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Turbulence of swarming sperm

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Collective motion of self-sustained swarming flows has recently provided examples of small-scale turbulence arising where viscous effects are dominant. We report the first observation of universal enstrophy cascade in concentrated swarming sperm consistent with a body of evidence built from various independent measurements. We found a well-defined $k^{-3}$ power-law decay of a velocity field power spectrum and relative dispersion of small beads consistent with theoretical predictions in 2D turbulence. Concentrated living sperm displays long-range, correlated whirlpool structures of a size that provides an integral scale of turbulence. We propose a consistent explanation for this quasi-2D turbulence based on self-structured laminated flow forced by steric interactions and alignment, a state of active matter that we call “swarming liquid crystal.” We develop scaling arguments consistent with this interpretation.

I. INTRODUCTION

Large-scale coherent dynamics in sessile drops of bacterial flows [1] or sperm [2,3] were reported many years ago. For spermatozoa confined between two plates [2], coherent vortex rings composed of as few as 10 individuals were observed to form. In the case of sperm, collective motility analysis based upon the observed turbidity in the vicinity of the sessile drop contact line has been used as a reliable index for male fertility scoring [3] in livestock. At a more fundamental level, there has been recent interest in the long-range spatiotemporal dynamics in concentrated suspensions of swimmers [2,4–11].

Collective motion of swarming flows is an emerging topic that involves challenging issues related to its continuum mesoscopic description, especially at high densities [12–21].

Recent observations in bacterial suspensions have shown that highly concentrated swarming flows display “turbulent-like” [4,8,22] behavior without inertia. While the use of the term “turbulent” might seem inappropriate in the context of inertialess flow (“chaotic” or “intermittent” are arguably more appropriate), it has been used in the present article due to its use in the majority of previous studies in the field [14,8,21].

II. EXPERIMENTAL METHODS, CALIBRATION, AND VALIDATIONS

A. Ram semen characteristics

Ram semen is highly concentrated with a volume fraction $\phi \approx 50\%$. This volume fraction is calculated based on the concentration and the ram spermatozoon dimensions. The ram spermatozoon length is approximatively 60 $\mu$m with 50 $\mu$m of it being the flagellum. Its head has an ellipsoidal shape, flat at the bottom and round at the top with a thickness of 3 $\mu$m, a 10 $\mu$m length, and 5 $\mu$m width.

B. Camera recording and microscope setup

For gray-level images, we use a high-speed camera PCO Dimax S4 (Photon Lines, France) mounted on a phase-contrast microscope (Nikon, Japan). For fluorescent images, we use a pco.edge camera (Photon Lines, France) mounted on a fluorescent microscope (Olympus BH2, Japan). Both microscopes are equipped with temperature control systems and with $\times 4$ magnification objectives. Recorded images are 2000 $\times$ 2000 pixels and have 12 bits of gray level. The frame rate is 50–100 Hz. Phase contrast is performed by a transparent phase plate in a phase-shift ring so as to enhance sample contrast. For the particle image velocity (PIV) measurement the pixel size is 1.08 $\mu$m $\times$ 1.08 $\mu$m, which gives recorded windows of 2 mm $\times$ 2 mm. The pixel size is 1.5 $\mu$m $\times$ 1.5 $\mu$m for particle tracking velocity (PTV) acquisitions giving 1.628 mm $\times$ 1.628 mm recorded windows. The ram sperm samples were confined in Leja (IMV) chambers of 100, 150, and 200 $\mu$m depth.

C. Particle image velocity (PIV) reconstruction

The standard image correlation PIV algorithm DavisTM has been used on the gray-level phase-contrast images. The projection of the Eulerian velocity field in the focus plane is reconstructed on a two-dimensional (2D) grid using a cross-correlation PIV algorithm. The algorithm estimates the most probable displacement of grid-centered small windows (identification window) between two successive frames.
The fluctuations velocity have a Gaussian similar fit. To improve the accuracy and reduce peak locking, iterative calculations are done to take into account the deformation of the identification window in order to increase the accuracy and reduce peak locking.

The spatial resolution is dictated by the size of this window, which in our case is 16 pixels × 16 pixels, which corresponds to 17 μm × 17 μm. This size is small enough to resolve spatial structures on the order of a few spermatozoa. The velocity field is obtained on a grid of size 126 × 126.

D. Particle detection, tracking, and PTV

Fluorescent microparticles (Spherotech, FH-2056-2) were mixed with semen at a final concentration of 3.9 × 10^6 beads/ml. The fluorescent microparticles have high-intensity Nile red fluorescence (excitation/emission 530/605 nm), size 1.7–2.2 μm and density 1.05 g/cm^3. They are spherical and uniform in shape.

Particle detection algorithms have been coded in house using MATLAB®. Individual beads are tracked along time from PTV of the fluorescent images as illustrated in Fig. 1.

1. Detection

Individual particles are first detected using a particle mask convolution analysis [23]. A particle template which consists of a 2D Gaussian distribution G having an isotropic standard deviation of σ = 5 pixels is scanned over the entire image in order to detect the peaks of image intensity which correspond to the particles. We compute the normalized convolution

\[ C(\Delta_i, \Delta_j) = \frac{\Sigma_{i,j} I(i,j) G(i + \Delta_i, j + \Delta_j)}{\sqrt{\Sigma_{i,j} I^2(i,j) \Sigma_{i,j} G^2(i,j)}}\]

where \( G(i) = \exp(-i^2/2\sigma^2)/\sqrt{2\pi}\sigma \) is the normalized Gaussian, and \( I(i,j) \) is the original image gray-level intensity. A 0.45 threshold is then applied to \( C(\Delta_i, \Delta_j) \) in order to provide an isolated binary island for each event detection. The center of the detected particle is then computed by performing a 2D Gaussian regression [24] into the convolution field \( C(\Delta_i, \Delta_j) \) to provide a subpixel accuracy of the position \( x_p^n \) of particle \( p \) of image \( n \).

2. Tracking

Since the mean displacement of the particle images is small compared to the mean inter-particle distance, the matching pairs between two successive frames can be found by pairing each particle in the first frame with its closest neighbor in the second frame. For each detected position \( x_p^n \) of image \( n \) we find any particle \( p' \) of image \( n + 1 \), whose position \( x_{p'}^{n+1} \) lies within a radius of \( R = 10 \) pixels, i.e., \( |x_{p'}^{n+1} - x_p^n| < R \). We then select the closest event for the pairing.

3. Particle tracking velocity (PTV)

For a particle \( p \) whose position, \( x_p^n \), in image \( n \), moves to position \( x_{p'}^{n+1} \) in the subsequent image \( n + 1 \), its velocity is \( \mathbf{v}_p = (x_{p'}^{n+1} - x_p^n)/\Delta t \) where \( \Delta t \) is the time between frames. By considering in each image \( n \) the relative distances \( D^2 = |x_{p'}^{n+1} - x_p^n|^2 \) and relative velocities \( V^2 = |\mathbf{v}_{p'} - \mathbf{v}_p|^2 \) of each \( (p, p') \) particle pair, we can build a very robust statistics of the relationship between \( D \) and \( V \). These data are shown in Fig. 10 below.

4. Comparison of PIV and PTV

Finally, we investigate whether phase-contrast gray-level texture PIV gives a sensible measurement of the real flow field from a direct comparison with PTV. Since small particles beads are perfect inertial Lagrangian tracers of the flow, they offer the most reliable estimate of the local flow velocity.

In addition to the fact that PTV and PIV fluctuations have similar PDFs as shown in Fig. 2, we directly investigate for each event, the relative error between PIV and PTV. On the same sequence of images of pure semen seeded with Lagrangian PMMA-Rhodamine B-particles of 1–35 μm diameter, a comparison was performed by interpolating the PIV to the beads particle locations. We found the relative error between PIV and PTV to systematically drop from 50% for particle diameters of 10–15 μm to 25% for particle diameters of 30–35 μm as shown in Fig. 3. This is consistent both with the optical averaging of the velocity field within the microscope.
field depth $\delta = 25 \pm 10 \mu m$, as well as the PIV window size and the resulting interpolation error. This observation provides strong evidence to support that the velocity field measured from the image gray-level texture correlation is indeed very close to the depth-averaged true velocity of the flow over the microscope field depth.

E. Sample collection and scoring

Ejaculates from adult Lacaune rams were collected with an artificial vagina [25]. All ejaculates used in this study scored over three for Massal mobility (MM), had a thick and creamy appearance, and were free of blood and urine contamination. Wave motion of semen was observed in a 10 $\mu l$ drop of raw semen on the warming plate of a BH-2 Olympus microscope with a 4× phase-contrast objective. The MM score was performed based on the Herman definition (13). The MM score was refined from its original definition to subjectively assess the rotation speed of dark waves (5 being the maximal speed). The sperm concentration was also assessed by measurement of absorbance using spectrophotometry. Such measurements provide a precalibrated estimation of the cell concentration inside a cylindrical tube (the calibration has been validated against other techniques such as manual counting). From the concentration, and knowing the volume of each individual, we infer the volume fraction.

F. Rheological behavior of pure and dilute semen

We perform a rheological analysis of pure and dilute fresh samples. Between 1 and 4 h after collection, samples maintained at 15 °C were placed into the lower plate of a cone-plate rheometer (Mars III Fischer Scientific). All dilutions were performed with isotonic NaCl. We mainly use a 1° angle, 35 mm diameter cone, since the quantity of fluid involved is obviously limited for semen collections (typically 1.5 ml). Nevertheless, some results have been confirmed and validated with a 1° angle, 60 mm cone diameter. Each measurement was performed during a steady state of the torque. The time lag associated with its stabilization is 180 s, and for each measurement, the integration time for the torque evaluation is 30 s. Each experiment has been performed at 37° C preventing the evaporation of the fluid during the measurement process (about 30 min) by confining the cone/plate setup into a controlled saturated water vapor enclosure. The shear rate $\dot{\gamma}$ has been varied from $10^{-3}$ to $2 \times 10^2$ s$^{-1}$.

Figure 4 illustrates the semen viscosity, which displays a typical shear-thinning power-law behavior. Since a very similar curve is observed for seminal plasma, it can be deduced that this shear-thinning behavior is typical of the plasma protein suspensions and weakly dependent on the presence of spermatozoa. This result is consistent with previous rheological analysis for other species [26,27]. This is also confirmed from the curve obtained for diluted plasma where we obtain the viscosity of water at large shear rate and the same power-law behavior at lower shear rates. Spermatozoon is a pusher-like swimmer, the presence of which decreases the semen viscosity [28–30], unlike to puller [31]. Due to the very reduced lifetime of spermatozoon activity, we were unable to perform viscosity measurements on active semen. All the presented rheology measurements were performed on weakly active or inactive spermatozoa suspensions. As found for suspensions of B. subtilis (pusher-like swimmer) [29], our results show that for nonactive cells the viscosity increases with concentration.

III. EXPERIMENTAL RESULTS

We experimentally investigate the statistical turbulence of semen with various complementary indicators.

After collection, a pure, fresh ram semen sample was introduced into a confined transparent glass chamber of 1 cm in length in each horizontal direction and a few hundred microns ($H = 100,150,200 \mu m$) in thickness. $H$ is much larger than cell size since the ram sperm head size is $a \approx 10 \mu m$. Direct
light phase-contrast reveals visible structures of many scales as shown in Figs. 5(a) and 5(b) (see movies S1 and S2 in Ref. [32]). The observed whirlpools in Figs. 5(a) and 5(b) are weakly sensitive to the cell gap $H$, but insensitive to the orientation of the glass chamber with respect to gravity. This dependence is quantitatively analyzed in Sec. IIIA. Hence in the following, we will ignore the influence of gravity, and the glass chamber is set up horizontally in the $(x,y)$ plane. The collective structures of the resulting swarming flow can be quantitatively analyzed using PIV measurements (see Sec. II D).

### A. Integral scale

An example of 2D flow field $\mathbf{u}(x) = (u(x), v(x))$ at a horizontal position $x \equiv (x,y)$ is provided in Fig. 5(c). Considering $\mathbf{u}'(x) = \mathbf{u}(x) - \langle \mathbf{u} \rangle$ the velocity fluctuations by subtraction of the mean flow $\langle \mathbf{u}(x) \rangle$, Fig. 5(d) shows the vertical component of the vorticity $\omega'(x) = \nabla \times \mathbf{u}' = \partial_x v'(x) - \partial_y u'(x)$ normalized by the enstrophy $\Omega' = \langle |\omega'|^2 \rangle/2$ where $\langle \cdot \rangle \equiv \int_S \cdot dS/S$ and $S$ is the total horizontal surface. We see that the largest flow structure is typically smaller than the recorded windows but much larger than individual cells (see movie S3 in Ref. [32]).

In order to further analyze this integral length scale the spatiotemporal correlation functions were evaluated. This provides insight into the structure cascade and its dynamics [7–11]. The energy-to-enstrophy ratio defines an integral length scale [8]

$$L^* = \sqrt{\frac{E_i}{\Omega'}};$$  \hspace{1cm} (1)

which is shown in Fig. 7(b). The relationship between $\Omega'$ and $E_i$ displays a linear trend, leading to a well-defined integral length scale $L^*$ that is independent of the initial energy of the active flow. As expected, $L^*$ is close to the observed maximum whirlpool sizes. Table I shows $L^*$ for different values of $H$. A

<table>
<thead>
<tr>
<th>$N$</th>
<th>$H$ (μm)</th>
<th>$L^*$ (μm)</th>
<th>$C$ (10^9 cells/ml)</th>
<th>$\phi$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>100</td>
<td>126 ± 16</td>
<td>5.2 ± 1.1</td>
<td>50 ± 11</td>
</tr>
<tr>
<td>1</td>
<td>150</td>
<td>133</td>
<td>5.4</td>
<td>52</td>
</tr>
<tr>
<td>1</td>
<td>200</td>
<td>146</td>
<td>5.4</td>
<td>52</td>
</tr>
</tbody>
</table>

TABLE I. The integral length scale $L^*$ for different gap spans of the glass chambers $H$. $N$ is the sample number and $\phi$ the evaluated volume fraction.
We then use the major axis of the ellipse as a weak dependence is found, the significance of which will be discussed in Sec. IV.

We wish to compare this integral length scale $L^*$ with the whirlpool structures present in Figs. 5(a) and 5(b). For this we need another quantification independent of the flow field, but related with image texture, so as to detect and quantify a typical length scale for the whirlpools. For this, gray-level phase-contrast images are thresholded at 70% of the gray-level mode. The resulting binary images are then dilated (using dilation morphological operator) with $5 \times 5$ square structural elements. The resulting images are filtered with a median filter whose size is between 31 and 41 pixels and again binarized with the same threshold. A typical example is illustrated in Fig. 6(b), which can be compared with the original gray-level display in Fig. 6(a). Connected components associated with the whirlpools are then detected. An ellipse is extracted, the properties of which preserve the second moments of the connected component pixel region [cf. Fig. 6(b)]. The center of each ellipse corresponds to the barycenter of the connected component. We then use the major axis of the ellipse as a typical length scale $l^*$ of the whirlpools and compare it with the integral scale $L^*$ defined in Sec. III A. $L^*$ and $l^*$ are compared in Table II where one sees they have similar values. An average 6% difference is found between both estimation of the integral length scale. This is an interesting observation since one is obtained from kinematics properties of the flow, and the other one results from the analysis and segmentation of gray-level texture.

### Table II. The integral length scale $L^*$ and the whirlpools characteristic length $l^*$ for different values of $H$.

<table>
<thead>
<tr>
<th>$H$ (μm)</th>
<th>$l^*$ (μm)</th>
<th>$L^*$ (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>119</td>
<td>109</td>
</tr>
<tr>
<td>100</td>
<td>125</td>
<td>120</td>
</tr>
<tr>
<td>100</td>
<td>112</td>
<td>108</td>
</tr>
<tr>
<td>100</td>
<td>129</td>
<td>127</td>
</tr>
<tr>
<td>100</td>
<td>150</td>
<td>137</td>
</tr>
<tr>
<td>100</td>
<td>130</td>
<td>150</td>
</tr>
<tr>
<td>150</td>
<td>175</td>
<td>133</td>
</tr>
<tr>
<td>200</td>
<td>183</td>
<td>146</td>
</tr>
</tbody>
</table>

B. Two-point correlation of the flow field

The PIV and PTV statistics of the velocity fluctuations, $u^*(x) = u(x) - \langle u \rangle$, is found to be isotropic and close to Gaussian (see Fig. 2) as in Refs. [8,22]. The two-point correlation between point $x$ and point $x + \xi$ of $u^*(x)$ is analyzed along the direction parallel to $\xi$, $u'_i = u^i / |\xi|$. The correlation function was found to be isotropic. Hence we consider only its dependence with distance $|\xi|$, as illustrated in Fig. 7(a). Considering the dimensionless distance $|\xi|/L^*$, provides a nice collapse of the correlation functions in Fig. 7(c). We see that the integral length scale $L^*$ also defines the typical correlation length that is independent of the initial energy of the active fluid [see Fig. 7(b)]. It is interesting to mention that an integral time scale $\tau^*$ can be also extracted from our measurements, as illustrated in Fig. 7(e). A nice collapse of the time-correlation profile is also obtained. This integral time scale displays a linear trend with the typical mesoscale time scale built upon the enstrophy $\Omega^* = \langle |\omega|^2/2 \rangle$, $\tau^* \sim 1/\sqrt{\Omega^*}$ as shown in Fig. 7(d).

We now analyze the “internal” length scale of the velocity field from the swimmer size $a$ to the integral scale $L^*$. For this purpose, we consider the energy spectrum, which is the signature of the cascade in the kinetic energy. From the observation of an isotropic velocity Fourier transform $\tilde{u}(\mathbf{k})$ in the wavelength $\mathbf{k}$ plane, we evaluate the averaged azimuthal kinetic energy spectrum in Fig. 8.

This spectrum displays three distinct regimes: at small $k$, it saturates to a white noise plateau, whereas, above a definite critical wavelength $k_c$, it displays a clear power law decay. At large $k$, it saturates again when $k$ reaches a second critical value $k_{\ast}$. The first regime corresponds to scales where the largest vortex reaches the size of the integral scale $L^*$, since $k_c = 2\pi/L^*$. At this stage, the fluctuations are uncorrelated in space and form a random set of coherent structures observed in Figs. 5(a) and 5(b) as already found by Refs. [4,5,8,10,11]. The second regime displays a well-defined, self-similar scale cascade, consistent with the $k^{-3}$ power-law decay illustrated with dotted lines in Fig. 8. More precisely, from fitting each
individual experiment, we estimate the error bars on the power-law decay to be $-3 \pm 0.4648$. Such $k^{-3}$ power-law decay is indeed found in quasi-2D turbulent flows. Then the last white noise plateau regime for $k > k_a = 2\pi/a$ is consistent with size of the order of individual swimmers. These suggest that the velocity correlations build up by coarse graining from the individual swimmer scale to reach broader collective scales within the range $[a, L^*]$. For dilute samples, the structures persist at large scale but turn cloudy at small scale (see movie S4 [32]). This qualitative observation is confirmed by the computation of the integral scale $L^*$ obtained from relation (1), while a linear trend is indeed observed between enstrophy and energy (not shown) during the time decay of the sample motion. This integral scale $L^*$ does not change with the dilution as we found $L^* = 118 \mu m$ for pure, $L^* = 90 \mu m$ for $\times 2$ dilution, and $L^* = 114 \mu m$ for $\times 4$ dilution. Hence, volume

FIG. 7. (Color online) (a) Longitudinal correlation function of the fluctuation velocity field, $\rho(\xi) = \langle u'(x)u'(x+\xi) \rangle / \langle u'^2 \rangle$ obtained from PIV measurements. The different set of curves corresponds to different semen samples recorded within chambers whose depths are $H = 100 \mu m; H = 150 \mu m; H = 200 \mu m$. (b) The 2D fluctuating kinetic energy $E' = \langle |u'|^2 \rangle / 2$ is plotted versus the fluctuating enstrophy $\Omega' = \langle |\omega'|^2 \rangle / 2$ for six different experiments each of more than 2000 images. Each circular symbol represents an experiment in a 100 $\mu m$ chamber where error bars represent the enstrophy standard deviation on each hole sequence. The dotted line shows the linear trend associated with a constant ratio $E'/\Omega'$, the square root of which provides the integral length $L^*=\sqrt{E'/\Omega'}$. (c) The collapse of the curves when considering dimensionless distance $\xi/L^*$ along the horizontal axis. (d) The linear trend between integral time scale $\tau^*$ obtained from the first zero of the temporal correlation function $\rho(\tau) = \langle u'(t)u'(t+\tau) \rangle / \langle u'^2 \rangle$ and the mesoscopic time scale $\tilde{\tau}$ obtained from enstrophy. (e) Temporal correlation function $\rho(\tau)$ plotted versus the dimensionless time $\tau/\tau^*$, where $\tau^*$ defines the characteristic time obtained from the first zero of the temporal correlation function.
fraction hardly affects the integral scale. This is confirmed by the computation of the power spectrum provided in Fig. 9, where it can be seen that the normalized spectrum amplitude is increased at small wavelength and decreased at larger wavelengths. This is the signature of a decrease of small-scale velocity correlations and lower velocity at small scales. The effect of the dilution on the small-scale structures is also visible in the movies (see movie S4 [32]).

Finally, it is important to report that semen dilution drastically decreases the optical phase contrast of the suspension, so that texture correlation computed by PIV is not reliable or difficult to handle. In many cases, we have not been able to get reliable PIV measurements from many diluted samples. For dilution larger than x8 the number of false vectors in the resulting PIV field becomes larger than 10% so that velocity field can no longer reliably be estimated. Hence, for diluted samples, the statistics of the presented results are less substantial than for pure semen ones.

C. Quasi-2D properties of the flow

The superposition of normalized instantaneous vorticity and streamlines, presented in Fig. 5(d), are in good agreement, qualitatively suggesting that the flow field is quasi-2D. This fundamental feature of the flow is more deeply investigated with two complementary indicators: (1) the ratio of the in-plane 2D kinetic energy to the out-of-plane kinetic energy and (2) the ratio between Lagrangian tracers’ curvilinear path length in the observation plane to the microscope field depth.

(1) We first evaluate the mean 2D kinetic energy \( E_{2D} = \langle |\mathbf{u}|^2/2 \rangle = \langle (u^2 + v^2)/2 \rangle \) and examine the time over which the semen energy decays. A slow decay is observed over a few minutes. During recording sequences of 40 s, the kinetic energy typically decreases by about 40%. Computing the 2D divergence of the flow, we evaluate the out-of-plane fluctuating kinetic energy. The divergence of the flow field can indeed provide an estimate of the vertical velocity at the edges of the microscope depth field. Since locally fluid incompressibility holds, the divergence of the 2D flow components is

\[
\text{div} \, \mathbf{u}' = -\partial_z w',
\]

where \( w' \) is the vertical component of the fluctuating velocity field. Integrating (2) in \( z \) from \( z = -\delta/2 \) to \( z = +\delta/2 \), where \( \delta \) is the microscope depth field, and assuming that the actual measured in-plane velocity is the depth-average velocity leads to

\[
\text{div} \int_{-\delta/2}^{\delta/2} \mathbf{u}' \, dz = -\delta w',
\]

where \( \delta w' = w'(+\delta/2) - w'(-\delta/2) \) is the out-of-plane fluctuation velocity component differences at the edges of the microscope field depth \( \delta \). Thus, by computing the divergence of the 2D PIV in-plane flow, we can estimate the fluctuating component of the vertical flow and the vertical fluctuating kinetic energy, \( E'_w = \langle |\delta w'|^2/2 \rangle \). Comparing \( E'_w \) to \( E_{2D} \) averaged for all time over 3000 images and six different experiments, we found that the ratio \( E'_w/E_{2D} = 8 \times 10^{-3} \). This is a clear indication that the out-of-plane to the in-plane velocity ratio \( |u'|/|\mathbf{u}'| \sim 0.0845 \sim 1/12 \) is typical of quasi-2D flows. (1) We seed the semen with highly diluted fluorescent particles (volume fraction equals 1.2 \times 10^{-5}). By tracking the particle positions within the flow (see Fig. 1 and Sec. II D) we evaluate their curvilinear path length in the horizontal plane \( \ell_{2D} \) and then compute its average value \( \bar{\ell}_{2D} \) for all particles denoted. We find \( \bar{\ell}_{2D} = 356 \pm 80 \, \mu m \), which is much larger than the estimated microscope field depth \( \delta = 25 \pm 10 \, \mu m \), as \( \delta/\bar{\ell}_{2D} \sim 0.0844 \sim 1/12 \). This is consistent with the previous estimate \( |u'|/|\mathbf{u}'| \sim 1/12. \)
FIG. 10. (Color online) Relative velocity $V$ of two particles versus their relative distance $D$ is plotted in bilogarithmic scales in order to challenge turbulent particle dispersion scaling predictions. One measurement is illustrated with data gathered from 3000 frames, with, in average, 575 detected particles per image pair. Symbols represent experiment on different males (♂: male no. 1; ♀: male no. 2; ♂: male no. 3), and solid black lines indicate a power-law fit whose slope is equal 1.05 consistent with $V \sim D$ scaling.

D. Separation law

Here we analyze the specific scaling of the particle tracer trajectory statistics. One interesting prediction in turbulent flows is related to the so-called separation law, also called relative dispersion [33], which is the relation between the average relative distance and the average relative velocity for each particle pair. This law for three-dimensional (3D) Kolmogorov turbulence differs from that from the enstrophy cascade scaling. For 3D Kolmogorov turbulence, the relative velocity $V$ [each particle pair $(p, p')$, $V^2 = |w_p - w_{p'}|^2$; see Sec. II D] for two particles in the flow scales as $V \sim D^{1/3}$ where $D$ is the relative distance [34,35]. In 2D turbulence associated with an enstrophy cascade scaling, the prediction becomes $V \sim D$ [33,36]. From PTV (Sec. II D), we evaluate the relative velocity $V$ of a very large number ($\approx 3 \times 10^5$) of particle pairs in order to build the histogram of the relative velocity versus relative distance $D$. By computing the average velocity $V$ over the interval $D, D + 8D$, we can test the scaling prediction for relative particle dispersion. Figure 10 shows that the linear prediction $V \sim D$ is consistent with our observations within more than one decade in distances. The quality of the experimental curve relies on the very large number of events: an average of 575 particles found correlated for each image pair (image pairs are needed to compute each particle velocity) and statistics are computed over 3000 images to give an excellent average. The scaling obtained in the log-log plot of Fig. 10 is highly reproducible from one sample to another (only three of them are illustrated). This very robust behavior is a reliable test for the enstrophy-cascade-like velocity spectrum, which is consistent with the spectrum obtained in Fig. 8.

IV. DISCUSSION

A. Scaling derivation

The development of continuum equations for concentrated suspensions is still an active field of research [12–17,22,37–40]. Here we provide simple insights for the leading asymptotic structure of a confined active suspension to examine whether the quasi-2D structure of the flow is imposed by the Hele-Shaw chamber or by some internal self-structured layering.

In the following, we show that self-structured layering gives a description consistent with our experimental findings, as a vertical length scale $a \ll H$ is shown to be a relevant length-scale. Since this length scale $a$ also equals the size of a spermatozoa head we show that layered sheets of horizontally aligned spermatozoa are slipping one over another and forming quasi-2D turbulent layers.

Obviously the standard 2D asymptotic reduction of Stokes flow leads to the Darcy-Brinkman equations [41]. But in the case of active fluid suspensions, the driving mechanism comes from a supplementary stress associated with the influence of active swimmers which drive the flow through dipolar forcing generates.

Fluid incompressibility associated with velocity field $(u, v, w) = (u, w)$ reads

$$\nabla \cdot (u, v, w) = \nabla \cdot u + \partial_x w = 0,$$

where $\nabla = (\partial_x, \partial_y, \partial_z)$. When considering an integral length scale $L^*$ for the typical in-plane variations along $(x, y)$, a typical swimming velocity of spermatozoa $u \sim v \sim U_2$, and a typical vertical length $a$, then incompressibility provides

$$w \sim a U_2 / L^* \sim \epsilon U_2,$$

and the definition of a small parameter $\epsilon = a / L^*$. The momentum balance equations which govern the motion of an active suspension on sufficiently large length and time scales at zero Reynolds number [37,38] are

$$\mu \Delta (u, v, w) - \nabla p = - \nabla \cdot \sigma^E = C_0 \nabla \cdot Q,$$

$$Q = \begin{pmatrix} p \mathbf{I} - \frac{1}{3} \end{pmatrix},$$

$$\sigma_0 = \frac{3 \phi \mu U_2 a^2}{4 \pi a^3},$$

where the extra-stress tensor $\sigma^E = - C_0 \nabla Q$ is composed of $C$, a positive constant for spermatozoa which are pushers, $Q$, the stresslet-nematic tensor averaged over the orientation of the swimmers, and $\sigma_0$, a dimensional active stress constant, $\sigma_0$ is composed of two lengths: $a$, associated with the spermatozoa head, and $\ell$, which is the flagellum length, and $\phi$, the volume fraction. The extra-stress tensor thus scale as $\sigma^E \sim \mu U_2 a^2 / a^3$. Using dimensionless length scale $L^* = \ell \approx 1/\epsilon(a, \epsilon, \epsilon, \ell, \ell)$, gradient $\nabla \equiv 1/a(\ell, \ell, \ell, \ell)$, and velocity scales $U_1(\ell, \ell, \ell, \ell)$, the leading order dimensionless form of (6) is

$$\frac{\mu U_2}{a^2} \left( \frac{\ell^2}{\ell^2 - \ell^2} - \partial \ell \right) \approx \frac{\mu U_2 a^2}{a^4} \left( \ell^2 - \ell^2 \right) \approx \frac{\mu U_2 a^2}{a^4} \left( \ell^2 - \ell^2 \right),$$

$$\frac{\mu U_2}{a^2} \left( \frac{\ell^2}{\ell^2 - \ell^2} - \partial \ell \right) \approx \frac{\mu U_2 a^2}{a^4} \left( \ell^2 - \ell^2 \right) \approx \frac{\mu U_2 a^2}{a^4} \left( \ell^2 - \ell^2 \right),$$

$$\epsilon \frac{\mu U_2}{a^2} \left( \frac{\ell^2}{\ell^2 - \ell^2} - \partial \ell \right) \approx \epsilon \frac{\mu U_2 a^2}{a^4} \left( \ell^2 - \ell^2 \right) \approx \epsilon \frac{\mu U_2 a^2}{a^4} \left( \ell^2 - \ell^2 \right).$$

The dimensionless pressure is chosen to balance the in-plane viscous forces, so at leading order $p \equiv (\mu U_2 L^* / a^2) \tilde{p}$, which is invariant along the gap span from the third momentum balance.
component, \( \partial_t \tilde{p} = 0 \). Balancing the viscous and extra-stress leading order contributions in each \((x,y)\) direction provides the same consistent scaling \( \epsilon \sim \frac{a^2}{\ell^2} = \frac{a}{\ell} \), so that one finds an estimate for the longitudinal length scale \( L^* \sim \frac{a}{\ell} \). One should also note that the dimensionless right-hand side of (7) implies that the horizontal orientation gradients \( \partial_x (p_x) \) are of same order of vertical one \( \partial_y (p_y) \). This indicates that the averaged orientation is mainly horizontal in the \((x,y)\) plane, i.e., \( (p_x) \sim (p_y) \sim O(1) \) and weakly vertical \( (p_z) \sim \epsilon \). One can also evaluate the resulting scaling for the in-plane kinetic energy \( E_{ij} \sim U_{\parallel}^2 \) and the enstrophy \( \Omega \sim (U_{\perp}/L^*)^2 \), so that \( L^* \) is also given by \( L^* \sim \sqrt{E_{\parallel}/\Omega} \), as evaluated in Fig. 7(b). This analysis is consistent with the following:

1. The ratio between the vertical and horizontal velocity \( \frac{|w|/|u|}{|w|/|U_\perp|} \sim a/L^* \sim \epsilon \), which also equals the ratio of vertical to horizontal bead displacements. Furthermore, since we measure \( |w|/|u| \approx 1/12 \), it gives \( \epsilon \approx 1/12 \), and \( L^* \approx 12a \). Choosing \( a \approx 10 \mu m \) for ram sperm leads to \( L^* \approx 120 \mu m \), which is consistent with the observed range \( L^* \approx 120-145 \mu m \). Furthermore, this leading order scaling predicts that \( L^* \) is independent of the width \( H \) consistent with the observation of a weak experimental variation as \( H \) varies (Table II).

2. Integral scale estimate \( L^* \sim \phi \ell^2/a \) with \( a \approx 10 \mu m \) and \( \ell \approx 50 \mu m \) for ram sperm, \( \phi \approx 1/2 \), predicts \( L^* \approx 125 \mu m \), which is also consistent with the observed range \( L^* \approx 100-150 \mu m \).

3. The ratio of \( (p_x)/(p_y) \sim a/L^* \) indicates a flow structured into preferentially horizontal layers the thickness of which is \( a \), reminiscent of swarming liquid crystals [19–21].

Finally, it is important to recall that the above derivation also includes the hypothesis of strongly confined layering of the suspension, which is consistent with most concentrated semen studied here, but not diluted ones. This observation provides strong evidence to support that the velocity field measured from the image gray-level texture correlation is indeed very close to the depth-averaged true velocity of the flow over the microscope field depth.

B. Discussion and perspectives

Let us first summarize the points raised by this study. We analyze the velocity of concentrated sperm suspensions with PIV reconstruction applied to gray-level textures obtained from phase-contrast microscope high-speed image recordings (as in Refs. [1,4,8,9]). In Sec. II D, we obtain a consistent comparison between PIV measurements on the optical field-depth averaged image and passive fluorescent beads, real local flow tracers velocity. This is strong and direct experimental evidence that PIV provides a measurement of the real velocity field. Since the precise physical origin of gray-level texture associated with phase-contrast images is unclear, such direct evidence was a necessary step. As a side remark, we should also mention that we have indirect evidence for a close link between the phase-contrast signal and local orientational order of nonaxisymmetric sperm cells. Furthermore, we put forward additional coherent evidence:

1. The instantaneous streamline and isocontour lines are nearly identical.

2. Direct and indirect (computing the flow divergence; see Sec. III C) PIV measurements provided a ratio between vertical to horizontal velocity close to 1/12.

3. This ratio has been found to be consistent with the ratio between the horizontal Lagrangian tracers meandering and the microscopic field depth.

4. The integral scale associated with the largest structures of the flow is found to be much larger than the individual swimmers (see Sec. III A) and weakly sensitive to the gap span of the experimental chambers.

5. We found consistent values of the kinematic estimation of the integral scale (provided by a clear experimental linear relationship between the energy and the enstrophy) and that image provided by based extraction from whirlpool characteristics, we found consistent values (see Sec. III A). This is experimental evidence that the typical maximal size of gray-level whirlpools depict the decorrelated units of the flow structure.

6. The horizontal energy spectrum is found to scale like an enstrophy cascade with \( k^{-3\pm0.46} \) power law (see Sec. III B).

7. When decreasing the volume fraction, the large structures of this turbulent flow are mainly unaffected, while the small scales display less energy.

8. The physical evidence for the flow field velocity to display such a specific power-law spectrum has been challenged with an independent experimental measurement based on the separation law of particle beads (see Sec. III D). By estimating the relative velocity \( V \) of each pair of beads with their relative distance \( D \) we found clear evidence for a \( V \sim D \) scaling. Since the theoretical prediction for a \( V \sim D \) scaling is based upon a \( 2Dk^{-3} \) power law of the energy spectrum, this is consistent with the \( k^{-3\pm0.46} \) behavior that we measure for the horizontal energy spectrum.

9. Finally, using a very simple model (see Sec. IV A) for active swimmers in a Newtonian flow where active swimming provides an extra stress, we found that a vertical scale close to the sperm size leads to a consistent scaling for the integral scale.

All these points provide a solid body of evidence that a layered 2D stratified turbulent flow is taking place in the described experiments. We believe that this feature is generic for concentrated suspensions of active and elongated swimmers. This layering can be explained by the combined effect of steric-interaction alignment and boundary influence. Highly concentrated elongated self-propelled particles, close to packing limit have no other choice than local alignment. This steric alignment mechanism applies in the horizontal direction because this is the only possible direction near the upper and lower glass plate. We believe that horizontal alignment propagates from upper and lower boundaries, down into the fluid bulk (the thickness of the chosen chambers are between 10 to 20 cell layers). A simple scaling analysis indicates that if the thickness of the layers are close to the micro-swimmers size \( a \), as suggested by a steric alignment mechanism of this layering, a horizontal integral scale can be predicted, the estimate of which is consistent with our measurements.

Since the proposed mechanism to explain the observed lamination is a steric repulsion alignment, we believe there might be some analogy between the system under study and “swarming nematic” systems (e.g., see Fig. 3 of [19],...
While we do not wish to push further this analogy in more detail, we mention it for completeness. As mentioned earlier, previous studies had already documented the statistical properties of swarming flows of active suspensions [4,5,11]. Most of studies were conducted at volume fraction much lower than ours with the exception of Ref. [8], mainly using bacterial suspensions. Furthermore, the energy spectrum found in these studies rarely presents a simple behavior. Reference [10] report a velocity spectrum which increases at low wave vectors and decreases at large ones. Moreover, the observed decay fitted with a $k^{-8/3}$ at large $k$ over a fraction of decade is indeed quite similar with the $k^{-3}$ reported here. This similarity supports the theory that the reported turbulence is quite generic to stratified or layered anisotropic concentrated active matter.

Finally, it is important to mention that we report in Sec. II F that the rheology of semen displays a power-law shear-thinning behavior. Such non-Newtonian rheology adds an important amount of complexity to the reported measurements. However, we believe that most qualitative features of the reported observations weakly depend on the fluid rheology. As a matter of fact, short-range (possibly lubricated) steric interactions between self-propelled elongated objects will always produce alignment. This is the alignment mechanism which produced at very high concentration the observed layering effect. Hence, the steric induced horizontal layering scenario evoked earlier which is responsible for the resulting 2D behavior of the flow would remain a salient property of anisotropic concentrated active matter flows.

Nevertheless, it is difficult to assess how much fluid rheology influences the observed power-law decay of the energy spectrum. It would certainly be of interest to better understand how the plasma rheology influences the observed dynamics. However, one should bear in mind that separating active cells from plasma is not easy, while, at the same time, preserving their mobility property. It will be experimentally challenging to access the impact of the separation methodology on the apparent changes of active suspension motion.

C. Conclusion

We have performed a systematic experimental analysis of the collective motion of fresh semen confined horizontally between two glass plates. PIV flow fields show “turbulent-like” behavior between the scale associated with the swimmer (spermatozoa) to a well-defined integral scale $L^*$. A $k^{-3+0.4648}$ behavior of the energy spectrum is reported which is consistent with the $V \sim D$ scaling of the separation law. That is typical of $k^{-3}$ 2D energy spectrum. Other indirect quantitative indicators have been put forward, all consistent with an apparent 2D turbulent flow in this concentrated active suspension. The reported results are interpreted to be associated with a laminated turbulent layer flow.

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