Hydrophilic-hydrophobic patterning is used by a variety of biosystems to direct the motion of fluids at surfaces. For example, hydrophobically hydrophilically patterned backs help desert beetles to capture water, and hydrophobic patches control water permeation in leaves [1]. This motif is also used to steer the motion and reaction of fluid droplets in liquid microchips [2] and is being utilized to design self-cleaning substrates [1]. Despite the utility of these designs, there have been surprisingly few theoretical studies into the dynamics of fluid flow over chemically patterned surfaces. In this study, we examine a conceptually simple system where two partially miscible fluids, A and B, are mechanically driven (by an imposed shear) to flow past patterned surfaces within a microchannel (see Fig. 1). The system exhibits two distinct steady states; however, in the transition between the two states, we uncover intricately complicated behavior, where monodisperse droplets of both A in B and B in A are formed periodically in time (as shown in Fig. 2), and the confined liquid displays regular, nondecaying oscillations in its structural characteristics. Furthermore, we isolate points where this system bifurcates between time-independent behavior and different types of oscillatory patterns. What is striking is that the observed phenomena occur even in the absence of hydrodynamic interaction; this is distinct from well-known instabilities in fluids [3,4]. Given that the system is relatively simple, the results suggest that complex transitions between well-defined steady states may well be evident in a broad variety of dynamical systems. In terms of technological applications, emulsions that contain monodisperse droplets are vital in the pharmaceutical, food and cosmetic industries [5]. Our results indicate that variations in surface potential can be exploited to create the desired solutions in a microchannel.

The observed complex oscillatory patterns arise from a competition between advection and thermodynamics as an imposed Poiseuille flow drives the phase-separated fluids to flow over the chemically patterned substrates. As shown in Fig. 1, the top and bottom of the microchannel are decorated with a checkerboard pattern. Each checkerboard is composed of two A(B)-like patches, which are preferentially wetted by the A(B) fluid. The first B patch (in yellow) is placed in the way of the A stream (in blue) and, correspondingly, the first A patch is located in the path of the B fluid.

The binary fluid is characterized by the order parameter \( \varphi(\mathbf{r}, t) = \rho_A(\mathbf{r}, t) - \rho_B(\mathbf{r}, t) \), where \( \rho_i(\mathbf{r}, t) \) represents the local number density of the \( i \)th component, \( i = A, B \). The thermodynamic behavior of the system is governed by the coarse-grained free energy functional, \( F = F_0 + \Psi_s \), where \( F_0 \) is the Ginzburg-Landau free energy for a binary mixture

\[
F_0 = \int d\mathbf{r} \left[ -\frac{a}{2} \varphi^2 + \frac{b}{4} \varphi^4 + \frac{k}{2} \nabla \varphi^2 \right],
\]

and \( a \) and \( b \) are positive constants. We consider the fluid to be in the two-phase coexistence regime where the equilibrium order parameter for the A(B) phase is \( \varphi_{eq} = \pm \sqrt{a/b} \). The term \( \frac{1}{2} |\nabla \varphi|^2 \) represents the cost of order parameter gradients. The free energy \( \Psi_s \) describes the interaction of a fluid element at a point \( \mathbf{r} \) with the patterned substrate. Specifically, we take [6]

![FIG. 1 (color). Schematic of the system.](image-url)
The imposed Poiseuille flow advects variations on the patterns and is proportional to the gradient of the chemical potential, \( \nabla \Delta \phi \). The velocity field \( \mathbf{v} \) obeys the Navier-Stokes equation in the overdamped limit, which is appropriate for low Reynolds number flow,

\[
0 = -\nabla p + \nabla^2 \mathbf{v} + \mathbf{H} + C \frac{\delta F}{\delta \phi} \mathbf{\nabla} \phi,
\]

where \( p \) is a Lagrange multiplier that guarantees the incompressibility condition, \( \nabla \cdot \mathbf{v} = 0 \), and \( H_x = (P_{\text{in}} - P_{\text{out}}) \xi_{\text{int}} / \eta L \) is the dimensionless form of the imposed pressure drop \( (P_{\text{in}} - P_{\text{out}}) \) along the channel of length \( L \); here, \( \eta \) is the shear viscosity. Because the pressure gradient is applied along the \( x \) axis, only the \( x \) component of the vector \( \mathbf{H} \) is nonzero, \( H_x = H \). The last term of Eq. (4) is the nondissipative part of the stress tensor \([9]\) (for the above dimensionless form, see [8]); this term represents hydrodynamic interactions. The constant \( C = \sigma \xi_{\text{int}} / (aM \eta) \) depends on the fluid properties, such as interfacial tension, \( \sigma = k \phi^{2}_c / \xi_{\text{int}} \), diffusivity \( aM \), and viscosity \( \eta \). The value of \( C \) determines the importance of hydrodynamic interactions for the specific fluid. For a fluid with a high viscosity, where \( C \ll 1 \), hydrodynamic interactions can often be neglected. In this work, we set \( C = 0 \); therefore, the velocity profile in our system is determined by the imposed pressure gradient. Thus, advection in a shear flow and diffusion of the fluids to the more wettable \( A \) or \( B \) domains control the evolution of the order parameter in the system and are responsible for the observed rich behavior [10].

Equations (1)–(3) are discretized and solved numerically by a cell dynamic system method [11] on a \( 120 \times 40 \times 40 \) grid. The following boundary conditions on the walls of the channel are imposed [12]: \( \frac{\partial \phi}{\partial n} \bigg|_{y=0} = 0 \), \( \frac{\partial \phi}{\partial n} \bigg|_{y=40} = 0 \), and \( \frac{\partial \phi}{\partial n} \bigg|_{x=0} = \int \partial \phi \bigg|_{x=0} \nabla \phi \bigg|_{x=0} \bigg|_{y=0} \nabla \phi \bigg|_{y=0} \bigg|_{z=0} \]. The last condition arises explicitly from the minimization of free energy in the presence of the substrate potential. At the entry of the channel, we have two-stream flow; at the exit, we assume free draining flow, i.e., \( \frac{\partial \phi}{\partial n} \bigg|_{x=0} = 0 \). For the velocity field, we assume no-slip boundary conditions on the walls [13].

We consider the order parameter evolution in the channel at different values of \( H \). At low \( H \) (low velocities), local thermodynamics dominate, and the fluid just mimics the underlying checkerboard pattern, with small distortions in \( \phi \) caused by the imposed flow. Higher velocities lead to more dramatic changes in \( \phi \) and yield the complex interfaces between the \( A/B \) fluids shown in Figs. 2 and 3. This behavior occurs when the scale of spatial distortions in \( \phi \) within the center of the channel (where the Poiseuille flow exhibits the maximum...
velocity, $v^{\text{max}}$) is comparable to the length of the patch, i.e., when $v^{\text{max}}l_{\text{diff}} = l$, where $l_{\text{diff}} = l^2$ is the characteristic diffusion time over the patch length $l$. This estimate yields a value of $H = 10^{-4}$. For higher values of $H$, the fluid flows through the middle of the channel with essentially no distortion while, near the top and the bottom substrates, $\varphi(r, t)$ is governed by the patches' wetting properties [see Fig. 3(e)].

For the intermediate $H$ values, a competition between the preferential wetting interactions and the imposed flow leads to fascinating behavior. On one hand, the wetting effects cause the fluids to diffuse to the respective patches to minimize the free energy and there is the general tendency to minimize interfacial regions between the $A$ and $B$ phases. On the other hand, the imposed flow carries fluid away from the favorable patches. If both substrates contained just the first half of the checkerboard (a single $A$ and $B$ patch), these patches would simply “switch” the location of the fluids, and the imposed flow would move the switched fluids along the channel. The presence of the second set of patches interrupts this flow because both the $A$ and $B$ streams again confront incompatible domains. In three dimensions, each component can avoid the second unfavorable region by diffusing into the bulk. However, recall that the first yellow ($B$) patch is on the blue ($A$) side (see Fig. 2). Thus, the $B$ fluid can extend only so far into the incompatible domain (similar arguments hold for the $A$ fluid). Each fluid forms “arms” that reach from the top and bottom of the walls; these arms can join and pinch off to form a bubble. Our coarse-grained modeling allows such a topology change without any ad hoc rules. Figures 2(a)–2(c) show the order parameter distribution at the “front” of the microchannel; the same behavior occurs for the $A$ fluid at the “back” of the channel (Figs. 2(d)–2(f)).

Between the limiting cases in Figs. 3(a) and 3(e) for relatively low and high $H$, respectively, three different types of behavior have been observed: two types of periodic behavior and a time-independent state asymmetric with respect to the top and bottom substrates [Fig. 3(b)]; which of the two morphologies shown in Fig. 3(b) is actually realized for fixed $H$ depends on the noise in the system [7]. The periodic cases exhibit “symmetric” [Fig. 3(c)] and “asymmetric” oscillations [Fig. 3(d)]. The arms in the figures and all the periodic behavior develop mainly near the sidewalls. In the middle of the box, the Poiseuille velocity field has a maximum and advection prevails. Near the wall, however, the velocity is much smaller and diffusion dominates, allowing the arms to move upward (downward) and join.

To analyze the complex dynamics, we define a parameter that characterizes the integrated changes in $\varphi(\vec{r}, t)$ near the sidewalls:

$$B_i(t) = \frac{1}{V_i} \int d\vec{r} |\varphi(\vec{r}, t) - \varphi(\vec{r}, 0)|.$$  \hspace{1cm} \text{(5)}

Here $i = \text{top}$, bottom indicates whether we integrate over the top (see red dashed box in Fig. 1) or bottom half of the sidewall region of volume $V_i$. (We choose the thickness of this region as $h/8$.) The evolution of $B_{\text{top}}(t)$ and $B_{\text{bot}}(t)$ for different values of $H$ is shown in Fig. 4. Note that $B_{\text{top}}(t) = B_{\text{bot}}(t)$ for all the symmetric cases [Figs. 3(a), 3(c), and 3(e)]. A case of asymmetric oscillations, where $B_{\text{top}}(t) \neq B_{\text{bot}}(t)$ is plotted in red. The maxima in the curves correspond to the largest distortions (where the bubbles are biggest); the minima correspond to the structure where the arms are separated by the greatest distance. The two curves for the asymmetric case [Fig. 3(d)] are similar to each other, but there is a phase shift between them. Each of these red curves displays two maxima and two minima in the periodic state. At early times, the system’s behavior is similar to the symmetric case but, at some time, spontaneous symmetry breaking occurs. The length of time before steady state is reached in the asymmetric case depends on degree of noise introduced in the strength of the fluid-substrate interaction [7]. The fact that one of the peaks becomes weaker than the other indicates that instead of the arms growing equally from both substrates and simultaneously forming a bubble in the middle, the top arm, for example, grows faster and contributes more to the bubble (higher maximum) than the bottom arm (smaller maximum). But for the next bubble, the situation is reversed, so the whole period encompasses both maxima; this period is roughly twice that of the symmetric case.

We also examined the system response as we change $H$ dynamically. For each value of $H$, the steady-state value of $B_{\text{bot}}$ or the maximum and minimum in the oscillatory regimes, are shown on the bifurcation diagram in Fig. 5. By abruptly increasing $H$ from below $H_1$ to above $H_6$, the system switches from the symmetric time-independent low-velocity regime [as in Fig. 3(a)] to the

![FIG. 4 (color). Evolution of $B_{\text{top}}$ and $B_{\text{bot}}$ for cases shown in Figs. 3(a) and 3(c)–3(e).](image-url)
Asymmetric oscillations, independent of the direction of the system is broken, we types of patterns [as in Fig. 3(b)]. Once the symmetry of symmetric with respect to top/bottom. Further gradual increases in $H$ we can observe all the different types of behavior shown in Figs. 3(b) and 3(d). Gradually decreasing $H$ leads to a transition from asymmetric to symmetric oscillations, and then to the time-independent asymmetric state (blue curve) [14]. Figure 5 clearly shows that the system displays hysteresis.

The nondecaying, time-periodic behavior in a simple binary fluid driven through the microchannel arises from interactions between the fluid and the patterned substrate, which introduces nonlinearity into the dynamical system. These interactions act as an “activator” in reaction-diffusion systems [3] and are responsible for the positive feedback. We note that the periods of the oscillations and the positions of the bifurcation points are dependent on the strength of the fluid-substrate interactions and the patch length. Thus, the system dynamics can potentially be controlled by varying these chemical features. In particular, other choices of patterns can potentially lead to new spatiotemporal patterns and dynamical behavior.

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**FIG. 5** (color). Bifurcation diagram. Open circles mark asymmetric oscillations, filled circles represent symmetric oscillations, and straight lines indicate time-independent behavior [except green lines that depict both symmetric ($H < H_1$) and asymmetric ($H > H_1$) oscillations]. All other parameters (except $H$) are the same as in Fig. 2. In the oscillatory regimes, the top and bottom curves correspond to the maximum and minimum of $B_r$.

**FIG. 3** (color). Oscillations (both types), time-independent oscillations (blue curve), and symmetric oscillations (green curve). The parameter regimes are marked by different symbols: filled circular markers indicate the critical values of $H$ for the symmetric oscillations, the top and bottom curves correspond to the maximum and minimum of $B_r$.

**FIG. 2** (color). Oscillations (both types), time-independent oscillations (blue curve), and symmetric oscillations (green curve). The parameter regimes are marked by different symbols: filled circular markers indicate the critical values of $H$ for the symmetric oscillations, the top and bottom curves correspond to the maximum and minimum of $B_r$.

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[4] We note that Thorsen et al. [Phys. Rev. Lett. 86, 4163 (2001)] experimentally demonstrated complex pattern formation involving water/oil/surfactant mixtures in microchannels that arises from a well-known competition between interfacial tension and shear forces. In the latter study, interactions between immiscible liquids introduce nonlinearity into the system while, in our case, nonlinearity is introduced and controlled by the patterned substrate.
[7] We introduce noise in the strength of the interaction at each point of the checkerboard pattern, i.e., $V(\delta) = V(1 + 0.03 \xi(\delta))$, where the independently distributed random numbers $\xi(\delta)$ lie in $[-1, 1]$.
[10] We have, however, carried out simulations with hydrodynamic interactions ($C = 10$). We observed similar periodic droplet formation, but the quantitative characteristics (the actual value of the imposed pressure gradient when droplet formation occurs, the period, and amplitude of oscillations) were altered by the specific value of $C$.
[12] Note that here and in the following text (including all the figure captions), all values are dimensionless. For simplicity, we use the same symbols for scaled values ($\mu$, $V$, $\varphi$, $h$, etc.) as we use for dimensional values in Eqs. (1) and (2). Here, $\mu$ is scaled by $(a \phi_0^2)$ with respect to its dimensional value (see, for example, [8]) and, correspondingly, $V$ is scaled by $a$.
[14] As a function of $H$, the period for symmetric oscillations is almost constant while, for asymmetric oscillations, it increases rapidly with increasing $H$. 

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