective tori with entrainment of fluid flow in concentric circular patterns.

**Self-organization in 20 mm depth Hele-Shaw cells**

This article reports mainly of fluid flow patterns generated from 20 mm depths of a 260 mm long Hele-Shaw cell. The reaction solution used, coupled with the heat capacity of the apparatus, gave a traveling wave with a maximum temperature jump after the complete reaction, of 2.1 K. This value of $\Delta T$ is lower than the temperature jump for solutions that gave convective tori and thermal plumes. Thus, these experiments were performed at higher Eötvös numbers than those previously reported,

$$E_0 = \Delta \rho gd^2 / \sigma. \quad (5)$$

This Eötvös number is effectively a ratio of thermogravitational and thermocapillary forces. The lower reduced Marangoni effect delivered hitherto unobserved patterns of self-organization: spatiotemporal inhomogeneities and Rayleigh-Taylor instabilities. Figure 6 shows the first waves generated after the initial initiation of the wave and full descent of the initial finger on the side of the container, $x = 0$. Finger A (faint) originates from the top of the solution and immediately fingers down the bulk of the solution. As opposed to Fig. 5, the initial wave at the top of the solution immediately starts to sink into the bulk solution without extensive movement in the x-direction. Finger A’s forward velocity stalls at 0.858 mm min$^{-1}$ as the mass of products behind the wave front cools and increase their specific gravity relative to the unreacted bulk solution. The descending wave velocity is oscillatory as it descends through the 4 images [Figs. 6(a)–6(d)]. It commences at 3.78 mm min$^{-1}$, accelerates to 3.87 mm min$^{-1}$, and finally decelerates to 3.83 mm min$^{-1}$ as it approaches the bottom of the vessel. This deceleration is due to the proximity of the other hot region at the bottom of the vessel, and, thus, heat loss and subsequent decrease in specific gravity of the product solution is reduced. The wave at the bottom of the vessel proceeds in the x-direction at a constant velocity of 4.02 mm min$^{-1}$. The rising plume is derived from the increase in temperature due to the autocatalytic reaction, causing the hot products to rise rapidly through the reagent solution as a volcanic plume. The plume rises through at 3.05 mm min$^{-1}$ in Fig. 6(b) and slows down to 1.54 mm min$^{-1}$ in Fig. 6(d) when it spreads horizontally back into finger A. From

FIG. 4. Schematic sketch of all the flows observed in this study. Wave initiation is from the top left hand corner of the vessel. Fingering, from the edge of the vessel, is rapid due to the efficient establishment of thermal equilibrium between products and the cold bulk reagents. Flow at the top aided by thermocapillary convection, and at the bottom, pure reaction-diffusion-convection propagation.

FIG. 5. Experimental data. The 4 consecutive series of pictures show the development of a finger from an initial Marangoni instability. This is repeated several times to give a series of convective tori in slightly shallower depths than one depicted in this figure. Temperature jump at the wave front for these series of experiments is much higher than in the experiments reported in this article. Adapted from Martincigh et al.1

FIG. 6. (a)–(d) The first observed finger and plume series. Development of finger A can be tracked from Figs. 6(a) to 6(d). It accelerates initially and then slows down, mainly due to the influence of the wave at the bottom of the vessel which reduces the thermal gradient, and hence thermogravitational effects. Plume also starts fast and slows down due to thermal equilibrium.
FIG. 7. (a)–(d) Another iteration of the finger development. As opposed to Fig. 5, the lower thermocapillary convection reduces the velocity and subsequently back-flow velocity. The wave then bifurcates at the bottom of the vessel to the right and the left.

The expected dynamics of the wave front, we expect the descending finger to be unstable (any defect at the wave front will be magnified) and the ascending plume to be stable (any deformities at the interface will be suppressed).

Figures 7(a)–7(d) show the further development of patterning from Fig. 6. Here, we concentrate on the development of a new finger. For a short period, wave moves with $v_x = 2.83 \text{ mm min}^{-1}$. Again, due to the lower $\Delta T$ at the wave front, the leading front of the wave quickly dips below the surface of the liquid and is propelled more by reaction-diffusion-convection mechanisms and not as much from the Marangoni instability. The formation of the fingering regime effectively halts the horizontal lateral instability of the effectively hydrothermal wave. Once a minor defect is created at the wave front interface, the denser products overcome the Marangoni convection and cause a rapid descent as the products force the wave front downward. This initial descent commences at 12.1 mm/min in the first one third of the reaction depth. Following this rapid descent, the wave front begins to curl back to the wave front at 1.16 mm/min due to availability of reagents and expected back flow, though not as pronounced as would have been expected from Fig. 4. After reaching the bottom of the container the wave front begins to rapidly propagate in both x-directions moving to the left at 4.70 mm/min and to the right at 2.93 mm/min. Propagation to the left is through a combination of RDC mechanisms and expected back flow (see Fig. 4), while propagation in the positive x-direction is solely through RDC mechanisms.

Figure 8 shows further progression of the hydrodynamical instabilities from Figs. 6 and 7. This is after three iterations of fingering and plume. The patterned changes in chemical wavelength, wave front speeds, and velocities of the plumes and fingers. There are two reactions proceeding in the reaction mixture: the catalyzed and noncatalyzed reactions. Thus, the chemical composition of reactants keeps changing with time. Figure 8 shows the usual initial lateral forward movement near the top of the solution, labeled as “A.” The finger “B” slants in the positive x-direction because of interaction with the plume movement just below it. Thus, the continuation of the plume in “C” has to proceed in the positive x-direction. Near the top of the solution, plume “C” bifurcates into “D” in the negative x-direction and “E” in the positive x-direction. The velocity of lateral instability “D” is much less than that of its positive counterpart “E.” D is slower because of the lower thermal gradient due to incoming instability A. Separation of A and D is maintained for extended periods. Instead of coalescing, they, instead, both finger into the unreacted region created by B and C resulting in very exotic heterogeneities in acid, sulfate, chlorine dioxide, and heat. After lateral instability denoted as “E,” the general sequence shown in Figs. 6 and 7 is repeated of finger and plume alternating. Our observation vessel is 260 mm long and thus not too many of these sequences are observed. Also, as wave approaches the end of the Hele-Shaw cell, boundary and eddy effects from the other wall begin to affect wave movement. In general, wave movement, propagation, and spatiotemporal organization are heavily dependent on vessel size and geometry. Figures 7 and 8 show entrainment of unreacted reagents, specifically the separation of fingers A and D in Fig. 8 and the isolated reagents in Fig. 6(d). If observed along the z-axis; the system will display spatiotemporal inhomogeneities in the x-axis. If the reaction vessel is wide enough, these spatiotemporal inhomogeneities will cover the x-y plane.

Overall system description

Chemical waves generated in an exothermic or endothermic reaction produce density inhomogeneities in the reaction medium. In considering double-diffusive flow between two parallel plates
that are perpendicular to the direction of gravity, the Oberbeck-Boussinesq approximation is used in which fluid density is assumed to depend linearly on two scalar fields, namely, the temperature,  \( T \), and the respective concentration of the different solution constituents, \( c_i \). In our nonlinear reaction system, the density will, thus, vary depending upon the extent of reaction and the temperature. The density, \( \rho \), can then be derived from

\[
\rho = \rho_0 \left( 1 - \alpha(T - T_0) + \sum \beta_i (c - c_i) \right),
\]

where \( \alpha = -(1/\tau) \delta \rho/\delta T; \) and \( \beta = (1/\rho) \delta \rho/\delta c_i \),

with subscripts "0" denoting initial conditions. \( \alpha \) is the thermal expansion coefficient and \( \beta \) is the solutal expansion coefficient. The first term in Eq. (6) represents the temperature dependence of density and the second term the concentration dependence. These two terms, in the system under study, deliver opposing effects, which give rise to the observed multicomponent convection (fingering regime). At isothermal conditions, the first term vanishes, leaving only the second term which, in this reaction system, \( \rho > \rho_0 \). This inequality will dictate the effect of buoyancy at thermal equilibrium. However, for both reactants and products, the buoyancy factors can be estimated by the Grashof number,

\[
Gr = -(\delta \rho/\delta T)g \Delta T d^3/\rho \kappa \nu = \alpha g \Delta T d^3/\kappa \nu,
\]

where \( d \) is the depth of the solution, \( \kappa \) is the thermal diffusivity, and \( \nu \) the kinematic viscosity. Since in our experiments we evaluated \( \delta \rho/\delta T = -0.4 \) kg m\(^{-1}\) K\(^{-1}\), then when \( T > T_0 \), the first term in Eq. (4) will pull the value of density, \( \rho \), to lighter values. With \( d \) raised to the third power, buoyancy factors are much more pronounced in deeper fluid solutions.

**MARANGONI CONVECTION**

At isothermal conditions, the product solution has a slightly lower surface tension than the reactant solution. Thus, both reaction-diffusion and thermocapillary convection act in the same direction with respect to wave propagation,

\[
\text{Surface tension, } \sigma = \sigma_0(y) = \tau (T - T_0) \quad \text{with } \tau = (\delta \sigma/\delta T)_p, c.
\]

Surface tension, in general, decreases with temperature, and our reaction solutions gave \( \tau = -9.5 \times 10^{-5} \) N m\(^{-1}\) K\(^{-1}\). Thermocapillary forces can be estimated from the Marangoni number, \( Ma \),

\[
Ma = -\tau \Delta TL/\rho \kappa \nu,
\]

where \( L \) is the length of the free surface. Although \( \Delta T \) changes over a reaction zone of about 2 mm (see Fig. 1), a reasonable Marangoni number can be calculated from our experimental data. The Eötvös number is effectively the ratio of thermogravitational and thermocapillary forces,

\[
\text{Eötvös number, } E_0 = Ra/Ma,
\]

where \( Ra \), the Rayleigh number \( = Gr \cdot \nu/\kappa \) is a better measure of the relative strengths of the forces responsible for the formation of the convective torus. In general, we calculate that the Eötvös number has a temperature-independent value of 0.40 for the same vessel and same conditions. \(^{34} \) This means that, for the same temperature gradient, the Marangoni effect should be 2.5 times larger than the thermogravitational-based Rayleigh effects. We expect, theoretically, a strong dominance of thermocapillary effects, with \( \Delta T \), over thermogravitational effects, and this was also observed experimentally in a previous publication from our laboratories. It is, thus, easy to investigate our pseudo-two-dimensional thermocapillary-buoyancy flows. Numerical analysis of the strongly-convective thermocapillary flows has shown that a finite temperature gradient remains over the entire free surface, leading to a global flow structure. Theoretical work by Smith and Davis also concluded that thermocapillary convection dominates in such cavity problems in the limit of fixed \( E_o \) and as Ma → infinity.

**Overall formulation of the problem**

Our system is a modified form of a Benard-Marangoni (BM) instability with internal heat generation. Convective instabilities driven by either buoyancy or thermocapillary effects have been the subject of numerous theoretical and experimental studies since the initial studies of Lord Rayleigh and Pearson. Considerably less work has been done on the more general and relevant case in which both mechanisms are acting simultaneously. Pioneering theoretical work on this BM instability was performed by Villiers and Platten in which they used acetone with lateral heating. \(^{35} \) They obtained qualitative agreement with experimental data with respect to the various convective regimes: steady or oscillatory. Velocity profiles were those calculated and shown in Fig. 9. Prandtl number for acetone was fixed at 4.24, and the major bifurcation parameter was the temperature jump imposed at the wave front. They modeled transient convection in a finite rectangular enclosure by using the stream function vorticity formulation of the two-dimensional Navier-Stokes and energy equations and assuming the Boussinesq approximation. By utilizing the relation

\[
\frac{\partial \sigma}{\partial x} = \frac{\partial \sigma}{\partial T} \frac{\partial T}{\partial x},
\]

and the Marangoni number [Eq. (10)] and the force for a non-deformable flat surface, they could utilize the no-slip adherence boundary condition and the Levich boundary condition

\[
\rho_0 \nu \frac{\partial V_z}{\partial z} = \frac{\partial \sigma}{\partial T} \frac{\partial T}{\partial x},
\]

to determine the commencement and nature of the lateral instability. Our reaction-diffusion-convection system requires several other parameters that come into play. Specifically, for a full modeling of the system, one needs evaluation and determination of the following parameters: Atwood number, Lewis number, Damköhler number, Schmidt number, Peclet number, the Nusselt number, and the Soret effect. \(^{36} \) Many phenomena and parameters like the Atwood numbers and Soret effect can be evaluated for our system. \(^{29, 30} \) We determined an Atwood number, \( A_t \), of \( 1.4 \times 10^{-3} \). In Rayleigh-Taylor instability (RTI), the penetration distance of heavy fluid bubbles into the light fluid is a function of acceleration time scale, \( Ag \), where \( g \) is the gravitational acceleration and \( t \) is the time. \(^{36} \) In our system, both natural and buoyant convections are important [see Eq. (4)]. There is no absolute cut-off value, but, generally, with \( A_t < 0.1 \), we can
FIG. 9. Images were converted to a HSV (hue/saturation/value) image, and HSV is used extensively in computer vision and image analysis for feature detection. The saturation channel of the HSV image was extracted and filtered. This is presented as a grayscale image, highlighting specifically the wave front. In subimage 1, the wave front is thickest at the top of the wave (A) at 4.30 mm, thinnest at (B) at 0.90 mm, and has an average thickness of 1.74 mm. Between subimages wave (A) moves laterally at an average rate of 7.32 mm min\(^{-1}\) and (B) moves at 2.71 mm min\(^{-1}\). In subimage 2, the thickest part of the wave front has moved to the bottom, and this can be attributed to the lower forward velocity and measures 2.57 mm. The central portion of the wave front shows no forward momentum and becomes the thinnest portion at 1.04 mm as thermocapillary convection draws more of the reacting front to the top of the wave, seen in arrow (C), and reaction-diffusion-convection causes an accumulation along (D).

Assume that the velocity field is solenoidal and thus use the Boussinesq approximation. At small Atwood numbers, the development of Rayleigh-Taylor instabilities proceeds initially through an exponential growth of infinitesimal perturbations that correspond to linear stability analysis. At amplitudes of about half of the wavelengths, the RTI saturates, and longer wavelengths take over. There will finally be nonlinear mode interactions for final successive wavelength saturation. We can define changes in temperature and concentration at the front as

\[
\Delta c = c_1 - c_0; \Delta T = T_1 - T_0 = -\Delta H_{\text{rea}} \cdot \Delta c/\rho_0 C_p. \tag{14}
\]

\(T_1\) is calculated from the extent of reaction and assumption that there are no heat losses. The Soret effect quantifies the fact that a flux can be generated solely by a temperature difference. This effect does not exist in pure fluids, and thus we can define a factor \(D_T\) for \(c_0(1 - c_0)\). \(D_T\) remains temperature dependent exactly as the original \(D\), before the temperature jump. Total flux is the standard Fick’s term, combined with a mass separation due to the temperature gradient. These two terms are of opposite signs, and when we are in steady state conditions, the terms will be equal with zero total flux, implying

\[
\frac{\partial c}{\partial x} = -\frac{D_T}{D} c_0(1 - c_0) \frac{\partial T}{\partial x}. \tag{15}
\]

We define the Soret coefficient as

\[
S_T = D_T/D. \tag{16}
\]

This effect can be positive or negative, depending on the sign of \(D_T\). We have attempted, initially, to estimate our Soret coefficient. The reaction commences with mainly organic compounds and has inorganic ionic products. We expect an absolute Soret coefficient for organics of approximately \(1.00 \times 10^{-3}\) to \(10^{-2}\) \(K^{-1}\). In our system, with a \(\Delta T\) of 1.5–2.5 K, we expect a negligible Soret effect since this will normally approximate to the extent of reaction within the linear region of temperature jump at reaction front. The Soret effect would be sizable, however, in fluid physics of crude oil reservoirs. We also expect thermal conductivity and kinetics constants to be essentially invariant after the temperature jump.

Approaches toward a reasonable model

We assumed no convection in the y-axis and accepted a 2D model as plausible for this system with the incompressible velocity field \(\mathbf{u} = (u_x, u_z)\). The system has length \(L_x\) and height/width \(L_z\). Gravity points along the z-axis. We then apply Darcy’s law [Eq. (17)] and continuity equation [Eq. (18)] and combine this with the reaction-diffusion-convection evolution equations for the autocatalytic production of the product species, \(c\), and for the elevation of the temperature, \(T\), derived from the exothermic reaction [Eqs. (19) and (20)]

\[
\nabla p = -\frac{\mu}{\kappa} \mathbf{u} + \rho(c, T)g, \tag{17}
\]

\[
\nabla \cdot \mathbf{u} = 0, \tag{18}
\]

\[
\frac{\partial c}{\partial t} + \mathbf{u} \cdot \nabla c = D\nabla^2 c + f(c), \tag{19}
\]
In Darcy’s law, density depends on concentrations and temperature as quantified in Eq. (6). “D” in Eq. (15) is the diffusion coefficient. In Eq. (20), $\alpha$ is the Newton heat loss coefficient. Since we worked only in the reaction front, this last term in Eq. (20) was ignored. We need to obtain dimensionless equations and parameters. We proceed to define the following:

$$
\frac{\rho \alpha c_p}{\theta} \left[ \frac{\partial T}{\partial t} + u \cdot \nabla T \right] = \kappa \nabla^2 T + |\Delta H_{\text{rxn}}(c) - \alpha(T - T_0)|. 
$$

In massive stars, there exists a semiconvection zone, a chemically inhomogeneous zone between the convective core and the radiative envelope. The governing equations mirror the equations we have derived here in this article. As the star evolves, a gradient of chemical composition develops at the outer border of the convective core. The atmosphere is convectively stable if the Ledoux criterion is met and unstable if the Schwarzschild criterion is met. More explicitly, in order for convection to occur, the adiabatic temperature gradient should be smaller than the actual temperature gradient of the surrounding gas (Schwarzschild), which is given by the radiative temperature gradient if convection does not occur (Ledoux). We have examined the development of plumes as analogous to the overshooting observed in Schwarzschild-type instabilities.

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**REFERENCES**

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