Effective Viscosity of Microswimmer Suspensions

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The measurement of a quantitative and macroscopic parameter to estimate the global motility of a large population of swimming biological cells is a challenge. Experiments on the rheology of active suspensions have been performed. Effective viscosity of sheared suspensions of live unicellular motile microalgae (Chlamydomonas Reinhardtii) is far greater than for suspensions containing the same volume fraction of dead cells. In addition, suspensions show shear thinning behavior. We relate these macroscopic measurements to the orientation of individual swimming cells under flow and discuss our results in the light of several existing models.

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In nature, organisms that can propel themselves in a fluid medium are ubiquitous. While larger organisms, such as fish, use inertia in their motion, microorganisms like spermatozoa, microalgae or bacteria, move at low Reynolds number, where viscous forces dominate over the effects of inertia. A recent and currently unresolved issue involves understanding the hydrodynamics associated with the individual or collective motion of microswimmers through their fluid-mediated interactions [1]. Microswimmer suspensions have been shown to lead to complex dynamics such as the so-called weak turbulence or bioconvection phenomenon [2–4]. Such active suspensions are made of self-propelled particles that create a force multipole either at the front of the body, in which case they are called pullers, or at the back, in which case, they are pushers. The flow induced by the force multipole is responsible for hydrodynamic interactions which are expected to have a dramatic effect on dynamics and particularly on the rheology of the suspension.

Recently, theoretical efforts have been made to model the effective viscosity of active suspensions. Stokesian dynamic simulations by Ishikawa and Pedley [5] show a difference in effective viscosity for suspensions of swimming spherical “squirmers” in a gravity field. Haines et al. [6] analytically showed that swimming results in a change in viscosity if the orientation distribution of swimmers is assumed to be anisotropic. More recently, Saintillan [7] used a simple kinetic model to study the rheology of a dilute suspension of self-propelled particles in a shear flow. He showed that suspensions of pullers exhibit increased effective viscosity compared to passive suspensions, while pusher suspensions exhibit a significant decrease in viscosity due to the motile activity; these results are consistent with previous predictions of Hatwalne et al. [8] who used coarse-grained active hydrodynamics to predict the rheology of active suspensions. Depending on swimmer types (puller or pusher), cell shape (spherical or ellipsoidal), and locomotion mechanisms, models can lead to very diverse results in terms of rheology. In order to better understand the effects of motility on viscosity, experimental work addressing the rheology of microswimmer suspensions is necessary.

Very recently, Sokolov and Aranson [9] measured the microrheology of suspensions of pushertype bacteria. They found, as predicted by [7,8], that the effective viscosity of such active suspensions decreases in comparison to passive particles at the same volume fraction. To our knowledge, no previous experimental data exist for pullertype microswimmer suspensions for which an increase of effective viscosity is predicted [7,8]. This Letter presents the first direct experimental macroscopic measurement of the effective viscosity of puller-type micro-swimmer suspensions: Chlamydomonas Reinhardtii (CR), a 10 μm motile unicellular alga. Rheological measurements performed on this system show a clear increase in effective viscosity when compared to a dead cell suspension. Shear thinning behavior is also measured. Based on models elaborated for dilute suspensions, we provide an interpretation of our results, invoking the anisotropic distribution of live cell orientations within the shear flow. We then discuss two hypotheses that could account for the origin of this anisotropy: a gravity torque or an effective elongated aspect ratio due to flagella beating.

CR microalgae [10] is a genus of green alga. It is a biflagellated unicellular organism. Chlamydomonas is used as a model organism for molecular biology, especially for studies of flagella motion, chloroplast dynamics, biogenesis and genetics. They are spheroidal in shape with two anterior flagella. Their back-and-forth movement produces a jerky breast stroke with a mean speed of $V \approx 40 \, \mu\text{m/s}$ in a waterlike viscous medium. Since the cell radius is $R \approx 5 \, \mu\text{m}$, Brownian motion is negligible. The swimming direction of the cells can be directed by stimulus gradients: a phenomenon known as taxis, such as chemo-taxis, rheotaxis, or phototaxis. Gradients are not used in our experiments in order to avoid any external tropism on the motility. Wild-type strains were obtained from the IBPC lab in Paris [11]. Synchronous cultures of CR were grown in a tri-acetate phosphate medium (TAP) using a 14 (10) hr light (dark) cycle at 25°C. Cultures were
typically grown for two days under fluorescent lighting before cells were harvested for experiments. These cultures were concentrated up to a typical volume fraction of 20 to 30\% by centrifuging for 20 minutes at 900g and resuspending them in a fresh culture medium to achieve the volume fractions required. Volume fractions were measured using hematocrit capillaries and a Neubauer counting chamber.

We started by characterizing cell motility. Microscopy imaging of chlamydomonas suspensions was carried out on an Olympus inverted microscope coupled with a Sensicam camera used at frame rates up to 30 Hz. Chambers made of glass and 240 \, \mu m spacers were coated with bovine serum albumine in order to reduce cell adhesion. Cells were imaged in bright field using a sensicam camera used at frame rates up to 30 Hz.

Two dimension trajectories of cells were recorded, typically for a few tens of seconds. Particle tracking was performed using interactive data language (IDL) [12]. The cell trajectories observed are correctly modeled by a persistent random walk [Fig. 1(c)]. Short term correlations in the direction of movement are observed: the swimmer then presents an almost fixed direction during a characteristic time. Hence, the mean square displacement of cells is well described by a random walk behavior with a persistence length \( \ell_p \) of approximately ten cell diameters: \( \langle x^2 + y^2 \rangle = \ell_p^2/4t/\nu - 0.5\ell_p^2[1 - \exp(-2t/\nu)] \) where \( \langle \rangle \) represents an ensemble average over more than a thousand independent measurements [12]. For \( t \ll \nu \), \( \langle x^2 + y^2 \rangle \sim V^2t \) with the ballistic velocity \( V = \ell_p/\nu \) and for longer times a random walk is observed, \( \langle x^2 + y^2 \rangle \sim 4Dt \) with \( D = \ell_p^2/4t/\nu \).

The algae velocity dependency on viscous drag was measured by adding a small amount of short chain dextran (molecular weight of 20,000) to the culture medium. This allowed variation of the viscosity of the medium between 1 and 2.5 mPa\,s. We checked that swimming velocity is inversely proportional to bath viscosity \( \eta \) (Fig. 1). This suggests that the stall force (the force needed to stop the swimming cell), which is proportional to the product \( \eta_0RV \), remains constant [Fig. 1(c)].

Let us now turn to the actual viscosity measurements. The effective viscosity of algae suspensions is measured on a Bohlin Gemini 150 rheometer equipped with cone-plate geometry (cone angle = 20°, diameter = 60 mm). Steady shear measurements were made at \( T = 20 \, ^\circ C \). If a shear stress \( \sigma \) is imposed and the associated shear rate \( \dot{\gamma} \) measured, effective viscosity is \( \eta_{\text{eff}} = \sigma/\dot{\gamma} \). Samples of different volume fractions were prepared and the effective viscosity measured for both live and dead cells. The cells were checked for motility after the rheometric measurement.

The effective viscosity \( \eta_{\text{eff}} \) of active suspensions is found to decrease with the shear rate [Fig. 2(a)] [16]. A clear global decrease of \( \eta_{\text{eff}} \) is observed from the maximum measured value at \( \dot{\gamma} = 4 \, s^{-1} \). As discussed below, it reveals competition between the shear rate and cells orientation in the flow. We then investigated the dependence of effective viscosity on the volume fraction of the suspension. To do so, viscosity was measured at a given shear rate of 5 \, s^{-1}, which is sufficiently high for rheometer resolution but low enough to allow viscosity to be affected by motility.

Figure 2(b) shows the relative viscosity \( \eta_{\text{eff}}/\eta_0 \) of live and dead cell suspensions as a function of the volume fraction. In both cases, viscosity is an increasing function of the volume fraction as it is for passive beads. Remarkably, the effective viscosity of swimming cell suspensions is quantitatively larger than the viscosity of dead cell suspensions (up to a factor of 2 for a 15\% volume fraction).

![image](image.png)

**FIG. 1** (color online). (a) Two dimensional trajectories of 50 swimming cells. Trajectory duration is 20 seconds. Trajectories starting positions have been all shifted to the origin. (b) Measured mean square displacement (MSD) of cells as a function of time. Averages have been extracted from 9 Hz sequences of about 700 cells trajectories. MSD fits well to a persistent random walk process with a correlation time \( \tau_c = 3.5 \pm 0.1 \, s \) and a coefficient \( D = 995 \pm 20 \, \mu m^2 \, s^{-1} \) (see text). Inset: Log-Log plot. (c) Measured mean velocity \( V \) of cells as a function of the inverse viscosity of the medium. The line represents a linear fit. Inset: Viscous drag force should be proportional to the product of the medium viscosity \( \eta_0 \), the mean radius \( R \), and the velocity \( V \) of the cells. This product \( \eta_0RV \) is shown to be independent of the medium viscosity. Error bars are smaller than the symbols.
The effective viscosity $\eta_{\text{eff}}$ of a suspension of passive spherical particles in a solvent of viscosity $\eta_0$ depends on its volume fraction $\phi$. Krieger and Dougherty’s semiempirical law [17] is shown to provide a reliable description of the measurements:

$$\eta_{\text{eff}} = \eta_0 \left(1 - \frac{\phi}{\phi_m}\right)^{-\alpha \phi_m}. \quad (1)$$

Here, $\phi_m$ is the maximal packing volume fraction. For a dilute regime, where $\phi \ll 1$, Eq. (1) reduces to $\eta_{\text{eff}} \approx \eta_0 (1 + \alpha \phi)$, where $\alpha$ is known as Einstein’s intrinsic viscosity ($\alpha = 2.5$ for passive and spherical particles) [18].

To quantify the effect of motility on the effective viscosity of cell suspensions, we fitted the measurements using Eq. (1). We set the maximal packing volume fraction to 0.62 and left $\alpha$ as a free parameter. This resulted in a large difference between active and passive suspensions. We found $\alpha = 2.5 \pm 0.1$ in the case of dead cells, which clearly behave as passive particles. However, in the case of swimming cell suspensions, we obtained $\alpha = 4.5 \pm 0.2$. Swimming cells induce a quantitative increase in the effective viscosity of the suspension.

In order to understand this increase in effective viscosity, we conducted complementary experiments which consisted of imaging the cells while subjected to a shear flow. Dead cells show a regular rotation at an angular velocity close to $\gamma/2$, as would passive spherical particles. We observed that swimming cells behave very differently: they resist the flow rotation for most of the time and eventually flip very rapidly. Figures 3(a) and 3(b) show picture sequences extracted from a fast-image film. High frequency acquisition (500 Hz) allows us to determine whether or not an alga is swimming by looking at the beating of flagella. In the case of a dead cell, flagella only move because of thermal agitation, whereas flagella of swimming cells beat at about 50 Hz. Figure 3 shows 50 Hz time sequences of cells subjected to a 10 s$^{-1}$ shear rate. The swimming cell spends about two thirds of the period in the $xOy$ plane and flips during one third of the period whereas the dead cell rotates at a constant velocity close to $\gamma/2$.

**Discussion.**—A puzzling question arises from our experiments: why should live cells and dead cells with the same aspect ratio and the same volume fraction respond so differently in a flow? In order to answer such a question, we propose some hypotheses based on existing models which might explain the increase of effective viscosity according to individual dynamics of live cells. As shown on Fig. 1(b), the typical time $t_0$ after which a chlamydomonas changes its direction of motion is about 3 to 4 seconds. This is much longer than the typical time associated with the shear flow: $t_f = \gamma^{-1} \approx 0.2$ s. This means that live algae resist the flow rotation [Fig. 3(a)] during approximately 70% of the time $t_0$, by being aligned in the flow. The fast flip between

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**FIG. 2.** (a) Effective viscosity of chlamydomonas suspensions as a function of shear rate. Data are shown for different volume fractions of the suspension, and star symbols represent the viscosity $\eta_0$ of the culture medium. The lines are shown for ease of viewing only. (b) Relative viscosity of microswimmer suspensions (measured at shear rate = 5 s$^{-1}$) as a function of volume fraction. Solid symbols represent live cell data and crossed symbols represent dead cell data. Measurements are fitted to Eq. (1) using $\phi_m = 0.62$ and $\alpha = 2.5$ (dead cells) and 4.5 (swimming cells).

**FIG. 3.** (a) Dead cell in a 10 s$^{-1}$ shear flow experiencing tumbling around the x axis. (b) Live chlamydomonas swimming in the same shear flow. Arrows indicate cells flips. Time between pictures is 20 ms. Cell diameter is about 10 micrometers. (c) Schematic view of the flow cell, where $Oz$ is the direction of observation, $yOz$ is the shear plane.
Brownian contribution is replaced here by hydrodynamic diffusivity due to hydrodynamic interactions between swimming CR.

In this Letter, we have experimentally shown that active suspensions of puller-type microswimmers present a dramatic increase in effective viscosity. We have correlated these macroscopic rheological measurements to the individual dynamics of cells under shear flow: swimming cells tend to resist the flow rotation unlike dead cells. More experiments are needed to explain the origin of this phenomenon: a gravity torque exerted on the cell or an effective large aspect ratio due to flagella beating.

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[16] Note that viscosity seems to increase slightly at higher shear rates.