

1 The human sperm head spins with a conserved direction during swimming in 3D

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3 Corkidi G.^{1*}, Montoya F.¹, González-Cota A.L.², Hernández-Herrera P.³, Bruce N.C.⁴,

4 Bloomfield-Gadêlha H.^{5*}, Darszon A.^{2*}

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6 ¹Laboratorio de Imágenes y Visión por Computadora, Departamento de Ingeniería Celular

7 y Biocatálisis, ²Departamento de Genética del Desarrollo y Fisiología Molecular, y

8 ³Laboratorio Nacional de Microscopía Avanzada, Instituto de Biotecnología, Universidad

9 Nacional Autónoma de México, Cuernavaca, 62210, México; ⁴Instituto de Ciencias

10 Aplicadas y Tecnología, Universidad Nacional Autónoma de México, Circuito Exterior S/N,

11 Ciudad Universitaria, 04510, Ciudad de México, México; ⁵Department of Engineering

12 Mathematics & Bristol Robotics Laboratory, University of Bristol, UK.

13

14 ***Corresponding authors:**

15 Emails: gabriel.corkidi@ibt.unam.mx, hermes.gadelha@bristol.ac.uk;

16 alberto.darszon@ibt.unam.mx

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18 **Abstract**

19 In human sperm, head spinning is essential for sperm swimming and critical for

20 fertilization. Measurement of head spinning has not been straightforward due to its

21 symmetric head morphology, its translucent nature and fast 3D motion driven by its helical

22 flagellum movement. Microscope image acquisition has been mostly restricted to 2D single

23 focal plane images limited to head position tracing, in absence of head orientation and

24 rotation in 3D. To date, human sperm spinning has been reported to be mono or

25 bidirectional, and even intermittently changing direction. This variety in head spinning

26 direction, however, appears to contradict observations of conserved helical beating of the

27 human sperm flagellum. Here, we reconcile these observations by directly measuring the
28 head spinning movement of freely swimming human sperm with multi-plane 4D
29 microscopy. We show that 2D microscopy is unable to distinguish the spinning direction in
30 human sperm. We evaluated the head spinning of 409 spermatozoa in four different
31 conditions: in non-capacitating and capacitating solutions, for both aqueous and viscous
32 media. All spinning spermatozoa, regardless of the experimental conditions spun
33 counterclockwise (CCW) as seen from head-to-tail. Head spinning was suppressed in 57%
34 of spermatozoa swimming in non-capacitating viscous media, though, interestingly, they
35 recovered the CCW spinning after incubation in capacitating conditions within the same
36 viscous medium. Our observations show that the spinning direction in human sperm is
37 conserved, even when recovered from non-spin, indicating the presence of a robust and
38 persistent helical driving mechanism powering the human sperm flagellum, thus of critical
39 importance in future sperm motility assessments, human reproduction research and
40 microorganism self-organised swimming.

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42 **Introduction**

43 Mammalian spermatozoa invariably spin as they freely swim through a fluid. Similarly to a
44 drill, coordinated helical motion of its whip-like flagellum causes the sperm to “corkscrew”
45 into the fluid, causing the sperm head to spin around its longitudinal axis during locomotion
46 (David et al., 1981; Denehy et al., 1975; Ishijima et al., 1992; Linnet, 1979; Phillips et al.,
47 1972; Rikmenspoel, 1965; Woolley, 1977). Sperm head spinning is a direct manifestation
48 of the cyclic molecular-motor activity shaping the flagellum into a helical beat in 3D
49 (Woolley, 1977; Woolley et al., 1984): a right-handed helical flagellum causes the sperm
50 head to spin clockwise (CW) whilst a left-handed sperm spins in the opposite direction
51 when seen from head-to-tail. Sperm spinning is suppressed if the flagellar waveform is

52 purely planar (2D), as observed for human sperm penetrating viscous medium (Smith et
53 al., 2009). The spinning ability has also been reported to be critical for a successful
54 fertilization, related to a complex cascade of trans-membrane ion channel transport that
55 coordinates this mode of motion (Miller et al., 2018; Zhao et al., 2022). Head spinning is
56 thus a fundamental feature linking the molecular workings of the flagellar beat with sperm
57 motion, and thus an important proxy of symmetry, or rather symmetry-breaking, of the
58 helical flagellar movement in 3D (Zaferani et al., 2021).

59 For the past 50 years researchers have attempted to quantify and define the spinning
60 direction of the human sperm head, though, until now, there is no consensus in the
61 literature of its spinning directionality. Human sperm spinning has been observed to be
62 mono-directed (Linnet, 1979; Smith et al., 2009b; Phillips, 1983; Woolley, 1977), bi-
63 directed (Ishijima et al., 1992; Dardikman-Yoffe et al., 2020; Drake, 1974), and even
64 intermittently directed (Bukatin et al., 2015). However, such reported variety in spinning
65 direction appears to contradict observations of a conserved helical beating of human
66 sperm flagellum (Bukatin et al., 2015; Ishijima et al., 1992; Linnet, 1979; Powar et al.,
67 2022; Zhao et al., 2022), and conserved chirality of structural components in mammalian
68 sperm flagella (Fawcett, 1975; Leung et al., 2021), with no agreement as to the direction of
69 rotation reported in earlier studies (Woolley, 1977; Bishop, 1958; Drake, 1974; Yeung and
70 Woolley, 1984; Woolley, 1979; Blokhuis, 1961; Daloglu et al., 2018).

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72 The human sperm head is translucent and has an axis-symmetric morphology around its
73 spinning axis (Figure 1), making it prone to optical illusions that may obscure accurate
74 detection of its spinning directionality. As such, the sperm head is prone to perception
75 bistability which can further disguise the true spinning direction in translucent objects (Liu
76 et al., 2012). This optical illusion causes spinning translucent objects to appear to oscillate

77 back-and-forth or spin with a switchable, and thus undefined, direction (Liu et al., 2012). At
78 microscale this difficulty is augmented by the contrast inversion effect due to spherical
79 aberrations in microscope objectives that cause a contrast-switch depending on the
80 object's relative position to the plane of focus (Keller et al., 2022; Goodman, 2005).
81 Switching the contrast of spinning objects and its subsequent dependence on the focal
82 plane used for imaging, have unknown consequences on the detected direction of
83 spinning (Figure 2), as we further detail in this paper. Altogether, these challenges indicate
84 that 2D microscopy may not provide accurate measurement of the directionality of sperm
85 head spinning (Muschol et al., 2018). Incidentally, *direct* detection of the head spinning
86 direction in human sperm has been limited thus far to single-plane 2D microscopy. Indirect
87 detection methods either exploit the sperm head centre-position traces (Ishijima et al.,
88 1992) or flagelloid tracks in 3D (Ishijima et al., 1992; Bukatin et al., 2015; Dardikman-Yoffe
89 et al., 2020), by following the trajectory of a certain point along the flagellum. In this case,
90 the head spinning direction is assumed to follow similar rotational movement of the
91 position traces of these counterparts. Indirect detection studies however have not yet been
92 validated against direct measurements of head spinning, as no ground truth is available for
93 this, highlighting this is an urgent gap in literature. Furthermore, different rotations are
94 present during sperm swimming and it is a challenging task to infer head spinning
95 indirectly from flagellar tracks that rotate around an average swimming axis, or from
96 flagelloid tracers that are ill posed and cannot define robustly a common rotation point.
97 Indeed, no modern 3D flagellar tracking research has attempted to directly track the head
98 spinning direction in human sperm. This is, however, critical for understanding sperm
99 swimming as human sperm precesses around its swimming axis: the sperm head spins at
100 the same time that the cell as a whole rotates around its swimming direction.

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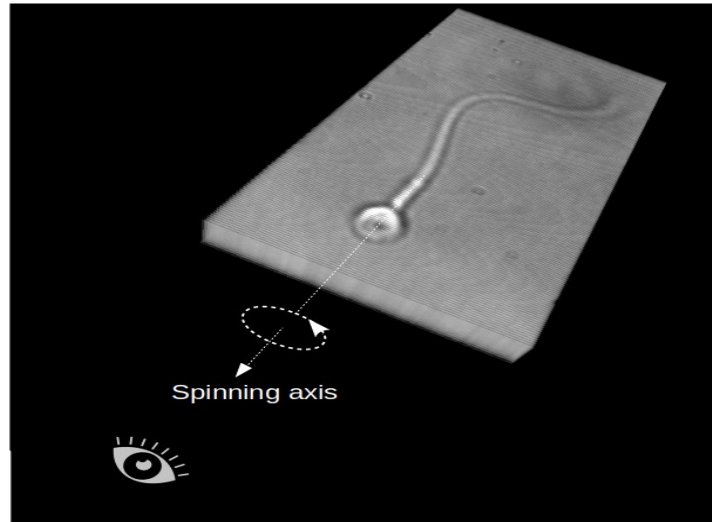
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113 Figure 1. The human sperm head is translucent and has an axis-symmetric morphology

114 around its spinning axis. The reference of observation in this work is from head to tail.

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116 Here we solve this half-century old problem using a high-resolution multi-plane 4D method
117 (Corkidi et al., 2008; Silva-Villalobos et al., 2014; Pimentel et al., 2012; Corkidi et al.,
118 2021; Gadêlha et al., 2020; Hernández et al., 2022) to directly detect the head spinning
119 direction of human spermatozoa. This method employs 4D bright-field microscopy with a
120 100x high-magnification objective that scans a 3D volume with high-speeds as sperm spin
121 and swim through the fluid. The high-precision 4D microscopy described here provides a
122 dense stack of multiple focal planes that bypass limitations of: (i) 2D microscopy that only
123 uses a single focal plane, (ii) the image dependence on focal plane positioning, (iii)
124 difficulties arising from contrast-switch of spherical aberrations of the lens, and (iv)
125 perception bistability effect of translucent spinning objects. We additionally show that head
126 sperm spinning direction cannot be confidently deduced from a single focal plane imaging.
127 We analyzed over 400 free-swimming human spermatozoa in four different conditions,
128 using capacitating and non-capacitating solutions within aqueous and viscous media.
129 100% of all sperm heads were observed to spin counterclockwise (CCW) when viewed
130 from head-to-tail, Figure 1. Our observations show that the spinning direction in human
131 sperm is conserved, even when head spinning is recovered from planar beating, indicating
132 the presence of a persistent helical driving mechanism powering the human sperm
133 flagellum. These observations may have important implications concerning the internal
134 machinery driving the head spinning, flagellum-powered cell motility and its physiology.

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136 **RESULTS**

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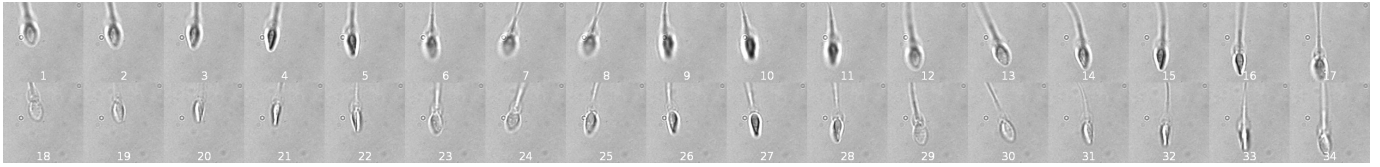
138 **a) 2D single-plane imaging cannot inform confidently the direction of head spin**

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140 Classical 2D single-plane imaging is not suitable to establish confidently the direction of
141 the head spin. Figure 2 shows an image sequence of 17 timepoints evolving from left to
142 right at two different focal planes containing the information of a complete head turn. As
143 can be appreciated, the visual information in these two planes is distinct and complex, with
144 changes in the brightness, contrast switches and head position relative to the focal plane.
145 Establishing the head spinning direction is not possible using single-plane information
146 alone (see Video 1). Visual inspection by different observers leads to different spinning
147 directions due to bistability perception.
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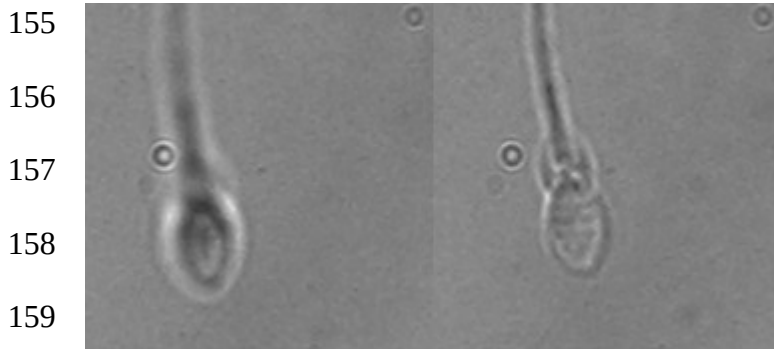
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151 Figure 2. Image sequence of 17 timepoints at two different focal planes separated by 4.8
152 microns (image 1-17 upper focal plane, images 18-34 lower focal plane), containing the
153 information of a complete head turn (see Video 1).

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161 Video 1. Establishing the head spinning direction is not possible using single-plane
162 information alone. Visual inspection by different observers leads to different spinning
163 directions due to bistability perception.

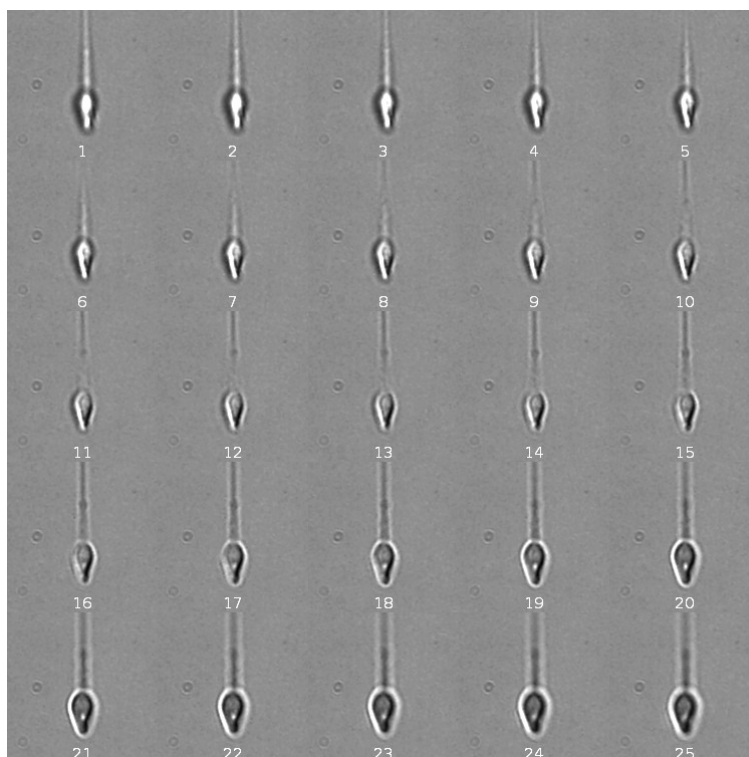
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165 **b) 3D bright field image stacks acquisition and contrast inversion for the human**
166 **sperm head.**

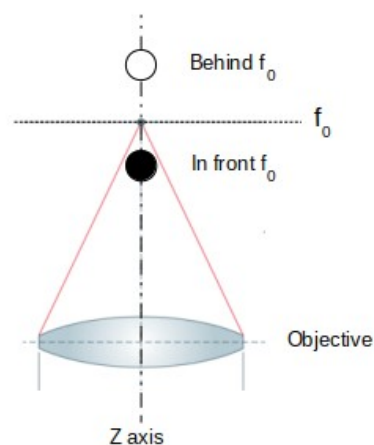
167 Figure 3 shows 25 consecutive focal planes (out of 50) from a single piezo rising slope and
168 assumed to correspond to a single instant. The experimental setup provides a temporal
169 resolution of 1/160 seconds and a spatial resolution along the z direction of 0.4 μm . In this
170 figure, different head elements and their contrast-inversion are revealed as the focal plane
171 moves through the sperm head, appearing as bright pixels inside the head with a dark halo
172 outside to the head behind the focal plane (Figure 3(1)), and as dark pixels inside the head
173 with a bright halo outside to the head in front of the focal plane (Figure 3(25)). Note
174 particularly how the left narrowest sides of the sperm head begin to appear as a bright
175 white border (as detailed in the next sub-sections) when images move farther from the
176 objective lens (15 to 11) in Figure 3. This contrast inversion effect will be explored in the
177 following section to track head spin directionality.

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197 Figure 3. Multifocal plane acquisition. From left to right and top to bottom, this figure shows
198 25 consecutive focal planes (out of 50) acquired while the piezoelectric device is rising.
199 Frames (150 x 150 pixels each) were cropped, for visualization purposes, from 640 x 480
200 pixels image sequences. Each focal plane is separated by 0.4 μm respectively (see
201 Materials and Methods). The schematic at the bottom depicts how the spherical aberration
202 of the optical system produces a contrast inversion over translucent objects depending on

203 their position with respect to the focal plane f_0 . The object appears dark if placed in front of
204 the focal plane f_0 , while bright if placed behind.

205 **c) Contrast inversion of the sperm head is due to the spherical aberration of the**
206 **objective lens using bright-field microscopy.**

207 The Rayleigh-Sommerfeld back propagation reconstruction method has been commonly
208 used to reconstruct the sperm flagellum in 3D (Bukatin et al., 2016), but this is based
209 solely on diffraction and cannot distinguish between defocus before or after the focused
210 object plane (Lee et al., 2007). Usually, the contrast inversion of the defocused object is
211 used heuristically to post process the results of the back-propagation method to produce a
212 final reconstructed object (Lee et al., 2007), alternatively, phase information from the
213 reconstructed object can be used to post-process the reconstruction itself (Wilson et al.,
214 2012). It is well known that microscope objectives are extremely well corrected for
215 conjugate planes, which are focused objects and image planes, but this is not true for non-
216 conjugate planes. Aberrations, particularly spherical aberrations, can strongly contribute to
217 image quality for non-conjugate planes (Keller et al., 2022; Kidger, 2002). Here we
218 demonstrate that it is the spherical aberration of the objective lens that produces a contrast
219 inversion as a function of the position of the object relative to the focused object plane
220 (Figure 3).

221 Diffraction effects that appear in the point spread function (PSF) are given by the Fourier
222 transform of the lens aperture function multiplied by the aberration function,

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$$PSF = \mathbf{F}(P(x, y) \exp(ikW(x, y))), \quad (1)$$

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226 where $\mathbf{F}()$ is the Fourier transform in the lens aperture plane, (x, y) is the position in the
227 image plane, $P(x, y)$ is the lens pupil (almost always taken as a circle), $k=2\pi/\lambda$ is the
228 light wave number, and $W(x, y)$ is the wave aberration function given by (Goodman,
229 2005)

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$$W(x, y) = A_d \frac{x^2 + y^2}{r^2} + A_s \left(\frac{x^2 + y^2}{r^2} \right)^2, \quad (2)$$

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233 where A_d is the amplitude of the defocus, A_s is the amplitude of the spherical aberration,
234 and r is the radius of the lens pupil. The first term in Equation 2 is the defocus contribution,
235 and the second term is the contribution of the spherical aberration. One important aspect
236 is the sign of the defocus term. From Goodman 2005, using the Gauss equation for a
237 defocused system we have:

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$$\frac{1}{d_0} + \frac{1}{d_i} = \frac{1}{f} \rightarrow \frac{1}{d_0} + \frac{1}{d_i} - \frac{1}{f} = -A_d, \quad (3)$$

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241 so that $A_d=0$ when the system is focused. Given that f and d_i (focal length and the
242 distance from the lens to the image) are fixed, and the distance from the object to the lens
243 (d_0) changes when the optical system is defocused, when d_0 increases (the object is

244 further away from the objective lens), the term $\frac{1}{d_0} + \frac{1}{d_i}$ is smaller and $A_d > 0$. When d_0

245 decreases (the object is closer to the objective lens), the term $\frac{1}{d_0} + \frac{1}{d_i}$ increases and $A_d < 0$.

246 We assume that the spherical aberration has the same sign for a defocus below and
247 above the geometrical focal plane. Simulations for a rectangular object are shown in
248 Figure 4. The diffraction from defocus alone in Figure 4A and B, does not give an inversion
249 of contrast between the object positions below and above the focused object plane. Figure
250 4C and D, however, where a small spherical aberration contribution has been included,
251 does show the contrast inversion effect. Finally, Figure 4E shows the case of an inclined
252 object where the left part of the object is further away from the objective lens and the right

253 part is closer to the objective lens, showing the contrast inversion within the object. It is
254 important to note that a change in sign of the spherical aberration contribution changes the
255 contrast inversion: with $A_s > 0$ an object closer to the objective lens than the focused
256 object plane would be dark and an object further from the objective lens would be bright.
257 The sign of the aberration depends on the specific design of the optical system and so
258 could vary between different laboratories working with different objectives. Our simulations
259 consider a spherical aberration within the range of the optical system used in our
260 experiments.

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262 The contrast-switch inside the object is a direct manifestation of the object's inclination
263 relative to the objective lens (Figure 4E). Here we exploit for the first time this unique
264 optical effect to extract the directionality of a spinning object, such as the sperm head, as
265 detailed below. Figure 5 shows how this contrast-switch of its planar projection varies
266 when the inclination of the object increases as the object spins around its long axis.
267 Clockwise spinning causes the bright part (further away from the objective lens, see white
268 arrows) of the inclined object to always move from right-to-left and the dark part to move
269 from left-to-right (dark arrows), whilst counterclockwise spinning causes the bright region
270 to always move from left-to-right and at the same time that the dark region moves from
271 right-to-left. Once a half-cycle is completed, and the flat object is parallel to the objective
272 lens, the bottom part of the object reaches the top and vice versa, re-setting the contrast
273 inversion: any part of the object reaching the top appears as bright, likewise any part of the
274 object reaching the bottom will appear as dark. In other words, after a half-cycle, a
275 bright/dark region abruptly re-appears at starting location, after reaching the end of the
276 object on the opposite side. For this reason, if an object spins in the same direction, this
277 will be manifested in the image as the bright or dark regions of the objects moving with a

278 persistent direction relative to the object's orientation (left-to-right or right-to-left). Likewise,
279 if the object spinning direction is reversed, the direction of motion of bright/dark regions will
280 equally reverse relative to the object's orientation (right-to-left or left-to-right). The
281 spherical aberration effect causing the contrast switch within the object's image allows for
282 robust, and yet simple, detection of spinning direction of axis-symmetric objects, such as
283 the human sperm head demonstrated below, that otherwise would not be possible, and
284 this feature is exploited here for the first time.

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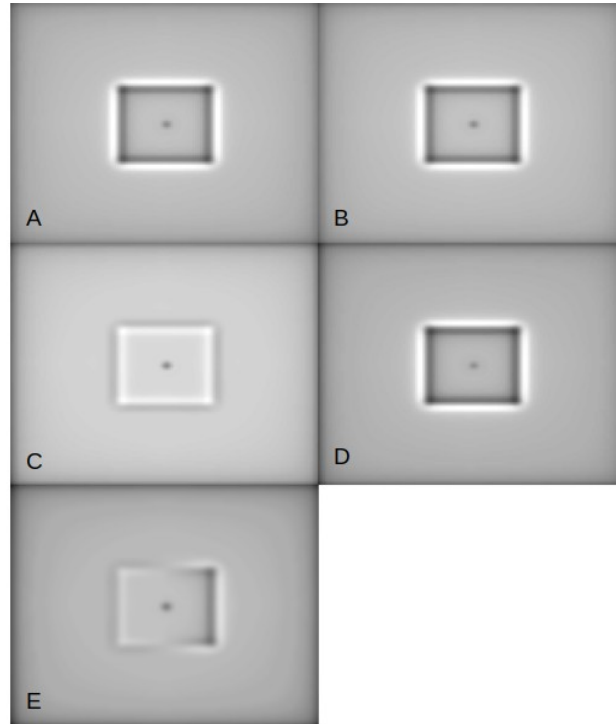


Figure 4. Results of the simulation of image formation for a weakly scattering phase object.

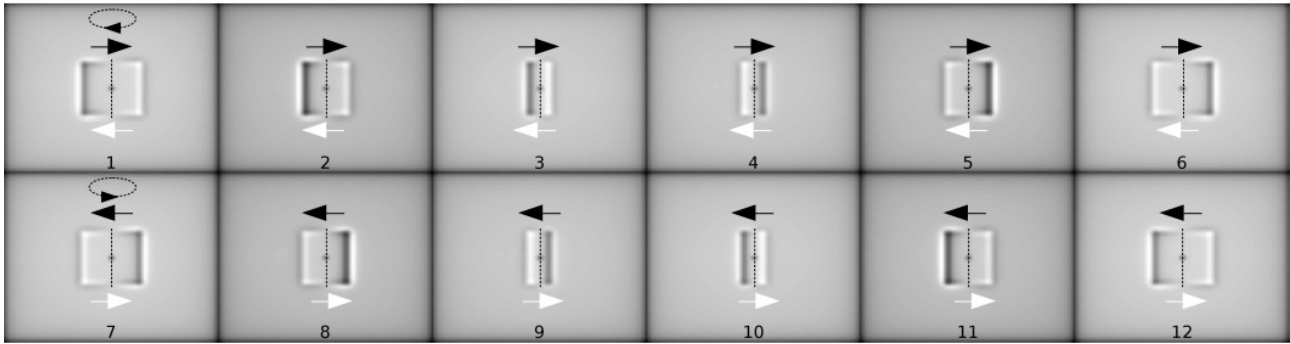
(A) image with $A_s=0$ and $A_d=0.3$; (B) image with $A_s=0$ and $A_d=-0.3$; (C) image with

$A_s=-0.0004$ and $A_d=0.04$; (D) $A_s=-0.0004$ and $A_d=-0.04$; (E) image for an inclined

planar object with $A_s=-0.0004$. The centre of the object plate is in the focused object plane

of the lens and the right part of the object is closer to the objective lens.

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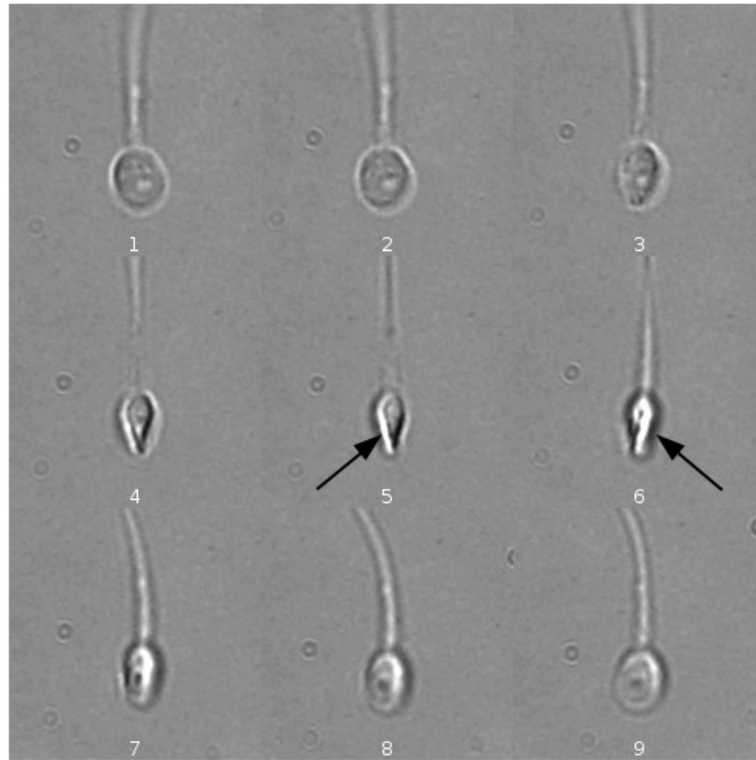
313 Figure 5. Simulation results of a spinning object, showing the direction of movement of the
314 dark and bright left/right edges from 2D imaging from a fixed focal plane projection. Top:
315 (images 1 to 6) a clockwise rotation viewed from the top of the images; bottom: (images 7
316 to 12) a counterclockwise rotation. $A_s = -0.004$ and the maximum defocus is $A_d = \pm 0.2$.

317 **d) Bright region induced by sperm head inclination moves with head spinning.**

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319 The human sperm head has a flattened side, and when the head spins around its
320 longitudinal axis, the narrow side behind the focal plane is clearly seen as a bright region,
321 whilst the part of the head above the focal plane appears as dark, when the sperm head is
322 inclined relative to the objective lens (Figure 6, images 4-6, see arrows). The contrast
323 switch inside the sperm head is due to the head inclination relative to the objective lens
324 (images 4-6); when the sperm head is parallel to the objective lens, no switch in contrast
325 occurs (images 8-9), as demonstrated in the above section. The time-sequence in Figure 6
326 (images 4-7), shows the bright region moving from left-to-right relative to the sperm head
327 orientation due to changes in inclination of the sperm head caused by the head spinning.
328 This switch in contrast when the head is inclined is barely perceptible when observed with
329 2D microscopy (Figure 2), as the sperm head continuously changes its position relative to
330 the focal plane when swimming freely in the fluid (see Video 1). A minimal focal change at
331 the micron scale is sufficient to prevent the detection of bright-dark regions induced by the
332 sperm head inclination, as shown in Figure 4E. Indeed, Figures 2 and 6 show the inversion
333 in contrast due to inclination of the sperm head is clearly visible for only a few focal planes
334 as the piezo rises. It would be a challenging task to change a single focal plane
335 dynamically, at the microscale, to keep the sperm head exactly in focus to enable the
336 detection of this change in contrast optical effect across the head using 2D microscopy, as
337 shown in Figure 5. Our multifocal system bypasses this challenge and allows unique
338 detection of the accumulated changes in contrast from bright-to-dark regions induced by
339 the sperm head inclination from a stack of multiple focal planes (Figure 3), as detailed
340 below.

341



342 Figure 6. The contrast inversion optical effect during a half turn of a sperm head. Nine
343 consecutive timepoints where the contrast inversion optical effect is clearly observed
344 (same focal plane), i.e. the narrowest, focally lower, border of the sperm head (behind the
345 focal plane) is enhanced resulting in a bright border while the sperm head is inclined
346 during head spinning (highlighted with arrows).

347 **e) Validation of the sperm head spinning detection using contrast switch of the**
348 **sperm head.**

349 We validated the use of the contrast inversion of the sperm head shown previously to
350 detect the head spinning direction against direct tracking of the motion of a particle
351 attached to a sperm neck during head spinning. Figure 7 shows a sperm cell with such a
352 particle rigidly attached to its neck, whilst spinning 360 degrees during free-swimming
353 motion. In the time-sequence for the Figure 7A, the focal plane is approximately placed
354 between the sperm and the particle along z, in such a way that when this particle is behind
355 the focal plane (images 1-4), it appears as bright, while when the particle is in front of the
356 focal plane (images 5-8), it appears as dark. The arrow on the particle shows the direction
357 of the displacement during head spinning; the black arrow indicates that the particle is
358 behind the focal plane (bright particle), while the red arrow indicates when the particle is in
359 front of the focal plane (dark particle). As such, the particle rotates in the CCW direction
360 following the sperm head spinning motion: the particle moves from left-to-right when
361 behind the focal plane (1-4), and moves from right-to-left when above the focal plane (5-8).
362 On the other hand, the bright border arising from the contrast inversion when the sperm
363 head is inclined relative to the objective lens always moves in the same direction relative
364 to the sperm head orientation, from left-to-right, as shown by the arrow on this bright
365 region, regardless of whether the particle is above (5-8) or below (1-4) the focal plane in
366 each of its half-cycle (see Video 2). CCW spinning (as seen from head-to-tail) causes the
367 sperm head bright region to always move from left-to-right relative to the sperm head, in
368 agreement with the direction of the particle rotation above.

369 When the focal plane is placed between the particle and the flagellum (Figure 7B), it
370 becomes evident their relative position: in the sequence 1-2 (top images), the particle
371 (bright) is located behind the flagellum (dark) while moving to the right, while in the

372 sequence 3-4, the particle is in front (while moving to the left). The movement direction of
373 the particle, combined with its relative position to the flagellum makes to confidently
374 conclude a CCW spin.

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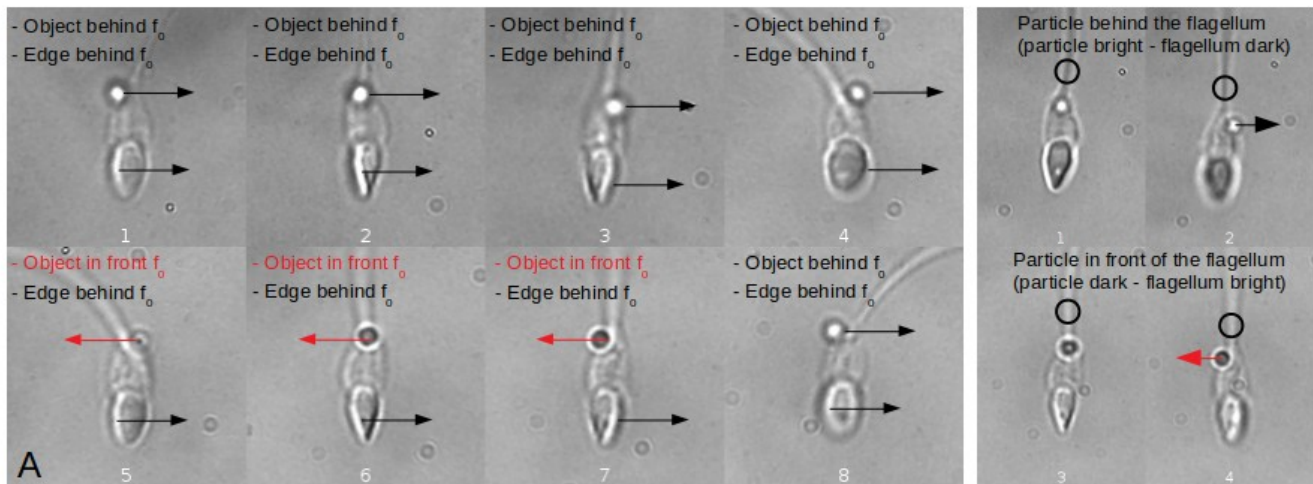
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384 Video 2. A sperm cell with a transluclid particle rigidly attached to its neck, whilst spinning
385 360 degrees during free-swimming motion.



386 Figure 7. Tracking a translucent particle stuck on a spermatozoon's neck, presenting the
387 contrast inversion optical effect while turning 360 degrees with a free-swimming sperm. (A)
388 Focal plane between Particle and Bright border: when the particle is above the
389 spermatozoon's neck it appears bright (behind the focal plane), while appearing dark when
390 under the neck (in front of the focal plane); simultaneously, the border of the spermatozoon
391 is always behind the focal plane appearing bright, showing a CCW spin direction when
392 seen from head-to-tail. Note the change of direction of the particle when it is dark, as the
393 particle rigidly follows the head spinning. (B) Focal plane between Particle and Flagellum:
394 when the focal plane is placed between the particle and the flagellum, it becomes evident
395 their relative position: in the sequence 1-2 (top images), the particle (bright) is located
396 behind the flagellum (dark) while moving to the right, while in the sequence 3-4, the
397 particle is in front (while moving to the left). A CCW spin is evident from these images. This
398 shows that the proposed method is capable of detecting spinning directions.

399 **f) Detection of sperm head spin direction from multifocal stacks**

400 Here, we exploit the volume stack and accumulate the contrast-switch caused by the head
401 inclination occurring at different focal planes by integrating this multifocal plane information
402 in a single image. This circumvents the fact that the contrast switch across the sperm head
403 is only observable for a few focal planes, as shown above. To this purpose, we calculated
404 a 2D Maximum Intensity Projection (MIP) image (Schindelin et al., 2012) for each piezo
405 rising slope z stack containing 50 focal planes through the whole acquisition time of 3.5
406 sec. The 2D MIP of the z-stack accumulates in a single image the maximum values of all
407 the focal planes, i.e. bright regions in the image shown in Figure 6 for a time sequence.
408 The integrated bright region induced by the switch in contrast when the sperm head is
409 inclined is manifested as a superimposed one-sided halo, only present when the sperm
410 head is inclined relative to the objective lens (images 4-7). Figure 8 shows the
411 accumulated bright region moving from left-to-right relative to the head as time progresses
412 from time sequence 4-7 (see also Video 3). The motion of the accumulated MIP bright
413 feature over the course of time follows the direction of head spinning, as demonstrated in
414 previous sections.

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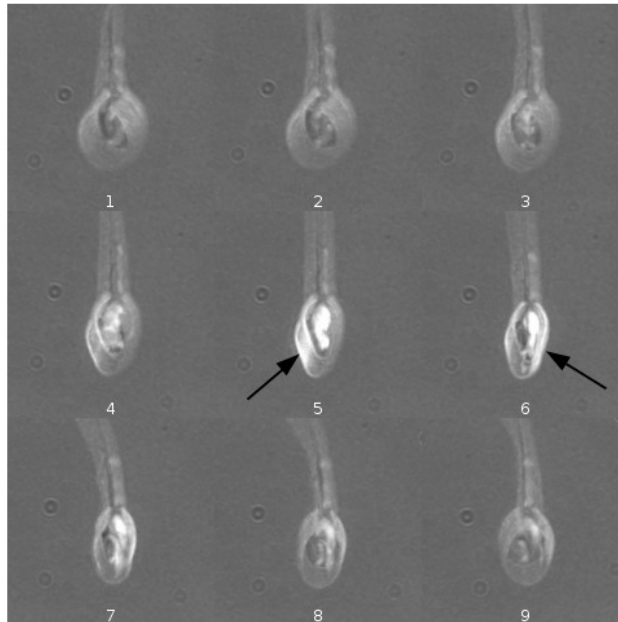
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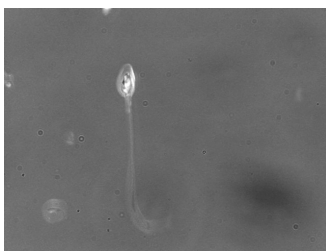
425 Figure 8. 2D MIP's (Maximum Intensity Projections) of 9 consecutive piezo rising slopes -
426 timepoints- (50 images each projection, images 1 to 9) during a half turn of a sperm head
427 (left to right, top to bottom). The brightest pixels over the z axis from 50 images
428 corresponding to one half cycle of the piezo device are projected into a single plane to
429 integrate the spherical aberration spread over the z range. The bright region moves from
430 left-to-right relative to the sperm head (arrows) (See Video 3).

431

432

433

434



435 Video 3. The 2D MIP of the z-stack accumulates in a single image the maximum values of
436 all the focal planes, i.e. bright regions for a time sequence. A bright region moves from
437 right-to-left relative to the head as time progresses. At the final part of the video, another
438 spermatozoon appears swimming in the opposite direction, showing the bright region to
439 move consistently in the inverse direction (see also Supporting Information and Figure S1).

440 The directional motion of the superimposed bright regions is extracted from the variations
441 on the intensity profile moving along the segment **bb'** (red line shown in Figure 9). To
442 detect the direction of the movement in which the one-side bright halo moves with respect
443 to the longitudinal axis of the head (as placed in front of the head -viewing the sperm from
444 the tip of the head to the flagellum-, line **C** Figure 9A), we have employed the following
445 steps:

446

447 1) To track the sperm head position, we used the method described in (Corkidi et al.,
448 2021). This method provides the spermatozoa dominant orientation over time (angle ϕ in
449 Figure 9A) relative to the microscope fixed frame of reference, depicted by the line **C**. The
450 center of the sperm head over line **C** is defined as a . These two experimental parameters
451 define the position and orientation of the major axis of the sperm head represented by **C**
452 (see Corkidi et al., 2021). By construction, **C'** is perpendicular to **C**, and the intersection
453 point is at a distance of d μm from a ($1/3$ the average size of the long axis of the human
454 sperm head). Finally, **b** and **b'** are located symmetrically along **C'** at a distance **b** from the
455 intersection of **CC'**. The pixels along **bb'** (represented by the red line) are used to measure
456 the intensity profile over the 2D MIP image (see Video 4). The direction of motion of the
457 maximum intensity of the profile series (by using variation in position of the weighted
458 average -see next section- along the segment **bb'**, as shown in Figure 9C and D) defines
459 the sperm head spin direction. The equation describing the points along the line **C** is

460 $y = (\tan\phi)x + (y_a - x_a \tan\phi)$, the intersection point between **C** and **C'** is given by the condition

461 $d = \sqrt{(x - x_a)^2 + (y - y_a)^2}$ and its coordinates (x_d, y_d) . The equation describing the points

462 along **C'** is defined by $y' = \left(\frac{-1}{\tan\phi}\right)x' + \left(y_d + \frac{x_d}{\tan\phi}\right)$. The points **b** and **b'** are positioned at

463 distances $b = \pm \sqrt{(x' - x_d)^2 + (y' - y_d)^2}$ along \mathbf{C}' centered at the intersection with \mathbf{C} . The set of
464 points that are embedded in the \mathbf{bb}' segment are $\{\vec{r}_i\}_{i=1}^N$, where N is the number of points
465 belonging to \mathbf{bb}' (red line in Fig 9.). From this, the gray level profile over the segment \mathbf{bb}'
466 can be quantified.

467

468 2) Determination of the sperm head spinning direction.

469 The position of weighted average of each intensity profile along \mathbf{bb}' , while the head is
470 turning, is quantified to establish the head spin direction. The weighted average position
471 over \mathbf{bb}' is \vec{r}_{wa} , analogous to the “center of mass” of the intensity levels along \mathbf{bb}' .
472 Tracking \vec{r}_{wa} over time reveals the movement of the brightest part of the sperm head 2D
473 MIP. This feature applied to the gray level profile over the line segment \mathbf{bb}' (see Figure 9)

474 is expressed as: $\vec{r}_{wa} = \frac{\sum_{i=1}^N I_{MIP,i} \vec{r}_i}{\sum_{i=1}^N I_{MIP,i}}$, where $I_{MIP,i}$ is the gray level value of the corresponding

475 2D MIP image \vec{r}_i . If \vec{r}_{wa} moves from \mathbf{b} to \mathbf{b}' the cell is rotating CCW and otherwise CW, as
476 shown in Figures 5 and 9.

477 Figure 9A and B shows the segment \mathbf{bb}' positioned over the sperm head in two
478 consecutive time points with the bright border moving from \mathbf{b} to \mathbf{b}' . The corresponding gray
479 level of the 2D MIP profile is plotted at the bottom of each image in C-D. Figure 9E shows
480 the values of the weighted average for two and a half head turns as shown in Video 3 and
481 Video 4. Red circles over the first minimum and maximum correspond to the first half turn.
482 Minima denote that the weighted average of the intensity profile is shifted towards the \mathbf{b}
483 side of the sperm head (taking \mathbf{b} (from \mathbf{bb}') as the origin for the profile), while maxima

484 denote the shift towards the opposite **b'** side. Figure 9F consists of the intensity profiles
485 (**bb'**) kymograph; lower dashed lines point out the bright profiles corresponding to Figures
486 A,B,C,D denoting a 180 degrees head turn. The upper time ascending arrow from left-to-
487 right of the kymograph denotes a CCW head turn.

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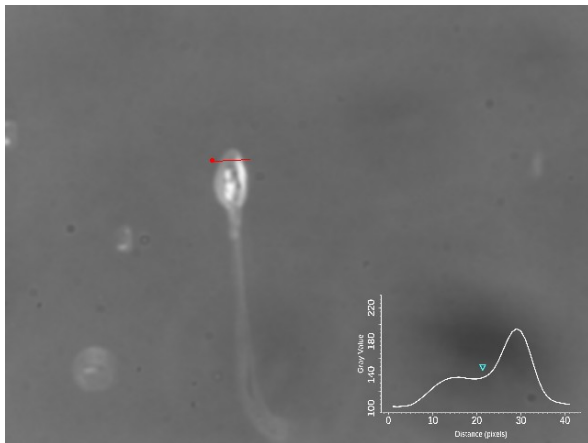
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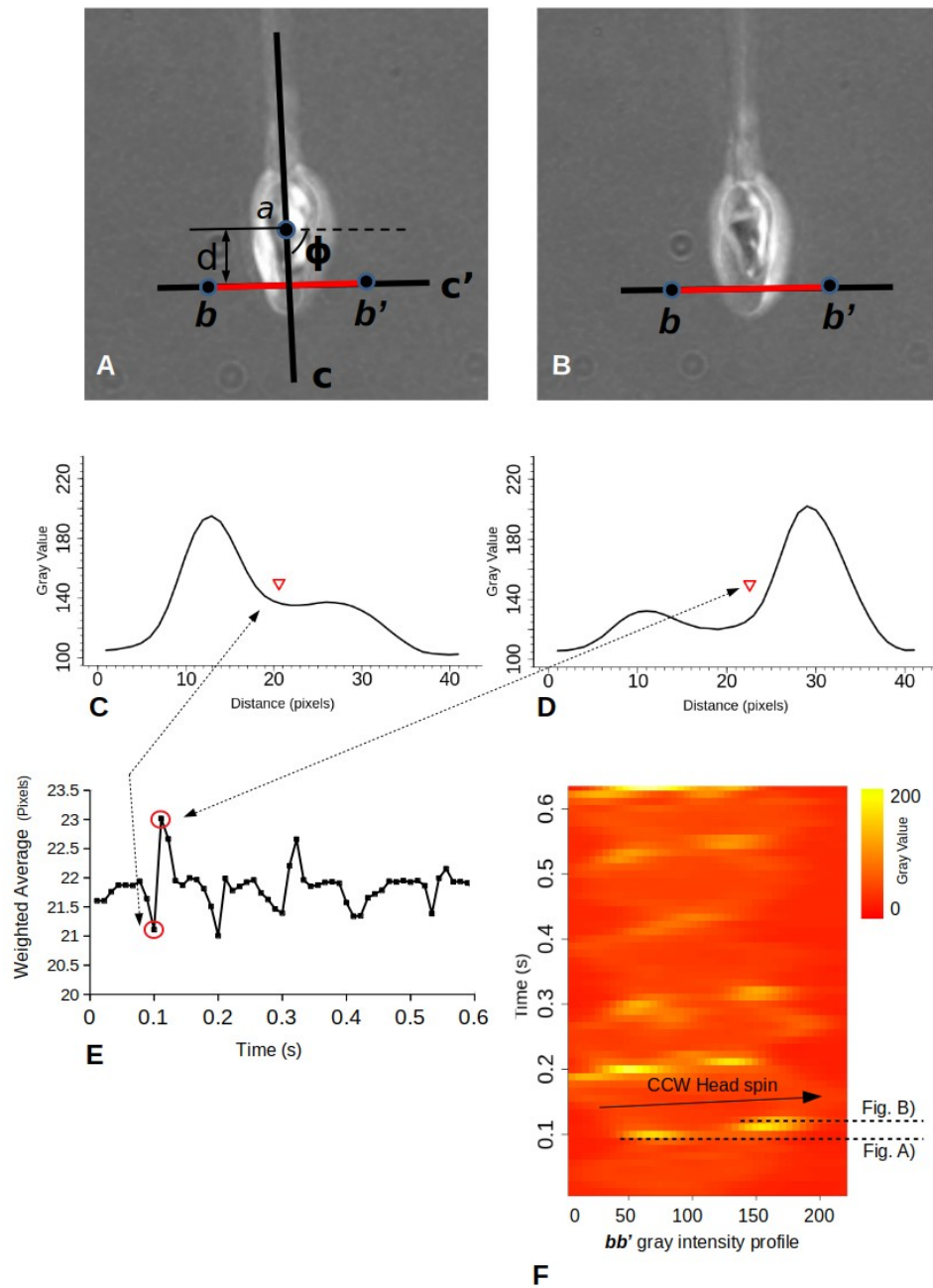
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496 Video 4. Gray level of the 2D MIP profiles (over the tracked red line **bb'** from Figure 9) with
497 its center of mass depicted in the graph by the blue triangle.

498



500 Figure 9. Sperm head bright region travels with a conserved direction. (A-B) The tracking
 501 process estimates the angle ϕ of the dominant direction of the spermatozoon relative to
 502 the microscope frame of reference and defines the orientation of the sperm (line **C**, see
 503 Corkidi et al., 2021). The straight-line **C'** is perpendicular to **C**, and the intersection point is
 504 at a distance of d μm from head center (1/3 the average size of the long axis of the human
 505 sperm head). **b** and **b'** are located symmetrically along **C'** at a distance **b** from the

506 intersection of **CC'**. The coordinates of **bb'** (represented by the red line) are used to
507 measure the intensity profile over the 2D MIP image (see Video 4). (C-D) The
508 corresponding gray level of the 2D MIP profiles (from A-B) with its center of mass depicted
509 by the red triangle. (E) Weighted average of 2D MIP profiles for two and a half head turns.
510 Minima denote that the weighted average of the intensity profile is shifted towards the **b**
511 side of the sperm head (taking **b** (from **bb'**) as the origin for the profile) while maxima
512 denote the shift towards the opposite **b'** side. The transition from a minimum to a
513 maximum corresponds to half a turn of the sperm head in a CCW direction (two and a half
514 CCW turns for this series). (F) Intensity profiles (**bb'**) kymograph; dashed lines point out
515 the bright profiles corresponding to Figures A and B denoting a 180 degrees head turn.
516 The lower time ascending arrow show waves of the brightness level traveling from left-to-
517 right, indicating CCW head spin when seen from head to tail.
518

519 **g) Human sperm spin with a conserved counterclockwise direction.**

520 We acquired 409 human spermatozoa while freely swimming in 3D; 180 under non-
521 capacitating conditions and 229 in capacitating conditions. Both groups were studied in
522 aqueous and viscous media (see Materials and Methods). We found that in aqueous
523 media, regardless of the medium used (non-capacitating or capacitating), all the sperm
524 spun in the CCW when seen from head to tail and swimming freely. In contrast, in viscous
525 and non-capacitating medium, 57% of the analyzed sperm were not spinning at all, while
526 the remaining 43% spun in the CCW direction. Interestingly, in capacitating conditions
527 these percentages inverted: 22% of the analyzed sperm did not spin at all, whilst 78%
528 spun with the conserved CCW direction. The results summarized in Table 1 suggest that
529 capacitating media also influences the ability of sperm to spin in a viscous fluid. All human
530 sperm observed to spin did so by turning in the CCW direction when seen from head to tail
531 regardless of the experimental condition.

532

533

Non capacitating media

Capacitating media

535

Aqueous

Viscous

Aqueous

Viscous

536 CCW

78 (100%)

65 (43%)

106 (100%)

101 (78%)

537 CW

0

0

0

0

538 No rotation

0

37 (57%)

0

22 (22%)

539

540 Table 1. Spin direction evaluation of 409 human spermatozoa while freely swimming in 3D;
541 180 under non-capacitating conditions and 229 in capacitating conditions (both groups in
542 aqueous and viscous media, see Materials and Methods). All human sperm in aqueous
543 media (non-capacitating or capacitating) spun CCW direction when seen from head to tail
544 regardless of the experimental condition, while in viscous and non-capacitating medium,
545 57% of the analyzed sperm did not spin at all, while the remaining 43% spun in the CCW
546 direction. Note that in capacitating conditions these percentages inverted: 22% of the
547 analyzed sperm did not spin at all, whilst 78% spun with the conserved CCW direction.

548 **Discussion**

549 Sperm flagellum elastohydrodynamics, mathematical modelling and image analysis have
550 indicated that the sperm head movement is highly dependent on the nature of forces and
551 torques imposed by the beating flagellum and subsequent interactions with the local
552 environment, among many other factors (Gadêlha et al., 2010; Gadêlha et al., 2019; Smith
553 et al., 2009b; Ishimoto et al., 2017; Gaffney et al., 2011). Indeed, self-organization flagellar
554 control models have demonstrated that mechanical attachment of the head (clamped or
555 hinged head conditions) can even dictate the travelling wave direction of the flagellum
556 (Riedel-Kruse et al., 2007; Camelet and Jülicher, 2020; Oriola et al., 2017; Sartori et al.,
557 2016). These matters highlight the critical importance of directly establishing the head
558 movement and its rotations in 3D. Furthermore, despite this critical importance and recent
559 advances in high-speed 3D imaging and 3D microscopy, for decades there is still no
560 consensus as to what direction human sperm spin during free-swimming motion (Muschol
561 et al., 2018). Reports include observations of: mono-directed CW or CCW (Linnet, 1979;
562 Smith et al., 2009b; Phillips, 1983; Woolley, 1977), bi-directed (Ishijima et al., 1992;
563 Dardikman-Yoffe et al., 2020; Drake, 1974), and even intermittently directed head spinning
564 (Bukatin et al., 2016). Against this background, direct detection of sperm head spinning
565 direction and its methodology are still lacking in the literature.

566 Phillips, 1983 has suggested that mammalian sperm spinning direction could be easily
567 detected with 2D bright field microscopy, by exploiting the fact that the sperm head is
568 flattened, and thus produces blinking “flashes of light” as the head spins around its
569 swimming axis (Phillips, 1997). It has been observed that the ‘flash of light’ travels from
570 left-to-right when the sperm head-to-tail is aligned with the vertical axis (similarly to the
571 sperm head orientation depicted Figure 2, and thus CW head spin was inferred. This
572 however postulates that such ‘flash of light’ *moves in the same direction as the sperm*

573 *head spinning*. We have demonstrated here this to be inconsistent with direct detection of
574 the head spinning, see Figure 7. Instead, we have found that the true head spinning
575 direction is opposite to the observed left-to-right movement of this “flash of light”, which
576 similarly to Phillips, 1983 also moves from left-to-right, as depicted in Figure 7. We have
577 shown that this mismatch of movement between optical brightness and the object spinning
578 motion is accounted for by spherical aberration effects of the lens. Furthermore,
579 translucent objects, such as the sperm head, are equally prone to perception bistabilities
580 Liu et al., 2012 that equally obscure the true head spinning direction with 2D microscopy,
581 in addition to other unknown image inversions within microscopy systems (see Methods).
582 These uncertainties, together with the combined use of 2D views of the sperm’s head
583 trajectory as a proxy to derive flagellar beat information, may have significantly contributed
584 to the contradictory observations that are documented in the literature regarding the sperm
585 head spinning direction.

586 In the present study, we evaluated the head spinning direction of more than 400
587 spermatozoa. Sperm suspended under four conditions were tested: in non-capacitating
588 and capacitating solutions of normal and high viscosity. The high viscosity value was
589 chosen to emulate that of the female cervical mucus (see Methods and Suarez, 2016) and
590 sperm were incubated in capacitating media for 6 hours. One hundred percent of the
591 spinning spermatozoa, in all experimental conditions, spun CCW, as seen from head-to-
592 tail. It is important to mention that the spinning direction of each single spermatozoon was
593 evaluated for a mean time of 3.4 s. A longer temporal analysis would be desirable to
594 evaluate whether spinning direction changes over periods longer than 3 sec, though
595 unfortunately not possible with our present high-resolution 4D set-up. Taken altogether, the
596 analyzed time of the 409 free-swimming sperm totalled 23 minutes, with no directional
597 change observed. Human spermatozoa in aqueous media, independently of their

598 capacitation state, all spun CCW. In contrast, in high viscosity, 57 % of spermatozoa in
599 non-capacitating media did not spin at all, and interestingly most of them recovered their
600 CCW rotation (80 %) when incubated in capacitating media. This striking head spinning
601 recovery phenomenon, as well as how sperm head spinning motion and flagellar rolling
602 influence swimming trajectories in 3D remain to be fully explored.

603

604 **Conclusions**

605

606 We studied here a half-century old problem by exploiting unique switch in contrast due to
607 spherical aberration effect in brightfield microscopy. We have shown that 2D microscopy
608 alone cannot distinguish spinning direction in axis-symmetric, streamlined, translucent
609 human swimming sperm; as well, methods employing such imaging technique may need
610 reassessment, whilst no methodology are currently available to directly measure human
611 head spinning. Indeed, previous studies mostly used visual inspection from video-
612 microscopy images or indirect measurement using flagellar tracing in 3D. We showed that
613 contrast inversion can be exploited to track head spinning but this requires finding the
614 appropriate focal plane in which the sperm head is, centred exactly at the focal plane; a
615 challenging task for freely-swimming spermatozoa as the head moves up and down during
616 cell progression. This is resolved by using a multi-plane detection system. The
617 methodology was validated as coherence and consistence prevailed between optics
618 theory and direct tracking of different sperm cells with particles attached to neck and head.
619 We have observed that human sperm head spins with a robust, conserved, and
620 recoverable counterclockwise spinning direction (when viewed from head to tail). Our
621 observations reconcile structural information of mammalian sperm that observe flagellar
622 architecture with conserved chirality in the axonemal driving unit with a conserved direction

623 of spinning for human sperm, regardless of viscosity and capacitating conditions. The
624 ability of human sperm to fertilize may be intimately related with head spinning, as
625 capacitating medium were observed to excite a larger proportion of the population to spin.
626 At last, the proposed methods can be applied to other free spinning objects and
627 microorganisms that possess similar axis-symmetric body architecture to human
628 spermatozoa.

629

630 **Materials and Methods**

631 ***Ethical approval for the human semen samples***

632 The bioethics committee of the Institute of Biotechnology, UNAM approved the proposed
633 protocols for the handling of human semen samples. Donors were properly informed
634 regarding the experiments to be performed and each donor signed and agreed to a
635 consent form. All samples fulfilled World Health Organization requirements for normal
636 fertile semen samples.

637 ***Media***

638 HTF (human tubal fluid) medium was used in this study. Non-capacitating HTF (pH 7.4)
639 contained (mM): 4.7 KCl, 0.3 KH₂PO₄, 90.7 NaCl, 1.2 MgSO₄, 2.8 Glucose, 1.6 CaCl₂, 3.4
640 sodium pyruvate, 60 sodium lactate and 23.8 HEPES. Capacitating medium (pH 7.4) was
641 HTF medium supplemented with 5 mg/ml BSA and 2 mg/ml NaHCO₃. Capacitating
642 recording medium (pH 7.4) was only supplemented with 2 mg/ml NaHCO₃, BSA was not
643 added. For viscous medium, 1% methyl cellulose was added to the non-capacitating or
644 capacitating recording medium, depending on the experimental condition.

645 ***Biological preparations***

646 Semen samples were obtained by masturbation from healthy donors after 48 h of sexual
647 abstinence. Highly motile sperm were recovered after the swim-up protocol. Briefly, 300 μ l
648 of semen were placed in a test tube, then 1 ml of non-capacitating or capacitating medium,
649 depending on the experimental condition, was added on top. Tubes were incubated at a
650 45° angle, at 37°C in a humidified atmosphere of 5% CO₂ and 95% air during 1 h. Then,
651 sperm from the medium on top were collected and concentration was adjusted to 10⁶ cells/
652 ml. To promote *in vitro* capacitation, sperm in capacitating medium were incubated for an
653 additional 5 h. Recordings were performed in (a) non-capacitating aqueous and viscous
654 medium, and (b) capacitating aqueous or viscous medium.

655 ***Sperm samples***

656 A total of 409 freely swimming spermatozoa (30 samples -one per day- from 9 different
657 donors) were analyzed: (a) 180 non-capacitated (78 in aqueous media and 102 in viscous
658 media) and (b) 229 in capacitating media (106 in aqueous media and 123 in viscous
659 media).

661 ***3D Imaging Microscopy***

662
663 Multifocal plane stacks were acquired with the system originally described in Corkidi et al.,
664 2008, consisting on an inverted Olympus IX71 microscope, mounted on an optical table
665 [TMC (GMP SA, Switzerland)], reconfigured with a piezoelectric device P-725 (Physik
666 Instrumente, MA, USA) which periodically displaces a high magnification 100x objective
667 (Olympus UPlanSApo 100x/1.4 na oil objective) at a frequency of 80Hz with a z
668 displacement of 20 μ m. A high-speed camera NAC Q1v (Nac Americas, Inc., USA)
669 acquired images at a rate of 8000 fps with 640 x 480 pixels resolution. Every rising
670 movement of the piezo device (half cycle i.e., 1/160 sec) contains 50 different focal planes

671 (1 image per focal plane). The high-speed camera can store 28,000 images (RAM is 8 Gb)
672 per cell, thus recording spinning motion for a total of 3.5 seconds. Since in this work the
673 rotation of the sperm head is a critical aspect, every possible inversion in each single
674 element conforming the optical-electronical pipeline's path had to be carefully considered
675 (inverted microscope, camera driver setup, image processing software and for
676 visualization -Fiji, Matlab, Paraview, etc.-). As a control test, we have placed in the
677 microscope stage (using a 4x objective) a known pattern (a character R in a piece of
678 paper) viewing the front face of the objective (upside down if seen by the top of the
679 microscope). We have verified that the character appeared upright in the computer screen
680 and that it moved accordingly with the horizontal and vertical stage movements (seen the
681 stage from the bottom-top direction where the objective is placed).

682

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688 **Competing interests**

689 The authors declare that no competing interests exist.

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697 **Author contributions**

698 Conception and design: G.C., F.M., H.G., A.D.; Acquisition of data: G.C., F.M., A-L.G.-C.,
699 Methodology: G.C., F.M., P.H.-H., A-L.G.-C, B.N.C., H.G., A.D.; Software: G.C., F.M.,
700 P.H.-H., B.N.C.; Resources: G.C., A.D.; Writing - review & editing: G.C., F.M., B.N.C.,
701 H.G., A.D.; Supervision, funding and project administration: G.C., A.D.

702 **Ethics**

703 The bioethics committee of the Institute of Biotechnology, UNAM approved the proposed
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708

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712 **Supporting Information**

713 **Invariance of the method to sperm orientation**

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715 To investigate whether the direction of sperm rotation is not an artifact of the illumination

716 setup of the microscope i.e., that the bright border of the sperm head edge is not an effect

717 of the illumination setup, we analyzed the rotation of non-capacitated sperm swimming in

718 four different directions in a Cartesian plane. As we explained previously, in the sequence

719 shown in Figure 6, it is clearly seen that the border of the narrowest part of the head is
720 naturally marked with a bright composed semi-circle. This bright feature turns in the
721 direction of the sperm head (note that this bright border is located behind the sperm head,
722 as explained before). We have verified that independently of the swimming trajectory of
723 sperm, the evolution on time of this bright semi-circle clearly defines the rotating direction
724 of the sperm head. Supplementary Figure S1 shows two consecutive time-points of 4
725 different sperm with outgoing trajectories from the center of each of four Cartesian planes.
726 As can be seen in this figure, the bright region (due to the optical spherical aberration
727 contrast inversion effect) in each sperm always appears first on the **b** side of the head and
728 then it moves to the opposite **b'** side at the subsequent time-point (see Figure 9). This
729 indicates that all the sperm rotate in a CCW (since the bright border is behind the head of
730 the sperm) direction independently of the direction of the sperm trajectory (four quadrants).

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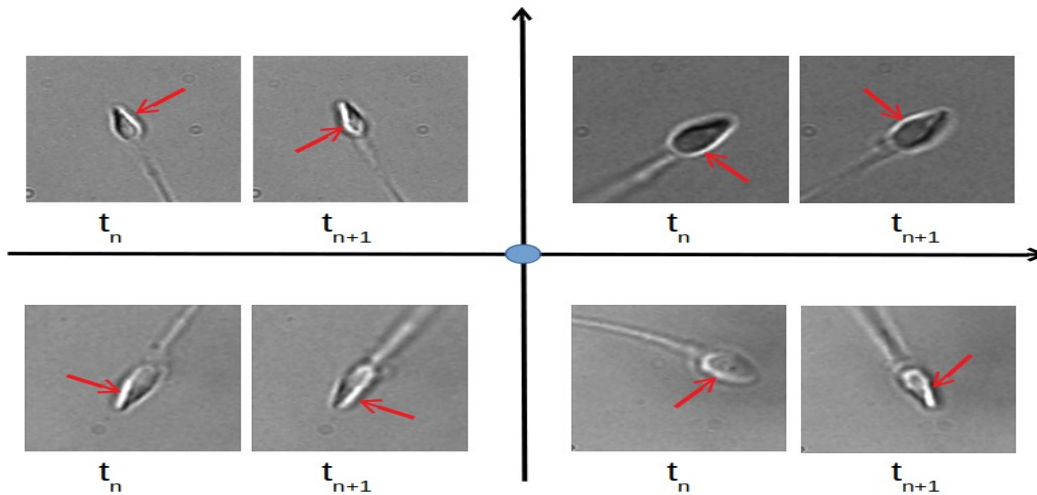
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740 Figure S1. Four different non-capacitated sperm with outgoing trajectories from the center
741 of each of the four Cartesian planes (blue circle). Two different time-points (t_n and t_{n+1}) are
742 shown for each sperm. The bright region (due to the optical spherical aberration contrast
743 inversion effect) always appears first in the left side of the head and then moves to the
744 right side at the subsequent time-point (from head to tail) indicating a CCW head spinning
745 as shown in Results. The red arrows indicate the bright border in each image.