Hydrodynamic pairing of soft particles in a confined flow

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Abstract

The mechanism of hydrodynamics-induced pairing of soft particles, namely closed bilayer membranes (vesicles, a model system for red blood cells) and drops, is studied numerically with a special attention paid to the role of the confinement (the particles are within two rigid walls). This study unveils the complexity of the pairing mechanism due to hydrodynamic interactions. We find both for vesicles and for drops that two particles attract each other and form a stable pair at weak confinement if their initial separation is below a certain value. If the initial separation is beyond that distance, the particles repel each other and adopt a longer stable interdistance. This means that for the same confinement we have (at least) two stable branches. To which branch a pair of particles relaxes with time depends only on the initial configuration. An unstable branch is found between these two stable branches. At a critical confinement the stable branch corresponding to the shortest interdistance merges with the unstable branch in the form of a saddle-node bifurcation. At this critical confinement we have a finite jump from a solution corresponding to the continuation of the unbounded case to a solution which is induced by the presence of walls. The results are summarized in a phase diagram, which proves to be of a complex nature. The fact that both vesicles and drops have the same qualitative phase diagram points to the existence of a universal behavior, highlighting the fact that with regard to pairing the details of mechanical properties of the deformable particles are unimportant. This offers an interesting perspective for simple analytical modeling.

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I. INTRODUCTION:

In the microcirculation, it is often observed that the red blood cells (RBCs) flow in single or multiple files forming small trains of cells, called clusters[1, 2]. The arrangement and organization of the RBCs depend on the diameter of the vessel and their concentration (hematocrit). Each RBC interacts hydrodynamically with the other cells[3]. RBCs can also interact via another mechanism, namely an interaction mediated by plasma proteins. The latter interaction is materialized either by bridging between RBCs or by a depletion force. In the bridging mechanisms proteins make a real bridge between two neighboring RBCs, while in the depletion mechanism osmosis is responsible for the cluster formation. We have recently discussed the implication of plasma proteins in the formation of RBC clusters in microcirculation [4]. The main objective is to gain further insight into the role of each mechanism. Therefore this paper is be directed towards numerical study of the effect of hydrodynamic interactions on cluster formation.

Several studies have been devoted to understanding the hydrodynamical interaction between suspended particles in the Stokes regime. Analytical models [5, 6] considered the motion of a linear array of rigid spheres at low Reynolds number in a cylindrical tube under a pressure-driven flow (i.e., an imposed Poiseuille flow). Wang and Skalak[5] estimated the range of the hydrodynamic interactions between the spherical particles to be of the order of the tube diameter. Leichtberg et al.[6] showed that the interparticle interactions were relatively small at weak confinements, reached a maximum at intermediate confinement, and were quickly damped out at strong confinement. More recently, colloidal particles confined between two parallel plates in a quasi 2D geometry have been studied experimentally and theoretically [7–9]. The complexity in these systems arises from the difficulty to decouple the effect of Brownian diffusion from hydrodynamic interactions. An antidrag between the moving particles attributed to a negative hydrodynamic coupling has been reported. A change of sign of the hydrodynamic coupling (from attraction to repulsion) in a cylindrical channel was also observed [7]. The effect of boundaries on the hydrodynamic interactions has been studied in the case of water-in-oil drops in quasi-1D microfluidic devices having a square section of the order of the size of the drop, so that the drops are constrained to move along the channel axis [10–12]. This study reported on a non-monotonous behavior of the hydrodynamic interaction resulting from an interplay between the plug flow and the
screening of the long-range hydrodynamic interaction induced by the confinement. Janssen et al. [13] have studied numerically pairs of rigid spheres and deformable drops driven by a Poiseuille flow through a three-dimensional (3D) rectangular channel in the Stokes regime. Due to the reversibility of Stokes equations, the interdistance between a pair of rigid spheres does not evolve in time. However, for a pair of deformable drops, due to the up-stream/down-stream shape asymmetry, hydrodynamic interaction leads to an attraction at long interdistances and a repulsion at short interdistances. The long-range attraction was attributed to the source-quadrupole flows induced by drop in the Hele-Shaw geometry. The pair of drops tends to the same stationary interdistance independently of the capillary number, a measure of the flow strength, which only affects the time needed to reach the steady state.

The rheological behavior and spatial organization of a suspension of vesicles and capsules in 2D and 3D under shear and parabolic flows have been widely studied numerically by several groups [14–34]. Still, a complete determination of the phase diagram of the paired states has not yet been achieved. For example, is the branch of the stationary solutions (say the stationary interdistance of a pair as a function of the confinement) unique, or are there many branches? If many branches exist, is there a coexistence domain, and how the topology of the bifurcation diagram evolves with parameters. The present study is focused along this question and reveals a phase diagram of a complex nature.

It has been shown in a recent study that stable clusters of vesicles can form in the absence of bounding walls under an imposed parabolic flow profile [35]. Our objective in this paper is to take this unbounded case as a reference and see how confinement affects the hydrodynamic interactions. We generically find repulsion at short interdistance and attraction at long interdistance. We also find that at a long enough vesicle interdistance the interaction can change sign, become repulsive, and then become attractive at longer interdistance. This points to the existence of an interaction that changes sign with distance, highlighting the nontrivial effect of hydrodynamic interactions. We further analyze the complex structure of the branches of coexisting stationary solutions. We mainly focus this study on vesicles, as a closer biomimetic counterpart for RBCs, but we also investigate the hydrodynamic interactions between drops in order to investigate whether or not there is a generic pattern behind our findings and assess how sensitive the hydrodynamic interactions are to the mechanical properties of the particles. We show that the bifurcation diagram for
drops is similar to that for vesicles (both qualitatively and almost quantitatively), pointing to an underlying universality.

A systematic 2D numerical study (based on a boundary integral formulation) is undertaken here in order to analyze the time evolution of a pair of vesicles or a pair of drops in a pressure-driven flow, by exploring several parameters, such as channel width, initial separation of the particles, and the flow strength. This paper is organized as follows. In Section II we introduce the model in detail. Section III presents the main results for vesicles, both for weak and strong confinement. Section IV gives an overview of the results for drops. Section V is devoted to a discussion of the results and the implication for real situations.

II. THE MODEL AND THE SOLUTION METHOD

We first present the method adopted in this study to solve the motion of two hydrodynamically interacting vesicles or drops confined between two parallel plates and subject to a Poiseuille flow. We start by introducing the theoretical model and then discuss the numerical scheme and the precision.

A. Mechanical model for vesicles and drops

Vesicles, drops, capsules (drops coated with polymers) are endowed with different mechanical properties leading to different responses to external stresses. Both vesicles[36–39] and capsules[40–42] are widely used to mimic RBCs under flow. RBC’s complex dynamics, such as tank-treading, tumbling and vacillating breathing (aka swinging or trembling) can be reproduced by both vesicles and capsules. Shapes exhibited by RBCs under a Poiseuille flow, such as parachute and slipper shapes, are also captured by the vesicle and capsule models (see review [18]). Vesicles resist bending, and possess an inextensible membrane (constant area in 3D and perimeter in 2D) but do not present a resistance to shearing, in contrast to RBCs and capsules, which are endowed with surface shear elasticity. Nevertheless, several features exhibited by RBCs are also captured by vesicles, such as the above-mentioned dynamics and shapes under flow.

The membrane energy of vesicles [43] is expressed in its 2D form as

\[ E_v = \frac{\kappa}{2} \oint c^2 ds + \oint \zeta ds, \] (1)
where $\kappa$ is the membrane bending modulus, $c$ is the curvature, $ds$ is the arc length element, and $\zeta$ is a Lagrange multiplier, which enforces membrane inextensibility.

Liquid drops resist deformation by surface tension (line tension in 2D) forces, which tend to restore their shape to a sphere (a circle in 2D). The tension energy is written as

$$E_t = \zeta_0 \oint ds,$$

where $\zeta_0$ is the surface tension. We can see, that eq. (1) is reduced to (2) by setting $\kappa = 0$ and by treating $\zeta = \zeta_0$ as a constant representing the surface tension, which is an intrinsic quantity for drops (while for vesicles, $\zeta$ is not intrinsic to the membrane, but is an auxiliary field to enforce a constant local arclength and thus can change according to membrane load).

The 2D membrane force is obtained by calculating the functional derivative of the energy (1)

$$f = \begin{cases} \kappa \left( \frac{\partial^2 \zeta}{\partial s^2} + \frac{c^3}{2} \right) n - c \zeta n + \frac{\partial \zeta}{\partial s} t & \text{for vesicles} \\ -c \zeta_0 n & \text{for drops} \end{cases},$$

where $n$ and $t$ are the outward normal and the tangent unit vectors, respectively. The details of the derivation can be found in Ref. [44].

Both for vesicles and for drops, the particle size $R_v$, which serves as the length scale of the problem, is defined by the expression

$$S = \pi R_v^2,$$

where $S$ is the area inside the particle contour. Here and below, we use generic term particle to denote both drops and vesicles when there is no need to make a distinction between them. Unlike drops, vesicles (and RBCs) are characterized by their reduced area

$$\nu = \frac{S}{\pi \left[p/(2\pi)\right]^2},$$

an intrinsic dimensionless parameter that expresses the ratio between the actual fluid area enclosed by the vesicle contour and the area of a disk having the same perimeter $p$ as the vesicle. For RBCs, the reduced volume (3D equivalent of the reduced area) lies in the range of 0.60 – 0.65. Accordingly, the reduced area $\nu$ is kept fixed to 0.65 in all our simulations. The equilibrium shape obtained by minimizing (1) for vesicles with reduced volume $\nu = 0.60 – 0.65$ is a biconcave shape similar to that exhibited by a RBC at rest [45].
B. Boundary integral formulation

We consider a system of two identical hydrodynamically interacting particles driven by a Poiseuille flow in a confined geometry between two laterally infinite plates (see Fig.1). The fluids inside and outside the particles have the same densities $\rho_{in} = \rho_{out} = \rho$, and the same dynamic viscosities $\mu_{in} = \mu_{out} = \mu$. The velocity in the absence of particles (i.e. the undisturbed velocity) is denoted by $u^\infty$ and its Cartesian components are given by

$$
\begin{align*}
  u^\infty_1(x) &= u_{max} \left[ 1 - \left( \frac{x_2}{W/2} \right)^2 \right] \\
  u^\infty_2(x) &= 0
\end{align*}
$$

where $x(x_1, x_2)$ is an arbitrary point in the whole domain, $u_{max}$ is the midplane velocity and $W$ is the channel width. $x_1$ and $x_2$ are the cartesian coordinates of $x$ along the flow and in the perpendicular directions, respectively. The Reynolds number associated with a vesicle (with typical size $10\mu m$, water viscosity and speed in the range of mm/s, as found in microcirculation) is of order $0.01$. The parameters for drops are chosen so that the Reynolds number remains small as compared to unity. Thus the motion of the fluids can safely be described by the Stokes equations

$$
\begin{align*}
  \nabla \cdot \sigma(x) &= -\nabla P(x) + \mu \nabla^2 u(x) = 0 \quad \text{(7)} \\
  \nabla \cdot u(x) &= 0 \quad \text{(8)}
\end{align*}
$$

where $\sigma$ is the stress tensor associated to the total velocity and $P$ is the pressure. The jump in the interfacial traction is balanced by the particle interface force $f = -[\sigma_{out} - \sigma_{in}]n$. 

![FIG. 1. Illustration of two vesicles driven by a Poiseuille flow and confined between two parallel walls situated at $x_2 = \pm W/2$.](image)
Besides the latter force balance condition, the solution of equations (7) and (8) must respect the following boundary conditions (B.C.)

\[
\begin{align*}
    \mathbf{u}(x) &= 0, \quad \text{when } x \text{ lies on the walls} \\
    \lim_{x \to \infty} [\mathbf{u}(x) - \mathbf{u}^\infty(x)] &= 0 \\
    \mathbf{u}_{\text{out}}(x) &= \mathbf{u}_{\text{in}}(x), \quad x \in \partial \Omega_i \quad (i = 1, \ldots, N_v)
\end{align*}
\]

where the subscripts “in” and “out” refer to the fluid inside and outside the particle, \( \partial \Omega_i \) is the contour of the \( i \)th particle and \( N_v \) is the total number of particles. The Stokes equations can be converted into a boundary integral equation [46] defined over \( \partial \Omega_i \), which reads

\[
\mathbf{u}(x) = \mathbf{u}^\infty(x) + \frac{1}{4\pi \mu} \sum_i \oint_{\partial \Omega_i} \mathbf{G}^{2W}(x, y) \cdot \mathbf{f}(y) ds(y)
\]

where \( \mathbf{G}^{2W} \) is a Green’s function satisfying the appropriate boundary conditions (i.e. it vanishes at the walls and at infinity), \( x(x_1, x_2) \) and \( y(y_1, y_2) \) are the target and source points, and \( ds \) is the element of membrane arclength. The whole right hand side in (12) is the total flow field. The integral on the right hand side is performed over the particles’ contours \( \partial \Omega_i \) and corresponds to the contribution of the particles to the total velocity field. This contribution is referred to as the induced flow field. We also focus on the flow field in the co-moving frame, that is the frame moving with the pair of vesicles (when they reach a steady-state interdistance).

C. Numerical method

1. Dimensionless form

Most of the governing equations are the same for vesicles and drops. The difference appears only in the interfacial force expression and in the characteristic time of shape relaxation to equilibrium. Dimensionless quantities are denoted with a star symbol and are defined as

\[
c^* = c R_v, \quad s^* = s/R_v, \quad \mathbf{u}^* = \mathbf{u} \tau_c/R_v, \quad x^* = x/R_v
\]
both for vesicles and for drops. The shape relaxation time $\tau_c$ is taken as a time unit and is defined as

$$\tau_c = \begin{cases} \frac{\mu R^3}{\kappa} & \text{for vesicles} \\ \frac{\mu R_e}{\zeta_0} & \text{for drops}. \end{cases}$$ \hspace{1cm} (14)$$

For vesicles, the Lagrange multiplier is non-dimensionalized as

$$\zeta^* = \frac{\zeta R^2}{\kappa}. \hspace{1cm} (15)$$

The dimensionless integral equation of the velocity along the contour is given by

$$C_a u^*(x^*) = C_a u^\infty(x^*) + \frac{1}{4\pi} \sum_i \int_{\partial\Omega_i} G^2 W(x^*, y^*) \cdot f^*(y^*) ds^*(y^*) \hspace{1cm} (16)$$

The dimensionless expression $f^*$ of the membrane force reads

$$f^* = \begin{cases} \left(\frac{\partial^2 c^*}{\partial s^*^2} + \frac{c^*^3}{2}\right) n - c^* \zeta^* n + \frac{\partial c^*}{\partial s^*} t & \text{for vesicles} \\ -c^* n & \text{for drops}. \end{cases} \hspace{1cm} (17)$$

In addition to the reduced area defined above for vesicles $(\nu = (S/\pi)/(p/2\pi)^2)$, we have a new dimensionless number associated to the flow in equation (16), which is given by

$$C_a = \begin{cases} \frac{\mu R^3 \dot{\gamma}}{\kappa} = \frac{\mu R^4}{\kappa} \frac{u_{max}}{(W/2)^2} & \text{for vesicles} \\ \frac{\mu R_e \dot{\gamma}}{\zeta_0} = \frac{\mu R^2}{\zeta_0} \frac{u_{max}}{(W/2)^2} & \text{for drops}. \end{cases} \hspace{1cm} (18)$$

We refer to this number as the capillary number. Here $\dot{\gamma} = 4R_v u_{max}/W^2$ is the shear rate defined as the value of $\partial u_1^{\infty}/\partial x_2$ at $x_2 = R_v/2$. We recall that the shear rate in a parabolic flow is position-dependent, unlike in a linear shear flow. Thus, the definition of the capillary number for a parabolic flow is not unique in the literature. For example, in [47] the chosen shear rate is that at the wall and is equal to $\dot{\gamma}_W = 4u_{max}/W$. The ratio between the present shear rate and that in [47] is equal to $R_v/W$. In a previous study [39] as a validation of the present code, we have reproduced the full phase diagram of the vesicle shapes obtained in Ref.[47] in the plane of capillary number and the degree of confinement. Here, we shall explore the ranges of parameters where the vesicle shape is of parachute type only (slipper shapes are excluded from our study). Similarly, the drops are maintained in the center of the channel in order to make an adequate comparison with the vesicle case.

The capillary number is the ratio between the flow stress and bending force density. It may be viewed also as the ratio between the characteristic shape relaxation time $\tau_c$ (eq.
(14) and the time scale of the flow $\tau_f = 1/\dot{\gamma}$. In order to have a reference for the conversion of dimensionless units into physical ones, the following dimensional numbers for RBCs can be used: $R_v = 3\mu m$, $\mu = 10^{-3}$ Pa·s and $\kappa = 10^{-19} J$. This leads to a characteristic time of shape relaxation of about $\tau_c \sim 0.2 - 0.3$ s. This is quite consistent with measured values for RBC[48–50].

In this study, we quantify the hydrodynamic interactions of two particles by tracking the time evolution of the distance between their centers of mass. Let us denote the leading particle by 1 and the following particle by 2, and let $X^{(i)}(t) = R_v X^{* (i)}(t)$ denote the instantaneous position of the center of mass of the contour of particle $i$ at time $t$. Then the distance between the particles (called interdistance below) is defined as

$$\Delta X(t) = R_v \Delta X(t)^* = X^{(1)}_1(t) - X^{(2)}_1(t).$$

(19)

Consistently, we define the velocity with which the particles approach (or separate from) each other as

$$\Delta U(t) = \frac{R_v}{\tau_c} \Delta U(t)^* = \frac{d\Delta X(t)}{dt}.$$  

(20)

2. Numerical method for the vesicle dynamics

The integral equation (16) is discretized using the trapezoid rule and derivatives are approximated using a finite difference scheme. Each particle is described by a collection of equispaced Lagrangian nodes advected by the flow. Their motion is obtained by solving the advection equation for each material node $x$ lying on the membrane

$$\frac{dx}{dt} = u(x)$$

(21)

We calculate the velocity of each discretized point on the membranes by solving the integral equation (16), and the position of each node is updated at each time step using an explicit Euler scheme.

$$x(t + dt) = x(t) + u(x(t), t)dt$$

(22)

A tension-like parameter is introduced as a penalty parameter instead of the Lagrange multiplier $\zeta$ (which enters both tangential and normal force; see equation (3)). In other words, each material point is linked to each of its two neighbors via a very stiff “spring”
in order to enforce the local conservation of arclength, as described in [51]. If the spring stiffness is denoted as $T_{\text{tens}}$, we can define a time scale $\tau_{\text{tens}} = \mu/(T_{\text{tens}}R_v)$. This numerical time scale is to be compared to the physical time scale $\tau_c$ defined in equation (14) and to the flow time scale $\tau_f = \dot{\gamma}^{-1}$. $\tau_{\text{tens}}$ must be taken small enough in comparison to $\tau_c$ and $\tau_f$ so that on the physical and flow time scales the local incompressibility of the membrane is safely satisfied. For most practical purposes $\tau_{\text{tens}} = 10^{-4} - 10^{-3} \tau_c$ has proven to be largely sufficient (see below).

There is no arclength conservation for drops. Therefore, the discretization points tend to accumulate in some regions of the membrane while depleting in the others if the simple advection of material points (21) is used. We resolve this challenge by applying an additional displacement of the discretization points every time step. The displacement field is chosen to be (i) tangential to the interface of the drop so that its shape remains the same and (ii) such that the discretization points approach the equispaced distribution of the interface.

Each membrane is described by $N_{\text{mem}} = 120$ nodes whose positions are updated each $\Delta t = 10^{-4} \tau_c$ (i.e. this corresponds to the time step). The relative errors corresponding to the area, the perimeter and the reduced area are around 0.07%, 0.035%, and 0.0009%, respectively. The steady-state value of the distance between the mass centers of the two vesicles $\Delta X_f^* \equiv \Delta X^*(\infty)$ is also reported (see Table.I). The calculations are performed on a cluster consisting of 32 dual-core AMD64 processors with 24GB RAM per node. OpenMP directives are used to parallelize the matrix-vector product computation. The time needed to complete $10^6$ iterations as a function of the number of cores is reported in Fig.2(a) using two vesicles in a channel of width $W/R_v = 2$ and a $C_a = 10$. Similarly, we have plotted the required run time to complete $2 \cdot 10^5$ steps using 12 cores as a function of the number of vesicles (see Fig.2(b)). It is important to underline that some of the cases reported in the phase diagram in the result section ran over more than two weeks on a 12-core node since we decided to avoid using any cut-off or periodic boundary conditions in our system due to the long-range nature of the hydrodynamic interaction. The use of the appropriate Green’s function (that vanishes on the walls) allowed us to avoid finite size effects, since we can consider literally an infinite domain along the flow direction.
TABLE I. Relative errors on area, perimeter and reduced area of vesicles in Poisuiille flow $N_{mem} = \{120; 180; 240\}$ and $\Delta t = 10^{-4} \tau_c$. Also shown is the final distance between the mass centers of the vesicles $\Delta X_f^*$. The other relevant parameters are $C_a = 10$ and $W = 2R_v$. We obtained almost the same results for $\Delta t = 5.10^{-5} \tau_c$.

| $N_{mem}$ | $\frac{(s-S_0)}{S_0}$ (%) | $\frac{(p-p_0)}{p_0}$ (%) | $\frac{(\nu-\nu_0)}{\nu_0}$ (%) | $\Delta X_f^*$ |
|-----------|-------------------------|-------------------------|-------------------------|----------------|
| 120       | 0.07                    | 0.035                   | 0.0009                  | 5.83           |
| 180       | 0.03                    | 0.015                   | 0.0004                  | 5.83           |
| 240       | 0.017                   | 0.008                   | 0.0003                  | 5.83           |

3. Numerical method for the velocity field inside and outside the particles

The velocity field in the fluid domains (inside and outside the particles) obeys the same boundary integral equation as that on the membrane (Eq. (12)), where now $\mathbf{x}$ is a location of any point in the $(x_1, x_2)$ plane. Once a steady-state configuration is reached (the final shape as well as the vesicle interdistance), the velocity field is evaluated as a post-processing task. We introduce a regular square grid (with a certain degree of refinement; the mesh size can be taken significantly smaller than $ds$ if need be). Since the Green’s function is singular when the target point coincides with the source point, a small stripe (of order $ds$ in width) around the membranes is excluded from the fluid domain in order to ensure a good behavior of the velocity field. The lattice points are in general not on the membrane, and we only need to evaluate the distance between the source point (lying on the membrane) and the target point (lying on the square grid). For each point $\mathbf{x}$, the velocity field is evaluated by using Eq. (12), where the integral along the membrane is performed exactly in the same way as in the previous section.

III. RESULTS FOR VESICLES

In this section, simulations of a pair of vesicles in channels of different widths are performed. Unless indicated otherwise, vesicles are initialized as elongated ellipses with major axis parallel to the flow direction. We quantify the hydrodynamic interaction between the vesicles by tracking the interdistance $\Delta X_f^*$. We investigate the role of the confinement, the capillary number, and the initial configuration (shapes and interdistances of vesicles) on
FIG. 2. The Total run time (in seconds) required to compute $10^6$ iterations as a function of the number of cores (a). The total run time needed to reach $2 \cdot 10^5$ steps as a function of the number of vesicles using 12 cores (b).

We shall see that in some cases, there are several coexisting stationary interdistances for a given value of confinement and capillary number.

**A. Weak confinement**

A previous study has been devoted to the cluster formation in the absence of walls [35]. We first study the behavior of a pair of vesicles in weak confinement in order to check that we can capture almost the same result. We have analyzed the time evolution of a pair of vesicles in a channel having a width $W = 20R_v$, which corresponds to weak confinement. We find that the steady-state interdistance is equal to about $2.53R_v$ for $W = 20R_v$. This result compares well with that obtained in an unbounded flow [35] where the steady-state interdistance is of about $2.4R_v$.

We have analyzed systematically the behavior of a pair of vesicles for different (but still weak) confinements and different initial conditions. Figure 3 shows a typical behavior of the pair interdistance as a function of time for different initial separations, which are denoted as $\Delta X_{init}^* \equiv \Delta X^*(0)$ in that figure. We see there that different initial conditions lead to the same final state. Figure 4 shows the final configuration of the pair of vesicles as well as the induced flow field, that is the total flow field from which we subtract the imposed Poiseuille flow. The total flow field is also shown.
FIG. 3. The pair interdistance as a function of time for different initial conditions (initial interdistance). $W = 20R_v$, and $C_a = 10$. Note that the horizontal axis is represented in log scale.

In order to further analyze the time evolution of vesicle pairs, we have studied systematically the behavior of the relative velocity of the two centers of mass of the pair of vesicles $\Delta U^*$ as a function of their interdistance $\Delta X^*$ by considering different initial interdistances. A positive value of $\Delta U^*$ means that the two vesicles repel each other, while a negative value means they attract each other. A stationary interdistance corresponds to $\Delta U^* = 0$. Figure 5 shows $\Delta U^*$ as a function of the dimensionless interdistance between centers of mass, $\Delta X^*$. We observe a repulsion at short interdistance and an attraction at long interdistance. There exists one stationary interdistance characterized by $\Delta U^* = 0$ (denoted as 1 within a dark circle in Figure 5). This stationary solution is unique and independent of the choice of the initial shapes explored so far (Figure 6). Furthermore, this stationary solution is unambiguously stable since for longer interdistances the velocity is negative meaning the vesicles attract each other, while for shorter interdistances, the relative velocity is positive and the vesicles repel. Generally, if the relative velocity crosses zero by going from positive to negative values (as $\Delta X^*$ increases), we have a stable stationary solution.

We performed this study for several other weak confinements and determined the corresponding stationary interdistances. Figure 7 shows the branch of stationary interdistance $\Delta X^*$ as a function of $W/R_v$. The stationary interdistance weakly depends on confinement,
FIG. 4. The induced flow field for a pair of vesicles in a weakly confined flow (a); $W = 20R_v$

We also represent the flow in the frame moving with the vesicle called hereafter “co-moving frame" (b). A zoom in the co-moving frame in the region located between the cells reveals the absence of bolus in case of $W = 20R_v$ where the final interdistance is about $2.4R_v$ (c); and the presence of quasi-circular bolus for $W = 13R_v$, where the final interdistance is of about $3.4R_v$ (d).

and remains close to about $3R_v$.

**B. Strong confinement**

Let us now examine the generic behavior under strong confinement. Consider the case
FIG. 5. The relative velocity as a function of the interdistance; $W = 18R_v$, and $C_a = 10$.

FIG. 6. The relative velocity as a function of the interdistance; $W = 18R_v$, and $C_a = 10$. Starting from different initial shapes, the pair of vesicles converges to the same stationary solution. The labels (A), (B), (C), and (D) correspond to the initial shapes depicted in the right figure.

$W = 3R_v$. Looking at the evolution of the interdistance as a function of time, we find that the vesicle pair settles into a bound, stable steady state for several initial conditions. Two
FIG. 7. The branch of solution representing the stationary interdistance as a function of the channel width.

examples are illustrated in Fig. 8. The first noticeable feature is a significantly larger stationary interdistance for $W = 3R_v$ than for weak confinement: the stationary interdistance of about $5R_v$ for $W = 3R_v$ is about two times the value found for weak confinement (about $2.5R_v$). A first interpretation would suggest the screening of the hydrodynamical interaction by the confining walls. However, since the screening is felt both in attraction and repulsion, this reasoning is \textit{a priori} not justified, leaving unclear the mechanism by which confinement would shift both attraction and repulsion zones.

To approach a more rigorous interpretation, let us first analyze the velocity field in a strong enough confinement regime. Figure 9 shows the induced flow field for two confinements $W = 6R_v$ and $W = 3R_v$, as well as the flow field in the co-moving frame (the frame moving with the pair). The main difference is that the flow lines which extend far away in the case of weak confinement (see Figure 4) are cut off by the effect of the walls in the case of strong confinement. As we shall see below, this partially hints to a weaker interaction magnitude. As for the weak-confinement case, we also analyze the relative velocity as a function of the interdistance. The result is shown in Figure 10. We obtain a stable stationary solution, which results for a short-range repulsion and long-range attraction. We find that the relative velocity amplitude is significantly smaller than that obtained for a weak confinement (compare with Figure 5). This can be traced back to the screening effect of the walls which weaken the interaction. We perform this study for several confinements
FIG. 8. The interdistance of a pair of parachute-like vesicles as a function of time for different initial interdistances $\Delta X_{\text{init}}^* = 3.9$ (solid line) and $\Delta X_{\text{init}}^* = 6$ (symbols). (b): The effect of the initial shape of vesicles in the pair on the steady-state solution is shown for ellipses (symbols) and parachutes (solid line). $W = 3R_v$, $C_a = 10$. Note that the horizontal axis is represented in log scale.

FIG. 9. The flow field in a co-moving frame (a,c) and the induced flow (b,d) field for a pair of vesicles for $W = 6R_v$ and $W = 3R_v$.

to determine the corresponding steady-state solution. Figure 11 shows the branch of stationary interdistance $\overline{\Delta X}$ as a function of $W/R_v$. In contrast to the weakly confined case (also shown in Figure 11), the stationary interdistance strongly depends on confinement, and varies from about $4R_v$ up to about $32R_v$. We have a new solution branch for small values of $W$ (the leftmost branch in Figure 11), distinct from the one discussed above for large $W$ (the rightmost branch in Figure 11). We have two distinct branches, one presenting a
FIG. 10. The relative velocity as a function of the interdistance; $W = 3R_v$, and $C_a = 10$.  

FIG. 11. The branch of solution representing the stationary interdistance as a function of the channel width for a strong confinement.

stationary interdistance that increases with $W$ and the other whose stationary interdistance decreases with $W$. We shall now dig further into the structure of the phase diagram.

C. Full phase diagram

We broaden our investigation in order to clarify the overall structure of the topology of the phase diagram. First, we analyze the basin of attraction of each branch by exploring a
wider range of initial conditions. We begin our discussion with the weak-confinement case. In the previous section, we saw that different initial conditions led to the same final solution. Exploring wider and wider regions of initial conditions reveals a different scenario. Keeping the same confinement $W = 13R_v$, as above, we find that beyond a certain initial pair inter-distance, the solution no longer converges to the same value. Figure 12(left) shows the time evolution of the interdistance, which is still evolving after 25000$\tau_c$ but converges ultimately to a value of about $23R_v$. For different initial conditions (shorter initial interdistances), we have seen a final interdistance of about $3.4R_v$ (Figure 12(right)). This clearly demonstrates the coexistence of two different stable solutions. In general, two stable solutions should be separated by an unstable solution. We seek to determine the location of the unstable branch by analyzing the relative velocity as a function of interdistance, as before.

Our results are shown in Figure 13. The locations where the relative velocity of the vesicles vanishes are indicated by points marked 1, 2, and 3. Points 1 and 3 correspond to a stable interdistance, whereas point 2 corresponds to an unstable one. Since the relative velocity in the vicinity of point 3 is very small, we zoom in to reveal the structure of the dependence of the relative velocity on the interdistance. Another systematic analysis done by varying the confinement (i.e. $W$) allows us to show the full diagram of stationary solutions (like points 1, 2, and 3) as a function of $W$. The results are summarized in Figure 14.

We see that the branch for weak confinement undergoes a fold singularity in the form of a saddle-node bifurcation, in which a stable solution (represented by solid line) merges with an
FIG. 13. Normalized relative velocity as a function of the interdistance for $W/R_v = 13$. (a), (b) and (c) show successive zooms. Solid dots and open circles represent, respectively, stable and unstable fixed points. A succession of stable and unstable fixed points is observed.

unstable solution (dashed line). The branch arising at strong confinement continues to exist (as a stable solution) beyond the saddle-node point, and does not show, for the values of $W$ explored so far, any sign of disappearance. Theoretically, this branch may either continue to exist for any $W$, or it may undergo a fold singularity, one of the scenarios expected from catastrophe theory. Since for weak confinement the relative velocity becomes too small to be of practical interest, the question of the behavior of the branches at long interdistances is only academic. We shall thus not dwell here any farther on this issue.

Finally, let us say few words on the strength of interaction as a function of confinement. As can be expected, confinement reduces the strength of the interaction by screening. To understand the difference in the pairing mechanism at large and small channel widths we first consider a pair of vesicles in two different channels of widths $6R_v$ and $15.25R_v$. The
FIG. 14. The full set of branches of stationary interdistance as a function of the channel width. Solid lines represent stable branches, whereas dashed lines unstable ones.

time evolution of the interdistance in each channel is shown in Figure 15. We looked at characteristic time needed to reach the steady state for different channel widths finding that, for the channel of width \( W = 6R_v \), the interdistance reduces by approximatively \( 1.4R_v \) in a time \( 7000\tau_c \), whereas in a channel of width \( W = 15.25R_v \), the interdistance reduces by \( 7.2R_v \) in only \( 120\tau_c \). Considering that the typical time \( \tau_c \) for RBC is about 0.1 s gives a time of about 10 min in the first case and 10 s in the second one. We must note that in reality there are always fluctuations and imperfections keeping the cells from remaining in the same lateral position. For example, if the leading cell is slightly off-centered its velocity may be sensibly different from that of the following cell, so that the characteristic time needed to reach the bound state may be on a significantly shorter time scale than 10 s.

D. Effect of the capillary number

In this section we describe the effect of the capillary number on the main results. We consider a pair of cells with four different capillary numbers \( C_a = \{5, 10, 25, 100\} \) and flowing in channels of widths ranging from \( 2R_v \) to \( 20R_v \) (\( \approx 6 \) to \( 60 \mu m \) for RBCs). Figure 16
FIG. 15. Time evolution of the interdistance between two vesicles in different channel widths. (a) and (b) \( W = 15.25R_v \). (c) and (d) \( W = 6R_v \). In (b) and (d), different initial shapes ellipses (symbols) and parachutes (solid line) are used. Note that the horizontal axis is represented in log scale.

depicts the stationary interdistance as a function of the channel width for different capillary numbers (we do not show the full branch as before due to computational cost, and especially because we do not see any significant changes). Globally, the capillary number seems to not significantly affect the behavior of the pair. For vesicles, higher values of \( C_a \) can be reasonably reached, such as 2000. For a vesicle of radius 20 \( \mu m \), a channel radius of 200 \( \mu m \), and a velocity of 1 \( cm/s \), one finds approximately \( C_a \simeq 1600 \). By assuming that the relative velocity scales with \( C_a \) (as shown in Figure 17), one finds that the relative velocity is of about \( 100R_v/\tau \sim 1R_v/s \), which is not devoid of experimental testability.
FIG. 16. Stationary interdistance as a function of the channel width for different capillary numbers (a). The phase diagram for $C_a = 10$ and $C_a = 100$ (b). Role of the initial shape (solid lines are ellipses and dashed lines are parachutes) on the time evolution of the pair interdistance for $C_a = 5$ and $W/R_v = 12, 13$ and 14 corresponding to the transition area (c). Same as (c) but for $W/R_v = 12$ and $C_a = 10, 25, \text{ and } 100$ (d). Note that the horizontal axis is represented in log scale in panels (c) and (d).

IV. RESULTS FOR DROPS

Since most of the behaviors observed for vesicles are also observed for drops, we give only a brief discussion of the phase diagram for drop pairs. We have checked, as for vesicles, that initial shapes do not affect the final state. Drops can break up into several smaller drops when subject to a strong enough shear flow. The same effect occurs in Poiseuille flow. For this reason, it was important to choose the capillary number for which the drops remain stable. We chose $C_a = 0.3$. Figure 18 shows the steady state of a pair of drops and the corresponding flow lines. Unlike the vesicle case, drops exhibit recirculation zones inside in...
FIG. 17. Relative velocity as a function of interdistance in weak confinement $W/R_v = 18$ for different capillary numbers. The dashed (grey), solid (blue), and dash-dotted (green) lines are for $C_a = 10, 100, \text{ and } 200$, respectively.

the form of two counter-rotating vortices. In addition, the velocity field at the interface is not constant along the contour as the drop interface is compressible.

The question naturally arises of whether or not the apparent difference in the overall flow patterns as compared to vesicles results in different phase diagrams. We have explored in a very systematic way the existence of branches of stationary solutions. The results are summarized in the phase diagram in Fig.19. This phase diagram is strikingly similar to that obtained for vesicles (Fig. 14). Not only is there a qualitative similarity but also the stationary interdistances obtained for drops in each confinement are close to those found for vesicles. The emergence of these strong similarities between the two systems points to the existence of a universal feature, where the details of the physical system do not matter too much. This lack of sensitivity of the hydrodynamic interaction to the mechanical properties of the interacting particles provides a prospect for further theoretical analysis.
FIG. 18. The velocity field in the co-moving frame of a pair of drops (c and d) and vesicles (a and b) in a weakly confined flow ($W = 20$ and $13 R_v$). $C_a = 10$ for the vesicles and $C_a = 0.3$ for the drops.

V. CONCLUSION

This study reports on a complex phase diagram regarding hydrodynamic interaction between two vesicles or two drops in a confined pressure-driven flow. It is found that several branches of stationary solutions coexist. An interesting fact is that vesicles and drops behave in the same way, despite the different nature of the underlying physics between
FIG. 19. Stationary interdistance for two drops as a function of the channel width. $C_\alpha = 0.3$.

the two systems. This offers a useful basis for analytical modeling of the main features of the pairing process. As we have pointed out, for some branches of the phase diagram, the relative velocity of the pair of particles may become very small, in particular, due to the exponential decrease of the hydrodynamic interaction with interdistance. Thus other branches for longer interdistances may exist but be quite difficult to resolve numerically due to the smallness of interactions. While the existence of higher branches is an interesting fundamental question in itself, the very small amplitude of hydrodynamic interactions at long interdistances reduces the significance of this question for practical applications.

It would be interesting in the future to extend this study to the 3D case both for vesicles and RBCs. Another extension of this work is the analysis of many cells and more precisely how the stable size of a cluster evolves as a function of confinement. It has been reported in [35] that in an unconfined Poiseuille flow, the cluster size depends on the flow strength: increasing the flow strength allows cluster of larger sizes to remain stable. It would be interesting to draw general conclusion about cluster stability in the presence of walls in order to complement the already existing literature on this topic [32]. A study of the stability of clusters would provide valuable information about the nature of modes that
destabilize them, as studied recently in Ref. [34]. Another important question is to analyze
the interplay between hydrodynamic interaction and that due to plasma proteins (following
our previous study [4]) and to study how the structure of the phase diagram reported here
evolves in this case.

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