Swimming sheet near a plane surfactant-laden interface
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Swimming sheet near a plane surfactant laden interface

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Abstract

In this work we analyze the velocity of a swimming sheet near a plane surfactant laden interface by assuming the Reynolds number and the sheet’s deformation to be small. We observe a non-monotonic dependence of sheet’s velocity on the Marangoni number ($Ma$) and the surface Péclet number ($Pe_s$). For a sheet passing only transverse waves, the swimming velocity increases with an increase in $Ma$ for any fixed $Pe_s$ while at large $Ma$ it increases and at small $Ma$ it initially increases and then decreases with an increase in $Pe_s$. This dependence of the swimming velocity on $Ma$ and $Pe_s$ is altered if the sheet is passing longitudinal waves in addition to the transverse waves along its surface.
I. INTRODUCTION

Our world is filled with motile microorganisms—*Helicobacter pylori* in stomach, *Escherichia coli* in intestines, *Chlamydomonas* in oceans and snow, *Paramecium* in stagnant basins and ponds etc. Understanding the motion of these organisms in complex flow conditions is essential for addressing several biophysical questions [1]. One such complex bacterial motion is concerned with its locomotion near an interface which has applications in the biofilm formation and bioremediation of an oil spill. Researchers working on this topic have successfully explained (i) how the swimming speed of an organism gets altered near an interface [2–5], (ii) why these organisms move in circles near a plane interface [6, 7] and (iii) why they reorient and get attracted towards an interface [8–11]. Recent works have also analyzed the change in the bacterial dynamics near an interface due to the (i) interface deformation [12–14], (ii) finite inertia of the swimmer or fluid [15, 16], (iii) non-Newtonian suspending fluid [17–19], and (iv) presence of surfactants [9–11].

We briefly review the works on locomotion near a surfactant covered interface. Near a plane surfactant laden interface, the attraction and reorientation of a swimming microorganism is similar to its behavior near a plane wall but its circling direction can be opposite to the one near a clean interface [9]. Considering the trapping of marine microbes onto drops, it was reported that the trapping dynamics outside drops is similar to that outside a rigid sphere, but the surfactant laden drops have better trapping characteristics than a rigid sphere or a clean drop [11]. This is nonintuitive as a surfactant laden drop usually has characteristics that are intermediate between a rigid sphere and a clean drop. Analogous to the interface deformations, finite inertia of the fluid or the organism and the non-Newtonian rheology of the fluid, the surfactant redistribution can enable a time-reversible swimming microorganism near an interface to achieve a net motion [20].

These works on the locomotion near a surfactant laden interface assumed the surfactant to be either incompressible (valid at large Marangoni numbers $Ma$, ratio of Marangoni stresses to the bulk viscous stresses) [21, 23] or compressible [24] but its surface advection being negligible compared to its surface diffusion (valid at small surface Péclet numbers $Pe_s$, ratio of surface advection to the surface diffusion of the surfactant). The objective of this work is to analyze the locomotion near a surfactant laden interface for all values of surface Péclet and Marangoni numbers.
To model the surfactant as generally as possible without losing the analytical tractability, we use a simple model microorganism. Hence, we model the organism as a 2D infinitely long swimming sheet that propels by propagating waves along its surface. This model was first proposed by Taylor [25] who considered a sheet passing transverse waves to represent the monoflagellated organism such as spermatozoon. It was later extended by Blake [26] who allowed the passage of both longitudinal and transverse waves along the sheet to represent the almost flat ciliated organisms such as Paramecium or Opalina. Due to the mathematical simplicity associated with this model, it has been used to study the locomotion (i) in a complex fluid [27–34], (ii) in a gel [35, 36], (iii) in a liquid crystal [37], (iv) in a porous media [38], (v) under confinement [24, 39–45], (vi) at finite inertia of the fluid or the organism [2, 46–48] and (vii) under transient effects [49]. The research on the locomotion under confinement has been restricted to the confinements caused by a rigid or soft wall [2, 3, 39–42, 45], a plane clean interface [4], a deforming membrane [43] or a gel [44] with a Newtonian or a non-Newtonian suspending fluid.

Noting that any interface is inevitably covered with impurities that act as surfactants, it is essential to generalize the theory of locomotion near a clean interface to accommodate the effects of surfactant redistribution. Even though the locomotion under confinement depends on the swimmer’s shape, most of the earlier research on the sheet’s motion under confinement is devoted to a sheet passing transverse waves along its surface (or a Taylor’s swimming sheet [25]). To emphasize the influence of the swimmer’s shape on locomotion and also to generalize the results associated with the sheet passing transverse waves, we analyze the motion of a sheet passing both longitudinal and transverse waves near a surfactant laden interface.

This paper is organized as follows. We provide the governing equations and boundary conditions associated with the motion of a sheet near a plane surfactant laden interface and present the perturbation technique and the solution methodology in Sec. II. We provide simple expressions for the swimming velocity under various limiting conditions in Sec. III. We then analyze the influence of surfactant redistribution on the swimming velocity of a sheet passing only transverse wave and that passing both longitudinal and transverse waves in Secs. IV A and IV B respectively. We finally provide several concluding remarks in Sec. V and present the expressions for the flow field, sheet’s velocity, and validate our results in appendices.
Consider a 2D infinitely long sheet near a surfactant laden interface (see Fig. 1 for schematic). We formulate this problem in a frame of reference moving with the swimming velocity of the sheet. The sheet deforms in such a manner that point \((x, 0)\) on its undeformed surface is located at \(\left(x_0, y_0\right)\) at time \(t\), where \(x_0\) and \(y_0\) are given by

\[
\begin{align*}
x_0 &= x + a \cos k (x + ct) + d \sin k (x + ct), \\
y_0 &= b \sin k (x + ct).
\end{align*}
\]

Here \(a, d (b)\) are the amplitudes of longitudinal (transverse) waves while \(c\) and \(k\) are the wave speed and wave number, respectively.

For analytical tractability, we make the following assumptions. We assume the interface is plane and non-deforming; such a non-deforming interface assumption is valid at small Capillary numbers (ratio of bulk viscous stresses to the capillary stresses). We assume the sheet is located symmetrically between two surfactant laden interfaces which can occur for locomotion in a film [50–52]. Hence, we only solve for the fluid flow above the sheet. We assume the amplitude of sheet’s deformation is much smaller than the wavelength, i.e., \(ak \ll 1, bk \ll 1,\) and \(dk \ll 1\). Hence, we can write \(ak = \varepsilon \tilde{a}, bk = \tilde{b},\) and \(dk = \varepsilon \tilde{d},\) where \(\varepsilon \ll 1\) while the magnitudes of \(\tilde{a}, \tilde{b},\) and \(\tilde{d}\) can be at most \(O(1)\). We denote the distance between the mean plane of the sheet and the interface by \(h\), which is at least as large as the wavelength i.e., \(hk \geq O(1)\). The surfactant is insoluble and we neglect any kind of interfacial rheology imparted by the surfactant to the interface. Insoluble surfactant limit is valid when the interfacial transport of the surfactant is much faster than the bulk transport and the adsorption-desorption between the bulk fluid and the interface. Hence in this limit, the bulk surfactant does not influence the interfacial surfactant transport. Also, assuming the local surfactant concentration on the interface \((\Gamma)\) is much smaller than the maximum possible surfactant concentration on the interface \((\Gamma_{\infty})\), we use a linear constitutive equation to relate the interfacial tension \((\gamma)\) to the surfactant concentration \([53]\) i.e., \(\gamma = \gamma_s - \Gamma RT\). Here \(\gamma_s\) is the interfacial tension of the clean interface, \(R\) is the ideal gas constant and \(T\) is the absolute temperature.

We use the following characteristic parameters to scale all variables to dimensionless variables: a reference length \(l_{ref} = 1/k\), reference velocity \(u_{ref} = c\), reference time \(t_{ref} = 1/(ck)\), reference pressure \(\left(p_{ref}^{(j)}\right)\) and stress \(\left(T_{ref}^{(j)}\right)\) in the \(j\)-th fluid \(p_{ref}^{(j)} = T_{ref}^{(j)} = \mu_j ck\),
FIG. 1: (Color Online) A schematic showing a swimming sheet located near a plane surfactant laden interface. The sheet propels by passing waves along its surface. The distance between the midplane of the sheet and the interface is $h$. We denote the fluid in which the swimmer is suspended as ‘Fluid 1’ while the fluid above the interface as ‘Fluid 2’. In the frame moving with the swimming velocity of the sheet, a uniform streaming flow exists in fluid 2 far away from the sheet with the velocity that is negative of the sheet’s swimming velocity. The origin of the coordinate system is located at the midplane of the sheet.

where $\mu_j$ is the dynamic viscosity of the $j$-th fluid. As the problem is two dimensional, we use the stream function ($\psi$) to solve it. We non-dimensionalize the stream function and surfactant concentration via $\psi_{ref} = c/k$ and $\Gamma_{ref} = \Gamma_{eq}$ respectively, where $\Gamma_{eq}$ is the equilibrium surfactant concentration. We hereby formulate the problem of sheet near a surfactant laden interface in dimensionless variables.

As the inertia of the flow due to the swimming microorganism is typically negligible, the flow field is governed by the Stokes equations and the incompressibility condition. Using the relation between velocity components and the stream function, Eq. (2), the governing equations are simply the biharmonic equations for the stream functions Eq. (3)

$$u^{(j)} = \frac{\partial \psi^{(j)}}{\partial y}; v^{(j)} = -\frac{\partial \psi^{(j)}}{\partial x}, \text{ where } j = 1, 2,$$

$$\nabla^4 \psi^{(j)} = 0,$$

where $u^{(j)}$ and $v^{(j)}$ indicate the $x$ and $y$ components of the velocity of $j$-th fluid while $\psi^{(j)}$ denotes the stream function concerning the $j$-th fluid. Similar to this work, the stream
function was governed by the biharmonic equation in the works of Taylor [25], Katz and [3] Man & Lauga [4]. Point \((x, 0)\) on the undeformed sheet is located at \((x_0, y_0)\) on the sheet’s surface at time \(t\), where \((x_0, y_0)\) is given by

\[
x_0 = x + \epsilon \left[ \tilde{a} \cos (x + t) + \tilde{d} \sin (x + t) \right],
\]

\[
y_0 = \epsilon \tilde{b} \sin (x + t).
\]

This sheet’s shape is the same as that considered by Katz but differs from that of Taylor because Taylor’s sheet passes only transverse waves along positive \(x\)-direction \((x_0 = x, y_0 = \epsilon \tilde{b} \sin (x - t))\). As the flow field in phase 1 should satisfy the no-slip and no-penetration boundary conditions on the sheet, we have

\[
\left. \frac{\partial \psi^{(1)}}{\partial y} \right|_{(x_0, y_0)} = \epsilon \left[ -\tilde{a} \sin (x + t) + \tilde{d} \cos (x + t) \right],
\]

\[
\left. \frac{\partial \psi^{(1)}}{\partial x} \right|_{(x_0, y_0)} = -\epsilon \tilde{b} \cos (x + t).
\]

This boundary condition differs from that of Taylor due to a different shape of the sheet and the direction of wave propagation but it also differs from that of Katz because Katz works in a frame moving with velocity \((U - 1) \mathbf{i}\) unlike the frame moving with velocity \(U \mathbf{i}\) considered by us and Taylor. Here \(U \mathbf{i}\) is the swimming velocity while \(\mathbf{i}\) being the unit vector along \(x\) direction. Either neglecting gravity or by assuming the density of the sheet is the same as the density of the fluid in which it is suspended, we find that the external force on the sheet is zero. Consequently, due to negligible inertia of the sheet, the net hydrodynamic force on the sheet must be zero i.e., \(\int_{\text{Sheet}} \mathbf{n} \cdot \mathbf{T}^{(1)} \big|_{\text{Sheet}} d\mathcal{S} = \mathbf{0}\) which simplifies to

\[
\int_{0}^{2\pi} \left\{ -\epsilon \tilde{b} \cos (x + t) \mathbf{i} + \left[ 1 - \epsilon \tilde{a} \sin (x + t) + \epsilon \tilde{d} \cos (x + t) \right] \mathbf{j} \right\} \cdot \mathbf{T}^{(1)} \big|_{(x_0, y_0)} dx = \mathbf{0},
\]

where \(\mathbf{j}\) denotes the unit vector along \(y\) direction. Also, \(\mathbf{n}\) is the normal to the sheet pointing into fluid 1 while \(\mathbf{T}^{(j)}\) is the stress tensor for \(j\)-th fluid. Using the Newtonian fluid constitutive equation, \(\mathbf{T}^{(j)}\) can be written in terms of pressure \((p^{(j)})\) and velocity fields \((\mathbf{v}^{(j)})\) of the \(j\)-th fluid as \(\mathbf{T}^{(j)} = -p^{(j)} \mathbf{I} + \left[ \nabla \mathbf{v}^{(j)} + (\nabla \mathbf{v}^{(j)})^\dagger \right]\), where \(\dagger\) stands for the transpose and \(\mathbf{I}\) is an identity tensor. The force-free condition is identically satisfied for the Taylor’s sheet in an unbounded fluid. Since Katz considers a sheet asymmetrically located between two walls, the sum of the hydrodynamic forces acting on the top and bottom surfaces of
the sheet must be zero and this force-free condition of Katz reduces to Eq. (6) for a sheet symmetrically located between two walls. Far away from the sheet, the velocity of fluid 2 should approach the negative of sheet’s swimming velocity \((U = U_i)\). In terms of stream function, this condition simplifies to

\[
as y \to \infty, \psi^{(2)} \sim -U y.
\]

A similar condition exists for the fluid in which the sheet is immersed in Taylor’s work but no far-field condition exists for a sheet bounded by walls as considered by Katz. Since the interface is non-deforming, the fluid velocity (in both phases) at the interface but normal to the interface must be zero,

\[
at y = h : \frac{\partial \psi^{(1)}}{\partial x} = 0; \frac{\partial \psi^{(2)}}{\partial x} = 0.
\]

A similar boundary condition is applied at the walls in Katz’s work to ensure that the fluid does not penetrate the walls. The fluid velocity at the interface but tangential to the interface must be continuous across the interface

\[
at y = h : \frac{\partial \psi^{(1)}}{\partial y} = \frac{\partial \psi^{(2)}}{\partial y}.
\]

Also, the jump in the tangential stresses across the interface should be balanced by the Marangoni stresses which using the relationship between interfacial tension and the surfactant concentration simplifies to

\[
at y = h : \lambda T_{yx}^{(2)} - T_{yx}^{(1)} = Ma \frac{\partial \Gamma}{\partial x}; \ Ma = \frac{RT\Gamma_{eq}}{\mu_1 c},
\]

where \(T_{yx}^{(j)}\) denotes the \(yx\) component of stress tensor in the \(j\)-th fluid and the viscosity ratio \(\lambda = \mu_2/\mu_1\). Finally, the transport of an insoluble surfactant on a non-deforming interface is governed by

\[
P_{es} \left[ \frac{\partial \Gamma}{\partial t} + \frac{\partial}{\partial x} \left( \Gamma u_{|y=h} \right) \right] = \frac{\partial^2 \Gamma}{\partial x^2}; \ P_{es} = \frac{c}{kD_s},
\]

where \(D_s\) is the surface or interface diffusivity of the surfactant and the slip of an interface is written as \(u_{|y=h} = \frac{\partial \psi^{(1)}}{\partial y}_{y=h} = \frac{\partial \psi^{(2)}}{\partial y}_{y=h}\). Instead of the continuous tangential velocity condition (Eq. (9)), shear stress balance condition (Eq. (10)) and the surfactant transport equation (Eq. (11)) at the interface, we simply have a no slip boundary condition at the wall in Katz’s work.
We need to solve Eqs. (3)-(11) to determine the swimming velocity of the sheet, $U_i$. As the surfactant transport equation is nonlinear, it is not possible to solve these equations analytically for an arbitrary value of $\epsilon$. However, for $\epsilon \ll 1$, we can use the following traditional technique to find the leading order approximation of the sheet’s swimming velocity: (i) express the boundary condition on the sheet’s surface as a series of boundary conditions applied at the mean plane of the sheet by doing a Taylor series expansion of any function $f(x_0, y_0)$ about $(x, 0)$ and (ii) expand all the variables as a power series in $\epsilon$

\[
\left\{v^{(j)}, T^{(j)}, p^{(j)}, U, \psi^{(j)} \right\}_{y=h} = \sum_{n=1}^{\infty} \epsilon^n \left\{v^{(j)}_n, T^{(j)}_n, p^{(j)}_n, U_n, \psi^{(j)}_n \right\}_{y=h}
\]

\[
\Gamma = \sum_{n=1}^{\infty} \epsilon^n \Gamma_n
\]

The resulting perturbed equations are linear and have the following general solution for the stream function and the surfactant concentration at $O(\epsilon^n)$

\[
\psi^{(1)}_n = \sum_{m=1}^{\infty} \left\{ \left[ (A^{(1)}_{n,m} + E^{(1)}_{n,m} y) \cos m(x + t) + (B^{(1)}_{n,m} + F^{(1)}_{n,m} y) \sin m(x + t) \right] \cosh my \\
+ \left[ (C^{(1)}_{n,m} + G^{(1)}_{n,m} y) \cos m(x + t) + (D^{(1)}_{n,m} + H^{(1)}_{n,m} y) \sin m(x + t) \right] \sinh my \right\}
\] + $\alpha_n y + \beta_n y^2 + \gamma_n y^3$

\[
\psi^{(2)}_n = -U_n y + \sum_{m=1}^{\infty} \left[ (A^{(2)}_{n,m} + E^{(2)}_{n,m} y) \cos m(x + t) + (B^{(2)}_{n,m} + F^{(2)}_{n,m} y) \sin m(x + t) \right] e^{-my},
\]

\[
\Gamma_n = \sum_{m=1}^{\infty} \left[ J_{n,m} \cos m(x + t) + L_{n,m} \sin m(x + t) \right],
\]

where $A^{(1)}_{n,m}$, $B^{(1)}_{n,m}$, $C^{(1)}_{n,m}$, $D^{(1)}_{n,m}$, $E^{(1)}_{n,m}$, $F^{(1)}_{n,m}$, $G^{(1)}_{n,m}$, $H^{(1)}_{n,m}$, $A^{(2)}_{n,m}$, $B^{(2)}_{n,m}$, $E^{(2)}_{n,m}$, $F^{(2)}_{n,m}$, $\alpha_n$, $\beta_n$, $\gamma_n$, $J_{n,m}$, and $L_{n,m}$ are unknown constants that need to be determined while satisfying the perturbed boundary conditions. The expressions of the constants that determine the $O(\epsilon)$ and $O(\epsilon^2)$ flow fields along with the expression of the leading order swimming velocity $U_2$ are provided in Appendix A. We also verify our calculation by comparing the leading order swimming velocity of the sheet $U_2$ with that reported in the literature in various limits. This validation is presented in Appendix B.
III. LIMITING CASES

We note that the swimming velocity depends in a complex fashion on $Ma$ and $Pe_s$ as can be seen in Eq. (A15) derived for a sheet passing transverse waves near an air-water interface. So, to understand the surfactants influence on the swimming velocity $U_2$, we need to numerically evaluate $U_2$ for various $Ma$ and $Pe_s$. Before doing this, in this section we report some simple expressions for the swimming velocity in the limits of small or large $Ma$ or $Pe_s$. Through such expressions, one can easily find out how the surfactant redistribution affects the swimming velocity. We report these limiting forms for a sheet near an air-water interface ($\lambda = 0$) by noting that the leading order swimming occurs at $O (\epsilon^2)$ i.e., $U = \epsilon^2 U_2 + O (\epsilon^4)$.

At small $Ma$, we get

$$U_2 = U_{2,\text{clean}} + \frac{Pe_s Ma}{(Pe_s^2 + 1) (\sinh (2h) - 2h)^2} \left[ \left( h^2 \cosh (2h) - 2h \sinh (2h) \right) \tilde{b} \left( \tilde{a} - Pe_s \tilde{d} \right) + h^2 + \cosh (2h) - 1 \right]$$

$$+ O \left( Ma^2 \right)$$

(16)

Here $U_{2,\text{clean}}$ is the swimming velocity near a plane clean interface whose expression for $\tilde{a} = \tilde{d} = 0$ is given in Eq. (B3). We see that the correction due to surfactant redistribution depends nonlinearly on $Pe_s$. As long as $\tilde{d} = 0$, we see that the surfactant redistribution increases the sheet’s velocity because $(h \sinh (2h) - \cosh (2h) + 1) > 0$ and $(h^2 \cosh (2h) - 2h \sinh (2h) + h^2 + \cosh (2h) - 1) > 0$ for all $h$. When $\tilde{a} = 0$, the surfactant redistribution can increase or even decrease the sheet’s velocity depending on whether the following ratio is respectively great than or less than 1.

$$\frac{(h \sinh (2h) - \cosh (2h) + 1) h}{(h^2 \cosh (2h) - 2h \sinh (2h) + h^2 + \cosh (2h) - 1) Pe_s d} \tilde{b}$$

(17)

In the more general case when $\tilde{a} \neq 0$, $\tilde{d} \neq 0$, and $\tilde{b} \neq 0$, the surfactant redistribution can increase or decrease the swimmer’s velocity depending on $Pe_s$, $h$ and relative order of magnitude of the wave amplitudes.

At small $Pe_s$, we get

$$U_2 = U_{2,\text{clean}} + \frac{Pe_s Ma \times \tilde{b}}{(\sinh (2h) - 2h)^2} \left[ \tilde{a} (h^2 \cosh (2h) - 2h \sinh (2h) + h^2 + \cosh (2h) - 1) \right]$$

$$+ O \left( Pe_s^2 \right)$$

(18)
From this expression, we see that the swimming velocity does not depend on ‘d’ modes. Also as \((h^2 \cosh (2h) - 2h \sinh (2h) + h^2 + \cosh (2h) - 1) > 0\) and \((h \sinh (2h) - \cosh (2h) + 1) > 0\), we see that the surfactant redistribution always increases the swimming velocity at small \(Pe_s\). This is in contrast with the observation that the surfactant redistribution can increase or even decrease the velocity of a swimming microorganism inside a surfactant laden drop at small \(Pe_s\)\(^{[20]}\). These different influences of the surfactant on the swimming velocity, reported in Ref. \([20]\) and this work, might be due to different shapes of the swimmer and the interface used in these two works.

At large \(Ma\), we get

\[
U_2 = U_{2,\text{wall}} - \frac{4b}{Pe_s Ma(2h^2 - \cosh (2h) + 1)^2} \left[ \left( \frac{h^2 \cosh (2h) - 2h \sinh (2h) + h^2 + \cosh (2h) - 1}{2} \right) \left( \vec{a} + \vec{d} \times Pe_s \right) + (h \sinh (2h) - \cosh (2h) + 1) \hat{b} \right] + O(Ma^{-2})
\]  

(19)

Here \(U_{2,\text{wall}}\) is the swimming velocity near a plane wall whose expression is given in Eq. \((B2)\). Again as \((h^2 \cosh (2h) - 2h \sinh (2h) + h^2 + \cosh (2h) - 1) > 0\) and \((h \sinh (2h) - \cosh (2h) + 1) > 0\), we see that the swimming velocity near a surfactant laden interface is always less than that near a plane wall at large \(Ma\).

IV. RESULTS

In this section, we examine the influence of surfactant redistribution on the swimming velocity of the sheet near a surfactant laden interface.

It was reported that the swimming velocity of a sheet, propagating longitudinal waves, near a plane wall is the same as its velocity in an unbounded fluid \(^{[3]}\) i.e., \(U = -\frac{\epsilon^2}{2} \left( \vec{a}^2 + \vec{d}^2 \right) + O(\epsilon^4)\). Our analysis suggests the presence of a surfactant laden interface instead of a wall near such a sheet, does not modify its velocity, i.e., velocity of a sheet propagating longitudinal waves near a plane surfactant laden interface is the same as its velocity in an unbounded fluid.

As we have analyzed the velocity of a sheet propagating longitudinal waves near a plane surfactant laden interface, we then proceed to examine the velocity of a sheet (near a surfactant laden interface) passing only transverse waves and that passing both longitudinal and
FIG. 2: (Color Online) The variation of the swimming velocity with the distance between the interface and (the mean plane of) the sheet for a sheet passing only transverse waves (blue solid line) and that passing both longitudinal and transverse waves (red dashed line).

Here the swimming velocity is normalized with the swimming velocity of a sheet passing only transverse waves in an unbounded fluid \((U_{2T∞})\). The viscosity ratio \(λ = 0\), Marangoni number \(Ma = 1\) and the surface Péclet number \(Pe_s = 1\).

transverse waves in the following two subsections. Even though the expressions for the leading order swimming velocity are derived for any arbitrary viscosity ratio \(λ\), we only report the influence of surfactant redistribution on this velocity for an air-water interface \((λ = 0)\) as experiments on locomotion in films are mainly performed for an air-water interface [50–52].

For a fixed \(Ma\) and \(Pe_s\), we found that the swimming velocity increases as \(h\) decreases (see Fig. 2). Similar trend with decreasing \(h\) was reported for confinements caused by a rigid plane wall [3], a plane clean interface [4] or a gel [44]. Also, the variation of the swimming velocity with \(Ma\) and \(Pe_s\) is qualitatively the same for any fixed value of \(h\). For this reason, we report and analyze this variation with \(Ma\) and \(Pe_s\) for a typical value of \(h = 1\).

We report here the typical values of \(Ma\) and \(Pe_s\). For the tail of a spermatozoon [54], the wave speed \(c \sim (200 − 1200) \times 10^{-6} \text{ m/s}\) and the wave number \(k \sim (1.14 − 4.18) \times 10^{5} \text{ m}^{-1}\). The maximum possible surfactant concentration at an air-water interface [55, 56] \(\Gamma_∞ \sim 10^{-6} − 10^{-4} \text{ mol/m}^2\). Assuming that the equilibrium surfactant concentration \(\Gamma_{eq} \sim (10^{-3} − 10^{-1}) \Gamma_∞\), we get \(\Gamma_{eq} \sim (10^{-9} − 10^{-5}) \text{ mol/m}^2\). The surface diffusivity of the surfactant [57] \(D_s \sim (1 − 10) \times 10^{-9} \text{ m}^2/\text{s}\). Using the definitions of \(Ma\), \(Pe_s\) along with the
viscosity of water ($\mu_1$) and the temperature of $T = 298$ K, we get $Ma \sim O(1) - O(10^5)$ and $Pe_s \sim 0.05 - O(10)$.

A. Sheet passing only transverse wave along its surface

![Graph](image)

FIG. 3: (Color Online) For a sheet passing transverse waves near a plane surfactant laden interface, the variation of the leading order swimming velocity with (a) $Ma$ for $Pe_s = 1$ and (b) $Pe_s$ for $Ma = 1$, 10, and 100. Here the swimming velocity is normalized with the swimming velocity of the same sheet in an unbounded fluid ($U_{2T\infty}$). The distance between the midplane of the sheet and the interface is $h = 1$, the viscosity ratio $\lambda = 0$, and the amplitudes of the longitudinal waves $a = d = 0$.

When a sheet passing only transverse waves is located near a plane surfactant laden interface, the dependence of its swimming velocity on $Ma$ (for fixed $Pe_s$) and on $Pe_s$ (for fixed $Ma$) is given in Fig. 3. We observe the following from this figure. As the minimum velocity occurs at $Ma = 0$ or $Pe_s \to 0$ and since $Ma = 0$ or $Pe_s \to 0$ represents a clean interface, we conclude that the swimming velocity near a plane surfactant laden interface is always more than that near a plane clean interface. Near a plane surfactant laden interface, (a) the swimming velocity increases with an increase in $Ma$ for any fixed $Pe_s$, (b) it increases with an increase in $Pe_s$ for any fixed but large $Ma$ i.e., $Ma \geq O(10)$ and (c) it initially increases and then decreases with an increase in $Pe_s$ for any fixed but small $Ma$ i.e., $Ma \leq O(1)$. 
B. Sheet passing both longitudinal and transverse waves along its surface

![Graph](b.png)

FIG. 4: (Color Online) For a sheet passing both longitudinal and transverse waves near a plane surfactant laden interface, the variation of the leading order swimming velocity with (a) $Ma$ for $Pe_s = 0.1, 1, 10, 100$ and with (b) $Pe_s$ for $Ma = 1, 10, 100$. Here the swimming velocity is normalized with the swimming velocity of a sheet passing transverse waves in an unbounded fluid ($U_2T_\infty$). The distance between the midplane of the sheet and the interface is $h = 1$, the viscosity ratio $\lambda = 0$, and the amplitudes of the waves $a = 0$, and $d/b = 2$.

In this section, we discuss how the effect of surfactant redistribution on the swimming velocity gets modified, in comparison to this effect presented in the earlier section, if a sheet is passing both longitudinal and transverse waves along its surface. For this purpose, we plot the variation of the swimming velocity with $Ma$ (for a fixed $Pe_s$) and $Pe_s$ (for a fixed $Ma$) in Fig. 4. From this figure, we observe the following. The swimming velocity for a sheet near a plane surfactant laden interface can be more or even less than that for a sheet near a plane clean interface ($Ma \to 0$ or $Pe_s \to 0$). Unlike the case of a sheet passing only transverse waves, the swimming velocity near a surfactant laden interface might not lie in between the swimming velocity near a plane clean interface and that near a plane wall. Desai et al. [11] reported a similar observation by noting that the critical trapping radius of a swimming microorganism outside a surfactant laden drop is always less than the critical trapping radius near a clean drop and that near a rigid sphere. When a sheet is in the vicinity of a surfactant laden interface, (a) the swimming velocity increases with an increase
in $Ma$ for a fixed but small $Pe_s \ (\leq O(1))$, (b) it initially decreases and then increases with an increase in $Ma$ for a fixed but large $Pe_s \ (\geq O(10))$, (c) it increases with an increase in $Pe_s$ for a fixed yet large $Ma \ (\geq O(10))$, (d) it initially increases and then decreases with an increase in $Pe_s$ for a fixed but small $Ma \ (\leq O(1))$.

It seems non-intuitive for the swimming velocity near a surfactant laden interface to lie outside the range corresponding to the one near a clean interface and a rigid wall. If we had a rigid particle moving near a surfactant laden interface due to some fixed force acting on it, its velocity always lies in between its velocities near a clean interface and a plane wall. So, the motion of organisms is defying the intuition built on the basis of particle motion. This is not the first time that such phenomenon is reported. Considering the motion of particles or organisms in complex fluids, we find that the particle experiencing a fixed force moves faster in a shear-thinning fluid than that in a Newtonian fluid but the velocity of a swimming microorganism in a shear-thinning fluid can be more or even less than that in a Newtonian fluid. Similar observation has been made for the velocity of a swimming microorganism in a viscoelastic fluid. Even though most of the observations in Figs. 3-4 can be qualitatively explained by analyzing the influence of surfactant redistribution on the amplitude of leading order interface slip (see Appendix C), such reasoning is not useful to understand the swimming velocity near a surfactant laden interface if this velocity lies outside the range bounded by the velocities near a clean interface and a plane wall. Consequently, the amplitude of leading order interface slip is not determinative of the swimming velocity changes. Hence to get a proper understanding of the variation of the swimming velocity with all values of $Ma$ and $Pe_s$, we need to correlate the surfactant induced changes in either (i) the amplitude and phase of the leading order slip (not just the amplitude) or (ii) the second order slip (instead of first order slip) with the swimming velocity which are not done in this work.

C. Apparent viscosity ratio

As the expression for the swimming velocity of a sheet near a surfactant laden interface is quite lengthy, we represent the surfactant laden interface as a clean interface with a modified or apparent viscosity ratio ($\lambda_{app}$) to convey the effects of surfactant redistribution in a succinct manner. The expression for $\lambda_{app}$ is simpler than the expression for the swimming velocity.
velocity. Using this expression for $\lambda_{app}$ and the dependence of sheet’s velocity on the viscosity ratio for a sheet near a clean interface \[4\], we can evaluate the sheet’s velocity at this apparent viscosity ratio to find its velocity near a surfactant laden interface. To determine the apparent viscosity ratio, we simply equate the sheet’s velocity near a surfactant laden interface with zero viscosity ratio to its velocity near a clean interface with a viscosity ratio that is equal to the apparent viscosity ratio. We then solve this equation for the apparent viscosity ratio to obtain

$$\lambda_{app} = \frac{-2 \left(\frac{1}{2} Ma \frac{\tilde{b}}{\tilde{d}} Pe_s + \tilde{a}\right) h -1/2 Ma Pe_s \tilde{a}}{\left(-2 Ma Pe_s \tilde{a} b h^3 + (2 Ma Pe_s \tilde{a} - 2 \tilde{b}) h^2 \right)} \sinh (h)$$

which simplifies for a sheet passing only transverse waves to

$$\lim_{\tilde{a} \to 0, \tilde{d} \to 0} \lambda_{app} = \frac{2 Ma Pe_s (- \sinh (2h) - 1/2 Ma Pe_s \cosh (2h) + Ma (h^2 + 1/2) Pe_s + 2h)}{(-4 Pe_s^2 - 4) \sinh (2h) - 2 Ma Pe_s \cosh (2h) + 8 Pe_s^2 h + (4h^2 + 2) Ma Pe_s + 8h}$$

(21)

In the limit $Ma \to 0$ or $Pe_s \to 0$, the surfactant laden interface behaves like a clean interface due to which $\lambda_{app}$ approaches the actual viscosity ratio that is zero. Hence, $\lambda_{app} \to 0$ as $Ma \to 0$ or $Pe_s \to 0$. In the limit $Ma \to \infty$, the surfactant becomes incompressible \[10, 58\] and the surfactant laden interface behaves like a rigid wall (see Appendix B for discussion on the incompressible surfactant). Hence, $\lambda_{app} \to \infty$ as $Ma \to \infty$. As $Pe_s \to \infty$, $\lambda_{app}$ approaches a constant value given by

$$\lim_{Pe_s \to \infty, \tilde{a} \to 0, \tilde{d} \to 0} \lambda_{app} = \frac{Ma^2 ((\cosh (h))^2 - h^2 - 1)}{4 \sinh (h) \cosh (h) - 4h}.$$  

(22)
for a sheet passing only transverse waves.

For a sheet passing transverse wave near a surfactant laden interface, typical variation of $\lambda_{\text{app}}$ with $Pe_s$ for various $Ma$ is shown in Fig. 5. From this figure, we observe that $\lambda_{\text{app}}$ increases with an increase in $Pe_s$ at large $Ma$ ($\geq O(10)$) while it initially increases and then decreases with an increase in $Pe_s$ at small $Ma$ ($\leq O(1)$). Combining this dependence of $\lambda_{\text{app}}$ on $Ma$, $Pe_s$ with the dependence of $U_2$ on $\lambda$ for a sheet near a clean interface (Fig. 6b), we expect the velocity of a sheet near a surfactant laden interface to increase monotonically (resp. vary non-monotonically) with an increase in $Pe_s$ at large $Ma$ (resp. at small $Ma$), consistent with the observations of Fig. 3b.

![Graph showing variation of apparent viscosity ratio with $Pe_s$ at $Ma = 1$, $Ma = 10$, and $Ma = 100$.](image)

**FIG. 5**: (Color Online) The variation of apparent viscosity ratio with $Pe_s$ at $Ma = 1$ (blue solid line), 10 (red dashed line) and 100 (yellow dash-dotted line). Here, the sheet is passing only transverse waves, hence $a = d = 0$. The actual viscosity ratio of the surfactant laden interface $\lambda = 0$ while $h = 1$.

V. CONCLUSIONS

We aimed to understand the dependence of a microorganism’s swimming velocity on the Marangoni number ($Ma$) and the surface Péclet number ($Pe_s$) when it is near a surfactant covered interface. For this purpose, we derived the velocity of a 2D infinitely long swimming sheet near a surfactant laden interface under the assumptions of zero Reynolds number, zero interface deformation and small sheet’s deformation. We observed that the swimming
velocity near a surfactant laden interface can be more or even less than that near a clean interface and this velocity varies non-monotonically with $Ma$ and $Pe_s$, these observations being highly sensitive to the type of wave passing through the sheet. Unlike the rigid particles near a surfactant laden interface, we found that the swimming microorganisms near such an interface can have a velocity that does not lie in between their velocities near a clean interface and a rigid wall. To succinctly express the effects of surfactant redistribution, we represented the surfactant laden interface as a clean interface with a viscosity ratio equal to apparent viscosity ratio whose expression is found by equating the swimming velocities near a clean and surfactant laden interface.

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**Appendix A: Expressions for constants that appear in stream function and surfactant concentration**

In this section, we provide expressions for the constants appearing in Eqs. (13)-(15) that enable us to determine the stream function and the surfactant concentration at various orders of $\epsilon$. Even though we derived these expressions for any arbitrary viscosity ratio $\lambda$ and for non-zero wave amplitudes $\tilde{a} \neq 0$, $\tilde{b} \neq 0$, and $\tilde{d} \neq 0$, as these expressions are lengthy, we provide these expressions only for the special case of an air-water interface ($\lambda = 0$) and sheet passing only transverse waves ($\tilde{a} = 0, \tilde{d} = 0$).

1. Expressions for constants appearing in $O(\epsilon)$ stream function and surfactant concentration

\[
A_{1,1}^{(1)} = 0, \quad (A1)
\]
\[
B_{1,1}^{(1)} = -\tilde{b}, \quad (A2)
\]
\[
-C_{1,1}^{(1)} = E_{1,1}^{(1)} = \frac{2(\sinh(h))^2 Pe_s^2 Ma \tilde{b} h^2}{\text{denom}_1}, \quad (A3)
\]
\[-D_{1,1}^{(1)} = F_{1,1}^{(1)} = \frac{-4Ma(cosh(h))^4Pe_s - ((Ma^2 + 4)Pe_s^2 + 4) sinh(h) (cosh(h))^3 \\
+ ((-Ma^2h + 4h) Pe_s^2 + (2h^2 + 4) MaPe_s + 4h) (cosh(h))^2 \\
+ Ma^2Pe_s^2 sinh(h) (h^2 + 1) cosh(h) + hPe_s Ma (Ma (h^2 + 1) Pe_s + 2h)}{denom_1} \,
\] (A4)

\[G_{1,1}^{(1)} = \frac{-2Pe_s^2hMa \bar{b} sinh(h) (cosh(h) h - sinh(h))}{denom_1} \,
\] (A5)

\[H_{1,1}^{(1)} = \frac{-sinh(h) \bar{b} \left( -4MaPe_s (cosh(h))^3 - ((Ma^2 + 4)Pe_s^2 + 4) sinh(h) (cosh(h))^2 \\
+ (4Pe_s^2h + (2h^2 + 4) MaPe_s + 4h) cosh(h) \\
+ Pe_s sinh(h) Ma (Ma (h^2 + 1) Pe_s + 2h) \right)}{denom_1} \,
\] (A6)

\[A_{1,1}^{(2)} = \frac{-2h^2e^h Ma sinh(h) Pe_s^2 \bar{b} ((cosh(h))^2 - h^2 - 1)}{denom_1} \,
\] (A7)

\[B_{1,1}^{(2)} = \frac{2h^2 \bar{b} \left( -Ma(cosh(h))^2Pe_s + (-2Pe_s^2 - 2) sinh(h) cosh(h) \\
+ 2Pe_s^2h + Ma (h^2 + 1) Pe_s + 2h \right) sinh(h) e^h}{denom_1} \,
\] (A8)

\[E_{1,1}^{(2)} = \frac{2e^h Ma sinh(h) Pe_s^2 \bar{b}h ((cosh(h))^2 - h^2 - 1)}{denom_1} \,
\] (A9)

\[F_{1,1}^{(2)} = \frac{-2h \bar{b} \left( -Ma(cosh(h))^2Pe_s + (-2Pe_s^2 - 2) sinh(h) cosh(h) \\
+ 2Pe_s^2h + Ma (h^2 + 1) Pe_s + 2h \right) sinh(h) e^h}{denom_1} \,
\] (A10)

\[J_{1,1} = \frac{2hPe_s sinh(h) \bar{b} ( -Ma(cosh(h))^2Pe_s - 2cosh(h) sinh(h) + Ma (h^2 + 1) Pe_s + 2h)}{denom_1} \,
\] (A11)

\[L_{1,1} = \frac{4Pe^2 \bar{b}h sinh(h) ( -cosh(h) sinh(h) + h)}{denom_1} \,
\] (A12)

where

\[denom_1 = ((Ma^2 + 4) Pe_s^2 + 4) (cosh(h))^4 + 4Ma(cosh(h))^3 sinh(h) Pe_s \\
+ (-4 + 4 + (-2h^2 - 2) Ma^2) Pe_s^2 - 4MaPe_s h) (cosh(h))^2 \\
- 4 (2Pe_s^2h + Ma (h^2 + 1) Pe_s + 2h) sinh(h) cosh(h) \\
+ ((h^2 + 1)^2 Ma^2 + 4h^2) Pe_s^2 + 4hMa (h^2 + 1) Pe_s + 4h^2, \]

\[U_1 = \alpha_1 = \beta_1 = \gamma_1 = 0. \] (A13)
2. Expressions for constants appearing in $O(\varepsilon^2)$ stream function

\[ \beta_2 = \gamma_2 = 0, \quad (A14) \]

\[
\begin{align*}
U_2 &= -\alpha_2 = \\
&= \frac{\beta_2}{\left[\left(2 \left(\frac{h^2+1}{2} \right) \text{Ma}^2 - 8 h \text{Pa}_s \text{Ma} \frac{h^2+1}{2} - 8 \frac{h^2}{2} \right) \text{Pe}^2 + 8 h \text{Ma} \left(\frac{h^2+1}{2}\right) \text{Pe} + 8 h^2 \right]}
\end{align*}
\]

\[ (A15) \]

Appendix B: Validation of results

In this section, we verify the expression for the velocity of a sheet near a surfactant laden interface by comparing it with similar expressions derived in the literature for several special cases. For instance, when a sheet is far away from the interface, its velocity should be the same as the sheet’s velocity in an unbounded fluid \[26\]

\[ \lim_{\lambda \to \infty} U_2 = 1 \]  \[\text{or} \quad \text{or} \quad \lim_{\text{Ma} \to \infty} U_2 = \] \[ \frac{1}{2} \left(-\bar{a}^2 - \bar{d}^2 + \bar{b}^2 + 2\bar{a}\bar{b}\right) \]  \[\text{B1}\]

As $\lambda \to \infty$ or $\text{Ma} \to \infty$, the sheet’s velocity should approach the velocity of a sheet placed symmetrically between two plane walls \[3\].

\[ \lim_{\lambda \to \infty} U_2 = \lim_{\text{Ma} \to \infty} U_2 = \frac{\left(\bar{a}^2 - \bar{b}^2 + \bar{d}^2\right) (\cosh(h))^2 - 2 \cosh(h) \sinh(h) \bar{a}\bar{b}}{2h^2 - 2\left(\cosh(h)\right)^2 + 2} \]

\[ \text{B2} \]

It is obvious that a surfactant laden interface behaves as a rigid wall in the limit $\lambda \to \infty$ but such a behavior in the limit $\text{Ma} \to \infty$ requires some explanation. As $\text{Ma} \to \infty$, the surfactant becomes incompressible \[10, 58\], i.e., $O(1)$ changes in interfacial tension are caused by infinitesimal changes in the surfactant concentration. So, treating $\Gamma$ as constant, the surfactant transport equation simplifies to $\frac{\partial}{\partial x} \left(u|_{y=h}\right) = 0$. The only solution of this equation that enables the velocity of fluid 2 to approach the negative of sheet’s velocity, far
FIG. 6: (Color Online) For a sheet passing transverse waves near a plane clean interface, the variation of the (a) leading order interface slip and (b) leading order swimming velocity with the viscosity ratio $\lambda$. Here the swimming velocity is normalized with the swimming velocity of the same sheet in an unbounded fluid ($U_{2T\infty}$). The values of other parameters $h$, $a$, and $d$ are kept the same as those of Fig. 3.

away from the interface, is $u|_{y=h} = -U$. The sheet near an interface with such interface boundary conditions $(u|_{y=h} = -U, v|_{y=h} = 0)$ is essentially equivalent to a sheet placed equidistantly between two plane walls.

In the limit $Ma \to 0$ or $Pe_s \to 0$, the sheet’s velocity should approach its velocity near a plane clean interface. This velocity was derived for a sheet passing transverse waves in the positive $x$-direction (Taylor’s sheet) (4). As we considered the sheet passing transverse waves in the negative $x$-direction, its velocity near a plane surfactant laden interface in the limits $Ma \to 0$ or $Pe_s \to 0$ should approach the negative of Taylor’s swimming sheet velocity near a plane clean interface.

$$\lim_{Ma, \tilde{a}, d \to 0} U_2 = \lim_{Pe_s, \tilde{a}, d \to 0} U_2 = \frac{\tilde{b}^2}{2} - \frac{\tilde{b}^2 h (h\lambda + 1)}{h^2 \lambda - (\cosh (h))^2 \lambda - \sinh (h) \cosh (h) + h + \lambda} \quad (B3)$$

**Appendix C: Effect of surfactant redistribution on the interface slip**

In this section, we analyze the influence of surfactant redistribution on the leading order interface slip. Let us first discuss how the viscosity ratio of an interface affects the interface
FIG. 7: (Color Online) Comparison of interface slip of a clean interface (blue solid line) with that of a surfactant laden interface (blue dotted line). Also plotted is the surfactant concentration on the surfactant laden interface (red dash-dotted line). The vertical dashed lines are just for reference. The blue (upper) and red (lower) arrows denote, respectively, the vector field of clean interface slip and the direction of Marangoni induced slip. The axis for the interface slip is on the left while that for surfactant concentration is on the right. Here, $Ma = 10$ and $Pe_s = 1$ for the surfactant laden interface. Also, $\epsilon = 0.1$ is used to calculate $\Gamma$ while the values of the other parameters $h$, $\lambda$, $a$, and $d$ are kept the same as those of Fig. 3.

slip and the swimming velocity for a sheet near a plane clean interface. We plot these features for a sheet passing transverse waves near a plane clean interface in Fig. 6. We observe that the interface slip decreases while the swimming velocity increases with an increase in the viscosity ratio of a clean interface. As an increase in the viscosity ratio corresponds to an increase in the viscosity of fluid above the interface (for a fixed viscosity of fluid below the interface), such high viscosity fluid above the interface hinders the interface slip; this explains the decrease in slip with an increase in $\lambda$.

When a sheet is near a clean interface, the interface slip varies in a sinusoidal fashion with the $x$-coordinate, i.e., $u_1|_{y=h} = A \sin(x + t + \phi)$, where $A$, and $\phi$ are the amplitude and phase of the slip, respectively (see blue solid line in Fig. 7). vector field of the interface slip is shown by blue (upper) arrows in this figure). When we compare slip (such as slip of clean and surfactant laden interface or slip of a surfactant laden interface at various values of $Ma$ or
Pe_s), we are essentially comparing the amplitude of the sinusoidal functions. Now, if a sheet is near a surfactant laden interface, the surfactant concentration varies non-monotonically with the position, as shown by red dash-dotted lines in Fig. 7. As the interfacial tension decreases with an increase in the surfactant concentration, this non-homogeneous surfactant concentration gives rise to a non-homogeneous interfacial tension which in turn causes the tensile stress imbalance on the interface, pulling the fluid from the regions of maximum \( \Gamma \) towards the regions of minimum \( \Gamma \). This Marangoni induced slip velocity is shown by red (lower) arrows in Fig. 7. The slip of a surfactant laden interface is the sum of the slip of a clean interface and the Marangoni induced slip. As this Marangoni induced slip is directed oppositely to the clean interface’s slip at most of the locations, the slip of a surfactant laden interface is less than the slip of a clean interface (see blue dotted line in Fig. 7 for the slip of a surfactant laden interface).

![Graph showing Marangoni stress and leading order interface slip with the Marangoni number Ma for a fixed surface Péclet number Pe_s = 1.](image)

FIG. 8: (Color Online) The variation of (a) the Marangoni stress and (b) the leading order interface slip with the Marangoni number \( Ma \) for a fixed surface Péclet number \( Pe_s = 1 \).

Here \( \epsilon = 0.1 \) is used to calculate the Marangoni stresses while the values of other parameters \( \lambda, h, a, \) and \( d \) are kept the same as those of Fig. 3.

With an increase in \( Ma \), the Marangoni stresses increase (see Fig. 8a) which in turn reduce the slip of an interface (see Fig. 8b). At large \( Ma \) like \( Ma = 100 \), with an increase in \( Pe_s \) the advective transport of surfactant increases in comparison to its diffusive transport; this increases the gradients in the surfactant concentration (see Fig. 9a) or Marangoni stresses...
FIG. 9: (Color Online) The variation of (a) the surfactant concentration and (b) the leading order interface slip with the surface Péclet number $Pe_s$ for a fixed Marangoni number $Ma = 100$. Here $\epsilon = 0.1$ is used to calculate surfactant concentration while the values of other parameters $\lambda$, $h$, $a$, and $d$ are kept the same as those of Fig. 3.

which in turn reduce the interface slip (see Fig. 9b).

We note that the Marangoni stresses increase with an increase in $Pe_s$ at any fixed $Ma$, not just at large $Ma$. But at small $Ma$, with an increase in $Pe_s$, the increasing Marangoni stresses reduce the interface slip during the initial increase of $Pe_s$ (see Fig. 10a) while they increase the interface slip during the latter increase of $Pe_s$ (see Fig. 10b). This is because the Marangoni induced slip is directed opposite to (resp. along) the slip of a relatively clean interface, especially in the regions where the absolute value of a relatively clean interface’s slip is maximum, during the initial (resp. latter) increase in $Pe_s$. Compare the red (lower) arrows with blue (upper) arrows in Fig. 10 to understand this observation. Here relatively clean interface is an interface with lower $Pe_s$.


FIG. 10: (Color Online) Comparison of the interface slip of surfactant laden interfaces with different $Pe_s$ but with a fixed $Ma = 1$. In each plot, the interface slip at low and high $Pe_s$ are denoted, respectively, by blue solid and blue dotted lines. Also plotted is the change in the surfactant concentration at high $Pe_s$ in comparison to that at low $Pe_s$ (red dash-dotted line). The dashed lines are just for reference. In subfigure (a) the low and high $Pe_s$ are 0.1 and 1 while in subfigure (b) they are 1 and 10, respectively. The blue (upper) and red (lower) arrows denote, respectively, the vector field of surfactant laden interface’s slip at low $Pe_s$ and the direction of Marangoni induced slip as $Pe_s$ is increased from its low to high value. The axis for the interface slip is on the left while that for the change in the surfactant concentration is on the right. The values of the other parameters $h$, $\lambda$, $a$, and $d$ are kept the same as those of Fig. 3.


[52] A. Sokolov and I. S. Aranson, “Reduction of Viscosity in Suspension of Swimming Bacteria,”


