Swelling of Diffusive Fluid Threads in Microchannels

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The dynamics of viscous fluid threads concurrently flowing with miscible solvents is experimentally investigated in square microchannels. Diffusive fluid threads are found to significantly swell at low flow velocities due to large specific interfacial area and hydrodynamic lubrication. An approach based on bounded function analysis of confined thread diameter is developed to model diffusive behavior of viscosity-differing fluids at the small scale. This work shows the determination of a critical flow rate associated with each fluid pair and the use of dynamic similarity to calculate diffusion coefficients between oils and organic solvents. The thread divergence is estimated based on the growth of diffusion layers and related to diffusion-induced buckling instabilities of viscous threads in parallel flows.

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Fluids threads comprise a broad class of slender viscous structures that are commonly encountered in industrial processes, including fiber synthesis [1] and electrospray [2], and often occur in natural situations, such as during the coiling of a liquid rope [3–5], or during the capillary breakup of confined jets [6–8]. While gravity and inertia play an important role at the large scale, the behavior of two liquids flowing at the small scale is closely related to the magnitude of their viscosity coefficients $\eta_1$ and $\eta_2$, as well as interfacial tension $\gamma_{12}$ for immiscible fluids or diffusion coefficient $D_{12}$ for miscible fluid pairs. Diffusive fluid displacement in porouslike media have shown the emergence of distinct miscible liquid structures, such viscous fingers and spikes, depending on the Péclet number $\text{Pe}$ [9,10]. Microfluidic systems provide precise control over flow geometry and flow rates of injection, and the use of hydrodynamic focusing sections are practical to investigate mass transfer phenomena between fluids having similar viscosities $\eta_1 \sim \eta_2$ [11–13]. In the case of large viscosity contrasts $\eta_1 \gg \eta_2$, lubricated threads of fluid L1 can readily form in a low-viscosity phase $L_2$ using a simple channel intersection [14]. Miscible core-annular flows display complex dynamics [15,16], including diffusive and inertial instabilities [17], as well as viscous buckling instabilities [18–20], which are of particular interest for the generation of cramped and helical microfibers [21–23].

Lubricated microfluidic threads also have large specific interfacial area compared to two-fluid layered flows and this property offers advantages for the development of continuous hydrodynamic methods for enhancing micromixing of viscosity-differing fluids [24] as well as for examining mass transfer mechanisms with complex fluids [25]. Determination of diffusion coefficients $D_{12}$ between fully miscible fluids in the context of large viscosity contrasts $\chi = \eta_1/\eta_2$, however, requires the use of advanced experimental techniques [26–28]. Steady-state measurements and theoretical modeling of $D_{12}$ at large $\chi$ are also challenging due to the spatial evolution of fluid viscosity and concentration gradients in the diffusive region [29]. In addition, numerous fluids of industrial interest, such as alcohols and polymers, are only partially miscible and form nonideal, conjugate solutions depending on their solubility [30]. Therefore, generic microfluidic methods are needed for the rapid characterization of $D_{12}$ between fully and partially miscible fluids made of chemically different species with various molecular affinities, such as between nonpolar viscous oils and polar organic solvents.

Here, the thread-forming ability of high-viscosity fluids is used to examine the relationship between convective and diffusive transport between high-molecular weight solutes and low-molecular weight solvents in confined microsystems. A method based on novel phenomenological and mathematical understanding of diffusive fluid threads in microchannels is developed to determine diffusion coefficients between fully and partially miscible fluids. The growth of diffusion layers around the thread is also shown to induce viscous buckling instabilities in straight microchannels.

Microfluidic platforms are made of glass and silicon where square microchannels of height $h = 250 \mu m$ form an orthogonal focusing section. The thick fluid $L_1$ is injected at volume flow rate $Q_1$ into the central channel and the thin fluid $L_2$ is symmetrically introduced in the side channels at $Q_2$ using syringe pumps [Fig. 1(a)]. The device is placed on top of an inverted microscope and illuminated with a fiber light for high-speed imaging. Fluids are made of conventional silicone oils and low-molecular weight alcohols.

Stable threads correspond to steady core-annular flows with a central stream of width $\varepsilon_0$. In the absence of interfacial tension and neglecting diffusion, solution of Stokes equation [15] in a circular tube of diameter $h$
At low flow rates, miscible threads are seen to significantly swell and a minimum diameter $\varepsilon_0$ is found in excess of $\varepsilon_0$ as shown on Fig. 1(a). Since the stable diameter $\varepsilon_0$ depends on $\varphi$, experiments are conducted by fixing the relative flow rate $\varphi = Q_1/Q_2$ while varying the absolute flow rate $Q_T = Q_1 + Q_2$. Figure 1(b) shows the spatial evolution of the normalized central stream width $\varepsilon/h$ for $\varphi = 0.25$ when $L_1$ is made 100-cS silicone oil and $L_2$ consists of isopropanol [Fig. 1(c)]. As $\varepsilon/h$ grows at the junction, a minimal value $\varepsilon_M$ is measured in the outlet channel before widening further downstream. Data show that $\varepsilon_M$ tends to $\varepsilon_0$ at large $Q_T$ while $\varepsilon_M$ tends to $h$ at small flow rates. The quantity $\varepsilon_M$ is practical as it provides a single measurement to characterize the role of diffusion on thread morphology. Therefore, we examine the influence of $Q_1$ and $Q_T$ on the variation of $\varepsilon_M$ for fixed values of $\varphi$. On the one hand, plotting $\varepsilon_M$ as a function of $Q_1$ [Fig. 1(d)] ungroups iso-$\varphi$ curves, which appear distinct and bounded with varying slopes. On the other hand, displaying $\varepsilon_M$ as a function of $Q_T$ [Fig. 1(e)] regroups iso-$\varphi$ data onto a master curve at low $Q_T$ and shows the progressive separation of each curve from the trend as $\varepsilon_M$ tends to $\varepsilon_0$ at large $Q_T$. The asymptotic behavior of $\varepsilon_M$ for large $Q_T$ is evident when represented as a function of $\varphi$ in conjunction with Eq. (1) on Fig. 1(f).

As experimental data show that, for a given fluid pair at fixed $\varphi$, $\varepsilon_M$ depends on $Q_T$ and ranges between 1 and $\varepsilon_0$, we briefly discuss the use of bounded functions to model the evolution of $\varepsilon_M$. A simple expression of $S$-shaped function corresponds to $f(x) = 1/(a + x^b)$ with $a > 0$. In the case where $a = 1$, symmetry properties indicate that $f(x) = f^C(x^{-1})$, where the complementary function $f^C = 1-f$. Therefore, a bounded function can also be described using a complementary function of a reciprocal variable. In general, as $f = a^{-1}$ when $x^b \to 0$, the sign of variable $b$ indicates whether $f$ tends to $a^{-1}$ when $x \to 0$ for $b > 0$ or when $x \to \infty$ for $b < 0$. The magnitude of $b$ corresponds to the steepness of the curve between the bounded values of 0 and $a^{-1}$ for $f$ or between 1 and $1-a^{-1}$ for $f^C$. Therefore, given the boundary conditions of $\varepsilon_M/h$, a complementary bounded function of the form

$$\frac{\varepsilon_M^*}{h} = 1 - \frac{1}{1 - \varepsilon_0/h + \left(\frac{Q_T}{Q_C}\right)^{-1.5}}$$

is proposed to model thread behavior. The steepness value of $b = -1.5$ is found to best fit the overall shape of curves for all fluid pairs and is therefore postulated as constant. By contrast, the value $Q_C$ corresponds to the offset of $Q_T$ and is specific to each fluid pair. The evolution of $\varepsilon_M/h$ along iso-$\varphi$ curves is calculated using Eqs. (1) and (2), and is displayed as a thin line for each $\varphi$ curve on Fig. 1(e). In practice, determination of the fitting parameter $Q_C$ is made along iso-$\varphi$ curves in the ($\varepsilon_M/h$, $Q_1$) parameter space displayed on Fig. 1(d) using $Q_T = Q_1(1 + \varphi^{-1})$ as individual curves are distinguishable and do not overlap with one another. Finally, observed values of $\varepsilon_M/h$ and predicted values of $\varepsilon_M^*/h$ are found to closely match as shown in Fig. 1(g). In conclusion, the method of analyzing bounded functions along iso-$\varphi$ curves of diffusive threads permits the determination of a single kinematic quantity $Q_C$ associated with each fluid pair.
To better understand the relationship between \( Q_C \) and the intermolecular coefficient of diffusion \( D_{12} \), we apply the iso-\( \varphi \) technique to fully miscible silicone oil fluid pairs and we examine a few soluble silicone oil and alcohol fluid pairs as shown in Fig. 2(a). In each case, a single critical flow rate \( Q_C \) is found to closely fit data for each fluid pair. The diffusion coefficient \( D_{12} \) between a solute made of large molecules having hydrodynamic radius \( R_1 \) and a solvent of low-molecular weight having viscosity \( \eta_2 \) can be inferred using the Stokes-Einstein equation, 

\[
D_{12} = \frac{kT}{6\pi\eta_2 R_1},
\]

where \( k \) is the Boltzmann constant and \( T \) is the temperature [31]. Hydrodynamic modeling of \( D_{12} \) is useful for the case of fully miscible fluids having similar chemical affinity, such as silicone oils of various molecular weights, where the hydrodynamic radius is assumed to scale as \( R_1 \sim \eta_1^{1/3} \) based on the manufacturer’s data sheet (Gelest) [32]. In this situation, the product \( D \eta_1^{1/3} \) is expected to remain constant at room temperature and can be estimated from tabulated values [26] to determine \( D_{12} \) for a given oil pair.

Here, the solvent viscosity of oil pairs is kept constant at \( \eta_2 = 0.49 \) mPa s (600 cS) and \( \eta_1 \) is varied according to 4.6 (5 cS), 48 (50 cS), and 487 mPa s (500 cS). The measured values of \( Q_C \) are plotted as a function of \( \eta_1 \) in Fig. 2(b) and data are well fitted with \( Q_C \sim \eta_1^{1/3} \), which in turn suggests \( Q_C \sim D_{12} \). Since the Péclet number \( Pe = Q/(hD_{12}) \) quantifies the relative influence of convection and diffusion, we compare \( Q_C/h \) to the values of \( D_{12} \) estimated using the hydrodynamic approach and find

\[
\frac{Q_C}{h} = Pe_C D_{12},
\]

where the critical \( Pe_C = 850 \) for fully miscible fluids [Fig. 2(b), inset]. The same method of determining \( Q_C \) is applied to the partially miscible oil and alcohol cases, where \( \eta_1 = 97 \) mPa s (100-cS oil) is fixed and \( \eta_2 = 2.25 \) (isopropanol), 2.83 (1-butanol), 3.57 (2-butanol), and 3.62 mPa s (1-pentanol). Over the limited range of solvent viscosity, data suggest \( Q_C \sim \eta_2^{1.7} \), which is in significant deviation from Stokes-Einstein prediction, as one would expect \( Q_C \sim \eta_2^{-1} \) [Fig. 2(c)]. Departure from theory is not unexpected as silicone oils and low-molecular alcohols have different chemical affinity and the configuration of a polymer in solution, such as its hydrodynamic radius \( R_1 \), is known to strongly depend on the interaction with the solvent [30,33]. In particular, alcohols can be considered as “poor” solvents for silicone oils since, for instance, we observe that a 100-cS silicone oil can be partially dissolved in butanol and pentanol but not in ethanol nor hexanol. Dynamic similarity, however, can be used to estimate the effective \( D_{12} \) for partially miscible oil and alcohol fluid pairs using Eq. (3), which yields the following estimates: \( D \approx 3.5 \times 10^{-10} \) (isopropanol), \( 2.5 \times 10^{-10} \) (1-butanol), \( 1.6 \times 10^{-10} \) (2-butanol), and \( 1.5 \times 10^{-10} \) (1-pentanol) m²/s with an evaluated uncertainty in the range of \( \Delta D \approx 10^{-11} \) m²/s based on the calculation of the critical flow rate with \( \Delta Q_C \approx 0.2 \mu\text{L}/\text{min} \). Hence, microfluidic determination of \( Q_C \) provides useful information about the initial flow interaction between chemically different fluids, which is characterized using an effective diffusion coefficient by analogy with fully miscible fluids.

Another aspect of the swelling of diffusive threads resides in the development of a diffusion layer \( \delta \) downstream from the location of \( \varepsilon_M \). As can be seen in Fig. 1(b), the spatial evolution of \( \varepsilon \) appears quasi-linear beside its bounded nature. To quantify thread swelling, we measure the location \( L_F \) where \( \varepsilon \approx h \) and the datum \( L_M \) where \( \varepsilon \approx \varepsilon_M \) as a function of the reduced flow rate \( Q_T/Q_C \) as indicated in Fig. 3(a). Both quantities depend on \( Q_T \) according to \( L_F/h = 5.3(Q_T/Q_C) \) and \( L_M/h = 1.5(Q_T/Q_C)^{1/2} \) [Fig. 3(b)]. Scaling relationships for \( L_M \) and \( L_F \) are used to estimate the thread divergence \( \nabla \cdot e \), where \( e = ei \) and \( i \) is the vector unit in the \( x \) direction, in the straight channel due to diffusion according to \( \nabla \cdot e = de/dx = (h - \varepsilon_M)/(L_F - L_M) \). Since \( de/dx \) depends on \( \varepsilon_M \), which in turn depends on \( \varphi \), we use the method of iso-\( \varphi \) curves with a few selected cases in Fig. 3(c). For \( Q_T < Q_C \), \( de/dx \) remains nearly constant while the divergence becomes inversely proportional to \( Q_T \) for \( Q_T > Q_C \). This behavior is expected as the exponent

\[
\frac{Q_C}{h} = Pe_C D_{12},
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FIG. 3. (a) Examples of measurement of $L_F$ and $L_M$ from micrographs. Fluid pair: 100-cS oil and isopropanol. Flow rates in $\mu$L/min (i) $(Q_T, Q_2) = (1, 2)$, (ii) (1, 4), and (iii) (1, 6). (b) Evolution of $L_M$ and $L_F$ as a function of $Q_T/Q_C$. Solid lines: $L_M/h = 1.5(Q_T/Q_C)^{1/2}$ and $L_F/h = 5.3(Q_T/Q_C)$. (c) Evolution of $\nabla \cdot \varepsilon$ with $Q_T/Q_C$ for various $\phi$. Inset: Comparison of $\nabla \cdot \varepsilon$ and $\nabla \cdot \varepsilon^\prime$ calculated from Eq. (4). Solid line: $\nabla \cdot \varepsilon = \nabla \cdot \varepsilon^\prime$. (d) Micrographs showing internal buckling. Fluid pairs: from (i) to (ii) 100-cS oil and isopropanol and from (iv) to (vi) 0.65-cS and 500-cS oils. (i) $(Q_1, Q_2) = (0.5, 17)$, (ii) (0.5, 25), (iii) (0.5, 40), (iv) (1, 60), (v) (1, 70), and (vi) (1, 90).

associated to the evolution of $L_F$ becomes dominant for large $Q_T$, which leads to the approximation

$$\nabla \cdot \varepsilon^\prime = a \left(1 - \frac{\varepsilon_0}{h}\right) \left(\frac{Q_T}{Q_C}\right)^{-1},$$

where $a = 1/5.3$. As can be seen in Fig. 3(c), inset, good agreement is found between data and Eq. (4). The thread divergence can also be expressed using Eq. (3) as $\nabla \cdot \varepsilon^\prime = b(\varepsilon_0^2/h) Pe^{-1}$, where the thread complementary diameter $\varepsilon_0^C = h - \varepsilon_0$ and $b \approx 160$. Thus, our analysis suggests the diffusion layer scales as $\delta \sim x/Pe$. This result compares with Taylor dispersions where $\delta \sim (x/Pe)^{1/3}$ in the upstream region and $\delta \sim (x/Pe)^{1/2}$ in the downstream region [34,35] taking into consideration the confined nature of $\delta$ and the use of scaling laws with varying exponents. Such “ultradiffusive” behavior of lubricated threads is interpreted as resulting from their large interfacial area and location at the centerline of parabolic stokes flows as opposed to stratified flows where diffusion primarily occurs near the solid walls.

Finally, the growth of the diffusion layer has implications on the structural stability of small threads. Previous analysis highlights the complex radial composition of miscible fluid threads having a highly viscous core of diameter $\varepsilon_0$ unsheathed within a swelled layer of diameter $\delta_M$ forming near the fluid contactor. In turn, the swelled thread is enclosed within a diffusive layer $\delta$ that further develops downstream. Thus, the highly viscous core experiences an overall deceleration as a result of swelling behavior of its envelope, which hints at the development of a dynamic compressive force along the thread axis to conserve momentum based on rudimentary control volume analysis. Experimental evidence of diffusion-induced buckling of viscous threads is displayed on Fig. 3(d) where threads are seen to coil within their diffusive envelope for small $\varepsilon/h$ at large $Q_T/Q_C$. Complementary theoretical and numerical work taking into account the development of transient stresses in the diffusive region would help further clarify internal buckling of diffusive fluid threads in parallel microflows.

Overall, the viscous regime of lubricated threads provides a useful reference to characterize mass diffusion phenomena between viscosity-differing fluids. Here, a thread-based method is developed in conjunction with bounded function analysis of iso-$q$ curves to identify a single kinematic quantity $Q_C$ associated with each fluid pair. In turn, information about $Q_C$ is employed to calculate an effective diffusion coefficient $D_{12}$ using similitude arguments. This approach of determining $Q_C$ does not require any particular assumption about $D_{12}$ and can be applied to a variety of fully and partially miscible fluid pairs having large viscosity contrasts. Techniques based on microfluidic viscous threads are promising for probing the role of fluid properties on diffusion as well as for characterizing diffusion-limited reactions with high-viscosity fluids. Future work could also examine the flow behavior of partially miscible fluids systems in relation with spontaneous emulsification phenomena to advance both practical understanding and modeling of viscous fluid interactions in confined microsystems.

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