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Chaotic mixing in effective compressible flows

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We study numerically joint mixing of salt and colloids by chaotic advection and how salt inhomogeneities accelerate or delay colloid mixing by inducing a velocity drift \( V_{dp} \) between colloids and fluid particles as proposed in recent experiments [J. Deseigne et al., Soft Matter 10, 4795 (2014)]. We demonstrate that because the drift velocity is no longer divergence free, small variations to the total velocity field drastically affect the evolution of colloid variance \( \sigma^2 = \langle C^2 \rangle - \langle C \rangle^2 \). A consequence is that mixing strongly depends on the mutual coherence between colloid and salt concentration fields, the short time evolution of scalar variance being governed by a new variance production term \( P = -\langle C^2 \nabla \cdot V_{dp} \rangle / 2 \) when scalar gradients are not developed yet so that dissipation is weak. Depending on initial conditions, mixing is then delayed or enhanced, and it is possible to find examples for which the two regimes (fast mixing followed by slow mixing) are observed consecutively when the variance source term reverses its sign. This is indeed the case for localized patches modeled as Gaussian concentration profiles.

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I. INTRODUCTION

Mixing of a scalar field \( C \) by laminar flows with chaotic trajectories in a bounded or periodic domain is often characterized by the evolution of its variance \( \langle C^2 \rangle \), a quantity known to decrease with time when the flow is incompressible and no source of scalar is present [1]. The case of mixing in compressible flows has received less attention: Vergassola and Avellaneda showed that a compressible flow can modify scalar transport either in stationary cellular flows or random delta correlated flows [2]; in the context of reactive flows, the dimension of a diffusion coefficient whose magnitude and sign depend on the precise nature of the salt and colloids [7,8]. Starting from a situation with initial concentration profiles \( C_0(\mathbf{r}) \) and \( S_0(\mathbf{r}) \), the time evolution of the concentrations is then given by the coupled advection diffusion equations:

\[
\frac{\partial S}{\partial t} + \nabla \cdot \mathbf{V} = D_S \nabla^2 S, \quad (1)
\]

\[
\frac{\partial C}{\partial t} + \nabla \cdot (\mathbf{V} + \mathbf{V}_{dp}) = D_C \nabla^2 C, \quad (2)
\]

where \( D_S \) and \( D_C \) are, respectively, the diffusion coefficients of the salt and colloids. In this situation, salt is mixed by the velocity field \( \mathbf{V} \) independently of the colloids, while the colloid concentration is also coupled to the salt concentration through the drift velocity \( \mathbf{V}_{dp} \). In order to study the impact of the salt gradients on colloid mixing, we restrict ourselves to two-dimensional (2D) situations; we achieve chaotic mixing using a time periodic velocity field, the so-called sine flow that has been a standard tool for studies of chaos in spatially smooth flows [1,11,12]. We recall it is \( T \) periodic with two subcycles for which the velocity field is \( \mathbf{V}(\mathbf{r}, t) = (\sin(\pi t), 0) \) for \( nT \leq t < (n + 1/2)T \) and \( \mathbf{V}(\mathbf{r}, t) = (0, \sin(\pi t)) \) for \( (n + 1/2)T \leq t < (n + 1)T \), which ensures the velocity field is divergence free at any time. For given initial concentration profiles \( C_0(\mathbf{r}) \) and \( S_0(\mathbf{r}) \), Eqs. (1) and (2) are solved with periodic boundary conditions using a pseudospectral method with a resolution of \( 512^2 \) for a square box of length \( L = 2\pi \).

We use an Adams-Bashford temporal scheme with order 2 and \( dt = 0.002 \) to ensure the scalar gradient and Laplacian are...
well resolved for diffusivities in the range $D_c \in [10^{-4}, 10^{-2}]$, which corresponds to Péclet numbers in the range $Pe = \max |V|L/D_c \in [0.065, 6.5]10^4$. For all simulations shown here we set $T = 0.8L/\max |V|$; in this regime of nonglobal chaos (see Poincaré sections in Figs. 3–10 of Ref. [13]), as in many natural situations, one can distinguish between thin structures developing due to stretching and folding and poor mixing in regular regions which govern the long time decay of scalar variance [1]. Figure 1(a) displays the colloid concentration field obtained in the reference case $V_{dp} = 0$, with Péclet number $Pe = 65 000$ and an initial profile $C_0(\mathbf{r}) = 1 + \sin(x)$, after five cycles of mixing.

When diffusiophoresis comes into play, the drift term $V_{dp} = \alpha V \log S$ depends on the instantaneous salt gradients. In order to be consistent with experiments [8,10], we set $\alpha = 0.001$ and $D_t = 0.01$ for all simulations, which corresponds to a Péclet number $Pe \sim 650$ for the salt. We define the initial salt concentration profile $S_0 = 1 + \sin(x) + S$, with $S = 0.01$ being an offset accounting for the unavoidable homogeneous background concentration of ionic species (buffer solution). As observed in experiments [10], one expects the drift term to modify colloid mixing depending on whether the initial colloid concentration field $C_0(x)$ is correlated, or anticorrelated, with the initial salt concentration profile $S_0(x)$. For all Péclet numbers we study the evolution of colloid concentration variance $\langle (C - \langle C \rangle)^2 \rangle$ for two different initial profiles $C_0 = 1 \pm \sin(x)$ and compare the results to the reference case $\alpha = 0$. The drift velocity, which does not depend on $C(\mathbf{r}, t)$, is displayed in Fig. 1(b) together with the corresponding salt concentration field computed at $t/T = 5$. With the chosen parameters, the modification of the total velocity field acting on the colloids, $V + V_{dp}$, is never larger than 1%, as shown in Fig. 1(c). However, we will demonstrate that this small variation leads to a strong alteration of mixing because $V_{dp}$ is not divergence free [Fig. 1(b)].

III. MIXING TIME

In Fig. 2, we display the evolution of the mixing time $T_{mix}$ needed to decrease the initial colloid concentration variance by a factor of 2, as a function of the Péclet number $Pe = \max |V|L/D_c$. As already observed in experiments [10], when salt and the colloid are initially injected together [salt-in configuration, $C_0(\mathbf{r}) = 1 + \sin x \sim S_0(\mathbf{r})$], mixing is approximately 15% slower than in the reference case $\alpha = 0$. On the contrary, when salt and colloids are injected separately [salt-out configuration, $C_0(\mathbf{r}) = 1 + \sin x$ and $S_0(\mathbf{r}) = 0.01 + \sin x$], the mixing is nearly 20% faster. This result supports the idea that the action of diffusiophoresis can be seen as a modification of colloid transport properties through an effective diffusivity $D_{eff}$ corresponding to an effective Péclet number $Pe = \max |V|L/D_{eff}$. Using the curves $T_{mix}(Pe)$, well fitted by a power law for the reference case (as usual for nonglobal chaos), one would then find $Pe_{eff} \sim 1.5Pe$ and $Pe_{eff} \sim 0.5Pe$ for the salt-in and salt-out configurations, respectively. This means that a correction to the velocity field smaller than 1% leads to a 50% change in the colloid effective diffusivity. This result proved to be quite robust when performing the same simulations with $T = 1.6L/\max |V|$ (not shown here). In this

FIG. 1. (Color online) (a) Colloid concentration field at $t = 5T$ in the reference case ($D_c = 10^{-4}$, $Pe = 65 000$) for an initial concentration $C_0 = 1 + \sin(x)$. (b) Salt concentration field at $t = 5T$ for $S_0 = 1.01 + \sin(x)$ ($D_t = 10^{-2}$, $Pe_t = 650$) together with the corresponding drift velocity field $V_{dp} = \alpha V \log S$ with $\alpha = 0.001$. (c) Corresponding time evolution of nondimensional maximum drift velocity, $max |V_{dp}|/max |V|$.

FIG. 2. (Color online) Mixing time $T_{mix}/T$ as a function of the Péclet number $Pe = \max |V|L/D_c$ as measured from time evolution of the variance $\sigma^2 = \langle C^2 \rangle - \langle C \rangle^2$ with and without diffusiophoresis. For all cases $S_0 = 1.01 + \sin(x)$ with $D_t = 10^{-2}$ and $D_c = \in [10^{-4}, 10^{-2}]$. Circles: reference case with no diffusiophoresis $C_0 = 1 + \sin(x)$ and $\alpha = 0$. Dashed line: power law fit with exponent $\gamma = 0.26$. Squares: salt-in case $C_0 = 1 + \sin(x)$, $\alpha = 0.001$. Diamonds: salt-out case $C_0 = 1 - \sin(x)$, $\alpha = 0.001$. 013027-2

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regime of global chaos we did not observe any significant difference concerning diffusiophoresis: if mixing is indeed enhanced with \( T_{\text{mix}} \) varying linearly with \( \log(\text{Pe}) \), we found for all configurations (salt-in, salt-out) that \( T \) has a weak impact on the measured effective Péclet numbers.

**IV. COMPRESSIBLE EFFECTS**

However, this mean field approach based on \( T_{\text{mix}} \), if interesting for interpreting the long time behavior, may not catch the short time evolution of colloid variance when concentration gradients are not generated yet, so that diffusion is negligible. As displayed in Fig. 3, if one always observes a decrease of scalar variance at small times in the salt-out and reference cases, the salt-in case is more appealing: the variance first increases for \( t/T \leq 2 \) before decreasing for \( t/T \geq 2 \). This short-time increase is incompatible with an effective diffusivity which would predict a variance decreasing with rate \( d(C^2)/dt = -2D_{\text{eff}} \langle (\nabla C)^2 \rangle \), where \( \langle \cdot \rangle \) stands for spatial averaging over one period of the flow. For a better understanding of the colloid variance evolution, one may then come back to Eq. (2) and derive an equation for the scalar energy \( C^2 \) valid for compressible flows. One obtains

\[
\frac{1}{2} \frac{dC^2}{dt} = \nabla \cdot \left[ (\nabla + \nabla_{\text{dp}}) \frac{C^2}{2} - D_c \nabla C \right]
\]

\[
= -D_c (\nabla C)^2 - \frac{C^2}{2} \nabla \cdot \nabla_{\text{dp}}.
\]

This equation, although similar to the classical scalar energy budget, contains an additional term proportional to the drift velocity field compressibility \( \nabla \cdot \nabla_{\text{dp}} \) that does not vanish in the present case. When averaging over one flow period, all terms written as a divergence disappear, and one obtains a new global scalar energy budget valid for compressible flows:

\[
\frac{1}{2} \frac{d(C^2)}{dt} = -D_c (\langle \nabla C \rangle)^2 - \left( \frac{C^2}{2} \nabla \cdot \nabla_{\text{dp}} \right).
\]

Because mean scalar concentration remains conserved even in the case of compressible flows, this equation also gives the evolution of scalar variance \( \sigma^2 = \langle C^2 \rangle - \langle C \rangle^2 \). Therefore scalar variance in compressible flows results in the competition between scalar dissipation \( \varepsilon_c = -D_c (\langle \nabla C \rangle)^2 \) and production \( P = -\langle C^2 \nabla \cdot \nabla_{\text{dp}} \rangle/2 \), the latter being proportional to the mutual coherence between \( C^2 \) and the total flow field compressibility \( \nabla \cdot \nabla_{\text{dp}} \). This equation, well verified in our case at any time step, helps us understand the evolution of scalar variance in the presence of diffusiophoresis. As seen in Fig. 4(a), dissipation alone cannot explain the observed differences in scalar variance at short times; indeed, for large-scale initial concentration profiles, \( \varepsilon_c \) remains weak until small-scale fluctuations have been created by stretching and folding. For the two first mixing cycles \( t/T \leq 2 \), the evolution of the variance is then governed by the production term \( P = -\alpha (\langle C^2 \rangle \log S)/2 \), positive for the salt-in configuration and negative for the salt-out configuration, as demonstrated in Fig. 4(b). This effect disappears when the salt has been mixed. Considering the Péclet number of the salt (\( \text{Pe}_s = 650 \)), Fig. 2 shows the mixing time is of the order of two mixing cycles, which means salt is totally mixed at \( t = 2T_{\text{mix}} \sim 4T \). For \( t > 4T \) both the drift velocity field and the production term have become very weak [Figs. 1(c) and 4], so that the decay of variance is mainly governed by scalar dissipation thereafter. We may then conclude that diffusiophoresis is globally impacting mixing because compressibility modifies the colloid concentration field at small times while regular mixing of this modified initial condition follows, shifted in time.

Up to now we have dealt with modal (\( \sin x \)) and therefore “nonlocalized” initial conditions for \( C_0 \) and \( S_0 \). However, since mixing in the presence of diffusiophoresis is governed by the coherence between salt and colloid concentration, it strongly depends on initial conditions. Therefore it is interesting to look at more realistic situations where the initial condition is a patch of fluid. For instance, one can compare situations of a patch of salt plus colloids introduced in pure water (salt-in situation) to
For short times for the salt-out case before becoming positive for the salt-in case as shown in Fig. 5(b), it is negative for initial configurations [Fig. 5(c)]. Indeed, if larger than dissipation at small times for these still large-scale scales. This again can be understood by looking at the evolution of two time scales for the salt-out configuration. One remarkable property of this compressible effects starts to unmix. To better understand the salt-out configuration we simulated a pure demixing configuration with an homogeneous initial colloid concentration $C_0 = 1$, for which initial variance, production, and dissipation vanish. These results, presented in log scale in Fig. 6(a), show that if the long time behavior of the salt-in and reference cases are identical, the variance decrease of the salt-out case is much slower, similar to the one of pure demixing. In this latter case, variance production increases at short time because the concentration increases in regions where $\nabla \cdot \mathbf{V}_{dp}$ is negative and decreases in regions where $\nabla \cdot \mathbf{V}_{dp} > 0$. As a consequence the concentration pattern created has a strong correlation with $\nabla \cdot \mathbf{V}_{dp}$, and the production term reaches a very high value after two cycles of mixing [Fig. 6(b)], leading to a very slow variance decrease, as already observed for the salt-out case.

**V. CONCLUSION**

We have studied 2D chaotic mixing of colloids under the action of diffusiophoresis, which produces a velocity drift and makes colloids no longer be tracers of the flow motion. We have demonstrated that this very small drift, which leads to a correction to the velocity field smaller than 1%, is responsible for an important change in mixing because it strongly modifies the topology of the total flow, which is no longer divergence free. This compressibility is at the origin of the scalar variance production term $P = -(C^2 \nabla \cdot \mathbf{V}_{dp})/2$, necessary to understand (i) the evolution of scalar variance at small times, not explained in the framework of effective diffusion for the salt-in case, and (ii) acceleration or delay of mixing depending on the coherence between salt and colloid concentration fields. One remarkable property of this compressible effects is the capacity to unmix an initially homogeneous colloidal solution by simply adding salt gradients. If the present results were obtained on the physical case of colloid mixing in the presence of chemical gradients, they are more general and may play an important role in cell dynamics because the

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**FIG. 5.** (Color online) (a) Time evolution of colloid concentration variance for a patch of salt $S_0 = 0.01 + \exp[-4(x^2 + y^2)]$ with $D_p = 10^{-2}$, $D_c = 10^{-4}$. Circles: reference case with no diffusiophoresis $C_0 = \exp[-4(x^2 + y^2)]$, $\alpha = 0$. Squares: salt-in case $C_0 = \exp[-4(x^2 + y^2)]$ and $\alpha = 0.001$. Diamonds: salt-out case $C_0 = 1 - \exp[-4(x^2 + y^2)]$ and $\alpha = 0.001$. (b) Time evolution of production term $P = -(C^2 \nabla \cdot \mathbf{V}_{dp})/2$. (c) Corresponding time evolution of dissipation term $|\varepsilon_c| = D_c (|\nabla C|^2)$.

**FIG. 6.** (Color online) (a) Circles, squares, and diamonds are the same as in Fig. 5(a), plotted in semilogarithmic coordinates. Triangles: corresponding demixing configuration with an homogeneous initial colloid concentration $C_0 = 1$. (b) Time evolution of production term $P = -(C^2 \nabla \cdot \mathbf{V}_{dp})/2$ for the same salt-out configuration as in Fig. 5(b) (diamonds) and the demixing configuration (triangles).
total flow is compressible [9]. Effective flow compressibility should also apply to temperature gradients (Soret effect) and may help us understand older experiments of DNA trapping and amplification in laminar thermally driven flows [6], the Laplacian of the temperature field being an image of the local DNA concentration. Finally, we note these compressibility effects are not a manifestation of only laminar mixing. They are also observed in particle laden turbulent flows providing the particles do not follow the fluid motions because they have inertia [14,15] or because they are sensitive to gravitational field [16]. Bridging between the different results, this suggests the possibility of a common frame of description for turbulent clustering and diffusiophoretic mechanisms via compressible effects.

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