Rotational motion and rheotaxis of human sperm do not require functional CatSper channels and transmembrane Ca\(^{2+}\) signaling

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Abstract

Navigation of sperm in fluid flow, called rheotaxis, provides long-range guidance in the mammalian oviduct. The rotation of sperm around their longitudinal axis (rolling) promotes rheotaxis. Whether sperm rolling and rheotaxis require calcium (Ca\(^{2+}\)) influx via the sperm-specific Ca\(^{2+}\) channel CatSper, or rather represent passive biomechanical and hydrodynamic processes, has remained controversial. Here, we study the swimming behavior of sperm from healthy donors and from infertile patients that lack functional CatSper channels, using dark-field microscopy, optical tweezers, and microfluidics. We demonstrate that rolling and rheotaxis persist in CatSper-deficient human sperm. Furthermore, human sperm undergo rolling and rheotaxis even when Ca\(^{2+}\) influx is prevented. Finally, we show that rolling and rheotaxis also persist in mouse sperm deficient in both CatSper and flagellar Ca\(^{2+}\)-signaling domains. Our results strongly support the concept that passive biomechanical and hydrodynamic processes enable sperm rolling and rheotaxis, rather than calcium signaling mediated by CatSper or other mechanisms controlling transmembrane Ca\(^{2+}\) influx.

Keywords Ca\(^{2+}\) signaling; CatSper; rheotaxis; rolling; human sperm

Subject Categories Cell Adhesion, Polarity & Cytoskeleton; Membranes & Trafficking; Signal Transduction

Introduction

In fluid flow, mammalian sperm realign their swimming path and move upstream—a mechanism called rheotaxis (Miki & Clapham, 2013; El-Sherry et al., 2014; Kantsler et al., 2014; Tung et al., 2014, 2015; Bukatin et al., 2015). In the oviduct, long-range navigation via rheotaxis directs sperm to the site of fertilization (Miki & Clapham, 2013). An important ingredient of rheotaxis is the rotation of sperm around their longitudinal axis, called rolling (e.g., Miki & Clapham, 2013; Kantsler et al., 2014; Bukatin et al., 2015), resulting in a cone-shaped beating envelope. Through this mechanism, vertical shear flow, e.g., near boundary surfaces, exerts a torque that aligns the longitudinal axis of sperm against the flow direction (Miki & Clapham, 2013; Kantsler et al., 2014; Bukatin et al., 2015). However, whether sperm rolling involves full 360° or incomplete rotations of alternating direction is debated (Miki & Clapham, 2013; Muschol et al., 2018).

The intracellular Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_i\)) controls the flagellar beat and swimming behavior of sperm (Kaupp et al., 2003; Publicover et al., 2008; Fechner et al., 2015). In most sperm species, [Ca\(^{2+}\)]\(_i\), is set by the voltage- and alkaline-activated CatSper Ca\(^{2+}\) channel (Quill et al., 2001; Ren et al., 2001; Kirichok et al., 2006; Lishko et al., 2010; Lishko et al., 2011; Strünker et al., 2011; Loux et al., 2013; Seifert et al., 2015). Mammalian CatSper comprises four homologous pore-forming subunits (CatSper 1–4) (e.g., Navarro et al., 2008) and at least six auxiliary subunits (CatSper β, γ, δ, ε, ζ, and Efcab9) (Liu et al., 2007; Wang et al., 2009; Chung et al., 2011, 2017; Hwang et al., 2019). The CatSper-channel complex is organized as quadrilateral threads along the flagellum; the CatSper threads encompass several other proteins, including...
Ca\textsuperscript{2+}-binding proteins and protein kinases, forming local Ca\textsuperscript{2+}-signaling domains near the membrane surface (Chung et al., 2014, 2017).

In \textit{Catsper1\textsuperscript{−/−}} mouse sperm, longitudinal rolling and rheotaxis were abolished (Miki & Clapham, 2013), suggesting that control of [Ca\textsuperscript{2+}], by CatSper is required for rolling and rheotaxis of mammalian sperm. For human sperm, it was specifically proposed that rolling is created by asymmetrical Ca\textsuperscript{2+} influx via CatSper channels, stimulated by local pH\textsubscript{i} signaling (Miller et al., 2018). The H\textsuperscript{+} channel Hv1 is organized along the flagellum of human sperm in two threads near two of the four CatSper threads (Miller et al., 2018). It was proposed that H\textsuperscript{+} influx via Hv1 organizes localized Ca\textsuperscript{2+} signaling that, ultimately, creates an asymmetry in calcium-dependent inhibition of dynein-powered microtubule sliding (Miller et al., 2018). However, the concept that rolling and rheotaxis are enabled by Ca\textsuperscript{2+} influx cannot be reconciled with the finding that rolling of mouse sperm does not require extracellular Ca\textsuperscript{2+} (Babcock et al., 2014; Muschol et al., 2018), and that exposure of human sperm to gradients of flow velocities does not evoke measurable changes in [Ca\textsuperscript{2+}]\textsubscript{i} (Zhang et al., 2016). Moreover, the inventory and regulation of signaling molecules are different among mammalian sperm (Kaupp & Strünker, 2017). For example, mouse sperm lack Hv1 channels (Lishko et al., 2010; Berger et al., 2017), and human CatSper is activated by nanomolar concentrations of prostaglandins and progesterone (Lishko et al., 2011; Strünker et al., 2011) that do not activate mouse CatSper (Lishko et al., 2011). Thus, if the quadrilateral arrangement of CatSper and its control by pH\textsubscript{i} were key to rolling and rheotaxis of mouse and human sperm, the underlying mechanisms ought to be vastly different.

Here, we show that human sperm undergo continuous full 360° rotations rather than incomplete rotations of alternating directions. Moreover, to scrutinize the role of CatSper and Ca\textsuperscript{2+} in rolling and rheotaxis of human sperm, we studied sperm of healthy donors and DIS patients (Fig 2D–G). For optically trapped control sperm from healthy donors, the rotation frequency decreased with increasing viscosity (Fig 2F), in line with previous results (e.g., Nosrati et al., 2015; Gallagher et al., 2019). To study rolling of single sperm cells with high time resolution and for long recording times, we combined bright-field microscopy with an optical tweezer (Ashkin et al., 1986) (Fig 2G). Sperm were trapped perpendicular to the optical axis (Fig 2H, Movie EV2), and the periodic intensity fluctuations of the laser light, which was back-scattered from the cell into the microscope objective, provided a measure of the rotation frequency (Fig 2I). The frequency distribution and mean frequency of trapped sperm (6.0 ± 2.1 Hz, \(n = 32\)) and freely moving sperm (7.0 ± 2.2 Hz, \(n = 1,097\)) were similar (compare Fig 2J and D). Trapping of sperm parallel to the optical axis allowed a frontal view onto the tip of the sperm head; this view reveals that human sperm display continuous full 360° rotations (Fig 2J, Movie EV3), in contrast to incomplete rotations of alternating directions that have been reported for mouse sperm (Babcock et al., 2014; Muschol et al., 2018). Remarkably, also CatSper-deficient human sperm displayed longitudinal rolling (Movie EV4): In freely moving CatSper-deficient sperm incubated under non-capacitating or capacitating conditions, the rotation frequency was normally distributed around a mean value of 6.0 ± 2.6 Hz.
(\(n = 1,009\)) and \(6.8 \pm 3.1\) Hz \((n = 946)\), respectively (Fig 2L and M). The CatSper-deficient human sperm swam progressively also in highly viscous media (Movie EV5), and, like in control sperm, the rotation frequency of CatSper-deficient sperm decreased with increasing viscosity (Fig 2N). When optically trapped, the CatSper-deficient sperm clearly displayed continuous full \(360^\circ\) rotations (Fig 2O and P, Movie EV6), and the rotation frequency remained constant over several tens of seconds (Fig 2Q). In conclusion, human sperm do not require functional CatSper for longitudinal rolling. If anything, the longitudinal rolling of CatSper-deficient sperm might be slightly enhanced in the absence of bicarbonate.
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Longitudinal rolling of human sperm does not require an influx of Ca$^{2+}$

We further examined whether Ca$^{2+}$ is required for rolling of human sperm, using both dark-field microscopy of sperm populations and optical trapping of single sperm cells. Control sperm from healthy donors held by the optical tweezer were dragged between parallel laminar flows of three different solutions (Figs 3A and EV2). This setup allows monitoring of the rotation frequency upon rapid switching of solutions. A stimulus buffer (stimulus stream) and sperm in control buffer (control stream) were separated by a barrier stream containing fluorescein in control buffer; the buffers were fed into a capillary via three inlets. The transfer from one to the other stream was monitored by changes in the fluorescence of fluorescein: When entering the barrier stream, the fluorescence rose and resumed basal values when sperm reached the stimulus stream (Fig 3B). Dragging of control sperm from healthy donors across the barrier stream was completed within ≤ 10 s (Movie EV7). Dragging itself did not affect rolling (Fig 3B and C): The mean rotation frequency before and after dragging between control buffers was 6.7 ± 2.8 Hz and 6.5 ± 2.8 Hz (n = 14), respectively. After dragging from bicarbonate-free to bicarbonate-containing buffer, the rotation frequency increased from 6.6 ± 2.9 to 11.3 ± 2.5 Hz (Fig 3D and E, n = 5). Next, the rotation frequency of trapped sperm cells before and after transition from 2 mM to < 20 mM extra-cellular Ca$^{2+}$ was studied. The rotation frequency was similar in the absence and presence of Ca$^{2+}$ (5.5 ± 3.6 Hz versus 5.9 ± 3.1 Hz, n = 5; Fig 3F and G). Although in dark-field microscopy of sperm populations, the fraction of motile sperm decreased in Ca$^{2+}$-free buffer with a time constant (t) of 5.3 min (Fig 3H) (Aaberg et al., 1989; Jin et al., 2007; Torres-Flores et al., 2011), at any time-point during the decay, motile sperm were also rolling (Movie EV8). The
mean rotation frequency and the rotation frequency-histogram (determined at ≤ 5 min in Ca²⁺-free buffer) were similar to those under control conditions (6.3 ± 1.9 Hz, n = 224; Fig 3I). These results show that Ca²⁺ influx is not required for rolling of human sperm.

Next, we studied the swimming behavior of human sperm in a glass capillary with and without fluid flow. Sperm were tracked in the field of view, and the starting point of each track was shifted to the origin.
of a coordinate system (Fig 4A, C, E, G, I, K). To quantify the rheotactic behavior, we determined the angular swimming directions and plotted the mean relative frequency of sperm swimming with angular directions of 45°–135°, 135°–225°, 225°–315°, and 315°–45° in a spider plot. Under no-flow conditions, control sperm swam randomly without any preferred directional angle (Fig 4A and B). In contrast, under flow conditions, a large fraction of sperm aligned their swimming path against the flow direction (Fig 4C and D, angular direction of the flow = 0°). The fraction of sperm swimming with directional angles between 135° and 225° was 25.7 ± 1.7% (n = 7, 1,301 sperm) and 44.3 ± 8.6% (n = 7, 1,083 sperm) in the absence and presence of a flow, respectively. For CatSper-deficient sperm under no-flow conditions, the angular swimming directions were random (Fig 4E and F; Movie EV9). Under flow conditions, like in control sperm, a large fraction of the CatSper-deficient sperm aligned their swimming path against the flow direction (Fig 4G and H; Movie EV10). The fraction of CatSper-deficient sperm swimming with directional angles between 135° and 225° was 27.6 ± 1.8% (no-flow; n = 6, 1,068 sperm) versus 47.8 ± 3.4% (flow; n = 7, 1,068 sperm). These results demonstrate that functional CatSper channels are dispensable not only for rolling, but also for rheotaxis of human sperm.
Rheotaxis of human sperm does not require Ca\textsuperscript{2+} influx

Finally, we studied the trajectories of CatSper-deficient sperm in Ca\textsuperscript{2+}-free buffer ([Ca\textsuperscript{2+}] < 20 nM). Under no-flow conditions, the angular swimming directions were random (Fig 4I and J). Under flow conditions, like in the presence of extracellular Ca\textsuperscript{2+}, a large fraction of the CatSper-deficient sperm aligned their swimming path against the flow direction (Fig 4K and L); in Ca\textsuperscript{2+}-free buffer, the fraction of CatSper-deficient sperm swimming with directional angles between 135° and 225° was 28.2 ± 2.7% (no-flow; n = 4, 442 sperm) versus 43.3 ± 3.6% (flow; n = 4, 620 sperm). These results demonstrate that Ca\textsuperscript{2+} influx in general is dispensable for both rolling and rheotaxis of human sperm and that rheotaxis of CatSper-deficient human sperm is similar in the absence and presence of extracellular Ca\textsuperscript{2+}.

Figure 4.
Figure 4. Rheotaxis of human sperm.
A Trajectories of human sperm in the absence of a fluid flow. Sperm were tracked for 1.86 s. The starting point of each trajectory was centered to the origin of a coordinate system, represented by the intersection of the dotted lines in the center of the circle.
B Spider-web plot of the mean (± SD) relative frequencies of sperm swimming with an angular direction of 315°–45°, 45°–135°, 135°–225°, and 225°–315° (n = 7; 1,301 sperm) in the absence of a fluid flow.
C Representative trajectories of human sperm in the presence of a fluid flow.
D Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions (n = 7; 1,083 sperm) in the presence of a fluid flow. The red arrow indicates the flow direction.
E Representative trajectories of CATSPER2−/− sperm in the absence of a fluid flow.
F Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions of CATSPER2−/− sperm in the absence of a fluid flow (n = 6; 1,068 sperm).
G Trajectories of CATSPER2−/− sperm in the presence of a fluid flow.
H Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions of CATSPER2−/− sperm in the presence of a fluid flow (n = 7; 1,068 sperm).
The red arrow indicates the fluid flow direction.
I Representative trajectories of CATSPER2−/− sperm in Ca2+-free buffer in the absence of a fluid flow.
J Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions (n = 4; 442 sperm) of CATSPER2−/− sperm in Ca2+-free buffer in the absence of a fluid flow.
K Representative trajectories of CATSPER2−/− sperm in Ca2+-free buffer in the presence of a fluid flow.
L Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions (n = 4; 620 sperm) of CATSPER2−/− sperm in Ca2+-free buffer in the presence of a fluid flow. The red arrow indicates the flow direction.

Rolling and rheotaxis persist in CatSper-deficient mouse sperm

Although both devoid of functional CatSper channels, rolling and rheotaxis are largely unaffected in human CATSPER2−/− sperm, but seem to be abolished in mouse Catsper1−/− sperm (Miki & Clapham, 2013). A major difference is that the quadrilateral Ca2+-signaling threads are disrupted in Catsper1−/− mouse sperm (Chung et al., 2014), but not in CATSPER2−/− human sperm (Fig 1G). This finding suggests that the supramolecular CatSper organization might be required for rolling and rheotaxis. To test for this possibility, we re-examined rolling and rheotaxis in mouse Catsper1−/− sperm. Surprisingly, not only wild-type (Fig 5A, Movie EV11) but also Catsper1−/− sperm (Fig 5B, Movie EV12) clearly displayed longitudinal rolling. The mean rotational frequency of wild-type and Catsper1−/− sperm was 2.9 ± 1.3 Hz (n = 24) and 2.6 ± 0.7 Hz (n = 24), respectively (Fig 5A and B). We studied the swimming behavior of wild-type and Catsper1−/− sperm in a glass capillary with and without fluid flow. We tracked sperm in the field of view, and the starting point of each track was shifted to the origin of a coordinate system (Fig 5C, E, G, I).

Figure 5. Rolling behavior and rheotaxis of mouse sperm.
A Left: Bright-field image series of a freely swimming wild-type sperm cell at t = 0, 151, and 303 ms. Scale bar represents 10 μm. Right: rotation frequency (mean ± SD) of freely swimming wild-type sperm (n = 24, three experiments).
B Left: Bright-field image series of a freely swimming Catsper1−/− sperm cell at t = 0, 151, and 294 ms. Scale bar represents 10 μm. Right: rotation frequency (mean ± SD) of freely swimming Catsper1−/− sperm (n = 24, three experiments).
C Representative trajectories of wild-type sperm in the absence of a fluid flow. The starting point of each trajectory was centered to the origin of a coordinate system, represented by the intersection of the dotted lines in the center of the circle. Each color represents one trajectory.
D Spider-web plot of the mean (± SD) relative frequencies of sperm swimming with an angular direction of 315°–45°, 45°–135°, 135°–225°, and 225°–315° (n = 3; 127 sperm) in the absence of a fluid flow.
E Representative trajectories of wild-type sperm in the presence of a fluid flow.
F Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions (n = 3; 175 sperm) of wild-type sperm in the presence of a fluid flow. The red arrow indicates the flow direction.
G Representative trajectories of Catsper1−/− sperm in the absence of a fluid flow. The starting point of each trajectory was centered to the origin of a coordinate system, represented by the intersection of the dotted lines in the center of the circle. Trajectories are magnified by a factor of 2.05 with respect to the plots C and E to compensate for the reduced swimming speed of the Catsper1−/− sperm.
H Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions of Catsper1−/− sperm (n = 4; 297 sperm) in the absence of a fluid flow.
I Representative trajectories of Catsper1−/− sperm in the presence of a fluid flow; trajectories are magnified by a factor of 2.05 with respect to the plots C and E to compensate for the reduced swimming speed of the Catsper1−/− sperm, and two trajectories were truncated (indicated by two parallel lines).
J Spider-web plot of the mean (± SD) relative frequencies of angular swimming directions (n = 4; 261 sperm) of Catsper1−/− sperm in the presence of a fluid flow. The red arrow indicates the flow direction.
do not require functional CatSper channels. For the time being, we cannot reconcile ours with previous results (Miki & Clapham, 2013). We suggest that other laboratories examine rolling and rheotaxis of CatSper-deficient mouse sperm independently.

Discussion

The Ca²⁺ channel CatSper has been implicated in rolling and rheotactic steering of mammalian sperm (Miki & Clapham, 2013;
Miller et al., 2018). We show that, in fact, rolling and rheotaxis of both human and mouse sperm do not require Ca\(^{2+}\) influx via CatSper. This conclusion agrees with other reports: Rolling of mouse sperm does not require extracellular Ca\(^{2+}\) (Babcock et al., 2014; Muschol et al., 2018), and exposure of human sperm to gradients of flow velocities does not evoke measurable changes in \([\text{Ca}^{2+}]_{i}\) (Zhang et al., 2016). Furthermore, it has been proposed that the quadrilateral organization of CatSper and associated signaling components provides the flagellar ultrastructure required for rolling and rheotaxis (Miller et al., 2018). The CatSper complex and the quadrilateral Ca\(^{2+}\)-signaling domains are abolished in Catsper1\(^{-/-}\) mouse sperm (Chung et al., 2014), but not in human CATSPER2\(^{-/-}\) sperm. Yet, both sperm species display rheotaxis and undergo rolling, suggesting that also the quadrilateral flagellar architecture along with potential asymmetric cytosolic Ca\(^{2+}\) gradients established by this structure is dispensable. Furthermore, mouse sperm lack Hv1 channels (Lishko et al., 2010; Berger et al., 2017), indicating that asymmetrical, spatially confined pH gradients established by Hv1 are not required for rolling and rheotaxis. Altogether, our results strongly support the concept that passive biomechanical and hydrodynamic processes enable rolling and rheotaxis rather than active spatio-temporally confined Ca\(^{2+}\) and H\(^{+}\) signaling.

On a broader perspective, rolling is not a unique feature of sperm that undergo rheotaxis: Sperm from marine external fertilizers also exhibit longitudinal rolling (Jikeli et al., 2015), although in their aquatic habitat, no gradients of fluid velocity exist and chemotaxis rather than rheotaxis is employed for navigation.

Of note, even in a viscous medium, rheotaxis of CatSper-deficient and healthy human sperm was rather similar (Fig 6A and B). Thus, at the particular flow velocity and viscosity that we used, and with respect to the parameters that we analyzed, the lack of functional CatSper does not seem to affect rheotaxis in more viscous fluid.

However, human sperm display rheotaxis within a broad range of physiological flow velocities and viscosities (Kantsler et al., 2014). The relation of upstream versus shear velocity is bell-shaped (Kantsler et al., 2014), demonstrating that the rheotactic performance peaks at a particular shear profile. Furthermore, the trajectories of human sperm swimming against a flow feature a transverse component, and in cylindrical tubes, sperm swim on spiral-shaped trajectories along the walls, thereby exploring the tube’s surface (Kantsler et al., 2014). The transversal component is positively and negatively related to the shear velocity and viscosity, respectively. Thus, subtle differences in the 3D beat of CatSper-deficient sperm might compromise rheotactic performance under some conditions encountered in the oviduct. For example, rheotaxis might be compromised at certain shear velocities and/or fluid viscosities, the relationship of upstream versus shear velocity might be shifted, and/or the transversal component might be altered. The beat envelope is a critical determinant of the rheotactic performance: Mouse sperm that lack CatSper suffer from a rather stiff proximal flagellum, altering the 3D flagellar envelope and hampering the reorientation in fluid flow (Chung et al., 2017). Future studies need to quantify the 3D flagellar beating pattern of control versus CatSper-deficient human sperm as well as their rheotactic performance over a broad range of fluid flows and viscosities.

Furthermore, in human sperm, CatSper translates stimulation with oviductal hormones, like steroids and prostaglandins (Lishko et al., 2011; Strünker et al., 2011; Brenker et al., 2012, 2018a; Miller et al., 2016; Mannowitz et al., 2017), into Ca\(^{2+}\) and motility responses that are important for human sperm chemotaxis and hyperactivation (Schaefer et al., 1998; Harper et al., 2003; Oren-Benaroya et al., 2008; Publicover et al., 2008; Baldi et al., 2009; Kilic et al., 2009; Alasmari et al., 2013; Schiffer et al., 2014; Tamburrino et al., 2014, 2015; Rennhack et al., 2018). In the absence and
presence of progesterone (100 nM), rheotaxis of control and CatSper-deficient human sperm was rather similar (Fig 6C and D), indicating that under our conditions, progesterone activation of CatSper does not affect rheotaxis. However, to decipher how the ligand control of CatSper and, thereby, [Ca$^{2+}$], are intertwined with rheotaxis, it is required to quantify rheotaxis of control and CatSper-deficient human sperm in the absence and presence of picomolar progesterone concentrations over a broad range of flow velocities and viscosities that emulate in vitro the complex physico-chemical landscape of the oviduct.

On a final note, it is unknown whether sub- or infertility in men correlates with the failure of sperm to undergo rheotaxis. Studying rheotaxis is technically demanding. Therefore, we propose to assess longitudinal rolling as a surrogate biomarker for infertility. The population analysis introduced in this study can be readily incorporated into existing computer-assisted sperm analysis setups for clinical diagnostics.

Materials and Methods

Reagents

Reagents were obtained from Sigma-Aldrich (USA) unless otherwise indicated.

Sperm preparation and buffer conditions

The studies were performed in accordance with the standards set by the Declaration of Helsinki. Samples of human semen were obtained from healthy volunteers and DIS patients with their prior written consent, under approval of the institutional ethical committees of the medical association Westfalen-Lippe and the medical faculty of the University of Münster (4INie). Ejaculates were allowed to liquefy at 37°C for 30–60 min. Motile sperm were purified by a “swim-up” procedure: Liquefied semen (0.5–1 ml) was layered in a 50-ml Falcon tube below 4 ml of human tubal fluid (HTF) medium containing (in mM): 93.8 NaCl, 4.69 KCl, 0.2 MgSO4, 0.37 KH2PO4, 2.04 CaCl2, 0.33 Na-pyruvate, 21.4 lactic acid, 2.78 glucose, 4 NaHCO3, and 21 HEPES, pH 7.35 (adjusted with NaOH). Alternatively, the semen was diluted 1:10 with HTF and sperm, somatic cells, and cell debris were pelleted by centrifugation at 700 g for 20 min (37°C). The pellet was resuspended in the same volume HTF, 50-ml Falcon tubes were filled with 5 ml of the suspension, and cells were pelleted by centrifugation at 700 g for 5 min (37°C). In either case, motile sperm were allowed to swim up into HTF for 60–90 min at 37°C. After swim-up, sperm were washed twice (700 g, 20 min) with HTF, the sperm concentration was adjusted, and HTF was supplemented with 3 mg/ml human serum albumin (HSA, Scientific Irvine, USA; referred to as HTF++); under these conditions, sperm are non-capacitated. For capacitation, sperm were resuspended after the second wash in HTF++ medium, containing (in mM): 72.8 NaCl, 4.69 KCl, 0.2 MgSO4, 0.37 KH2PO4, 2.04 CaCl2, 0.33 Na-pyruvate, 21.4 lactic acid, 2.78 glucose, 25 NaHCO3, and 21 HEPES, pH 7.35 (adjusted with NaOH), and supplemented with 3 mg/ml HSA. Sperm were capacitated in HTF+++ for at least 3 h. Alternatively, swim-up and washing were directly performed in HTF++. To study non-capacitated sperm in the absence of bicarbonate and the motility response evoked by a step increase in bicarbonate, swim-up and washing were performed in bicarbonate-free HTF containing (in mM): 97.8 NaCl, 4.69 KCl, 0.2 MgSO4, 0.37 KH2PO4, 2.04 CaCl2, 0.33 Na-pyruvate, 21.4 lactic acid, 2.78 glucose, and 21 HEPES, pH 7.35 (adjusted with NaOH). HSA (3 mg/ml) was added prior to the experiment to prevent attaching of sperm to the surface of the recording chamber. C57BL/6 wild-type and CatSper1−/− mice were handled and sacrificed in accordance with the German Animal Welfare Act and the district veterinary office under approval by the LANUV (84-02.05.13.115). Mouse epididymis was obtained from at least 21-week-old male mice that were anaesthetized with CO2 or isoflurane (AbbVie Deutschland, Ludwigshafen, Germany) and sacrificed by cervical dislocation. Mouse sperm were isolated by incision of the cauda epididymis in modified TYH medium containing (in mM): 138 NaCl, 4.8 KCl, 2 CaCl2, 2.9 KH2PO4, 1 MgSO4, 5.6 glucose, 0.5 sodium pyruvate, 10 sodium isocitrate, and 21 HEPES, pH 7.4. Sperm were allowed to swim out for 30 min at 30°C and 10% CO2. CatSper1−/− mice (Ren et al, 2001) were generously provided by David Clapham (Janelia Research Campus, USA).

Rolling analysis in sperm populations

The longitudinal rolling of human sperm was recorded in glass chambers (depth of ~100 µm) under an inverted microscope (IX73; Olympus, Germany), equipped with a condenser (IX2-LWUCD; Olympus, Germany) with a custom-made dark-field filter, a 10× objective (UPLFLN10X2PH1; Olympus, Germany), and additional 1.6× magnification lenses (16× final magnification). The samples were illuminated with a red light-emitting diode (LED; M660D2; Thorlabs, Germany) and a custom-made power supply. To study sperm rolling in bicarbonate-free HTF or HTF++, sperm were diluted 1:9 in the respective buffer 5–30 min prior to the experiment. To study rolling in the absence of extracellular Ca$^{2+}$ ([Ca$^{2+}$]o < 20 nM), sperm in HTF++ were diluted 1:9 only prior to the experiment into a Ca$^{2+}$-free HTF medium (HTF−C), containing (in mM): 69.8 NaCl, 4.69 KCl, 0.2 MgSO4, 0.37 KH2PO4, 5 EGTA, 0.33 Na-pyruvate, 21.4 lactic acid, 2.78 glucose, 25 NaHCO3, and 21 HEPES, pH 7.35 (adjusted with NaOH), and supplemented with 3 mg/ml HSA. To study sperm rolling in viscous medium, sperm in HTF++ were diluted in HTF++ fortified with methyl cellulose. Over 4–10 min, short movies (~725 ms) of sperm in different fields of view in the observation chamber were recorded at 124 Hz with a high-speed sCMOS camera (Zyla, 4.2 plus, Andor, UK). Longitudinal rolling was assessed with a custom-made program written in the ImageJ macro language (Rasband, 1997–2016). In brief, moving sperm heads were tracked, and rotation was monitored by an oscillating change in head brightness. The rotation frequency was computed from the average temporal distance between two intensity peaks. The sperm head is approximately plane symmetrical with planes intersecting the length axis and, therefore, lights up twice per 360° rotation. Thus, the rotation frequency (Frot) is given by Frot = Flight × 0.5. Sperm that displayed less than three relative maxima within the observation time were excluded from the analysis, yielding a cut-off for Frot of about 1.5 Hz; immotile sperm were excluded from the analysis. The rolling of mouse sperm in TYH was studied in observation chambers (depth of ~400 µm; Ibidi, Germany) under an
inverted microscope (IX73; Olympus, Germany) equipped with a condenser (IX2-LWUCD; Olympus, Germany), a 10× objective (UPLFLN10X2PH1; Olympus, Germany), and, optionally, an additional 1.6× magnification lens. The sample was illuminated by a red LED (M660D2; Thorlabs, Germany). Movies of sperm were recorded with a high-speed sCMOS camera (Zyla, 4.2 plus; Andor, UK). The rolling frequency of sperm was determined by visual frame-by-frame analysis.

**Optical trapping of sperm**

Unless otherwise indicated, we used capacitated sperm in HTF++ for optical-trapping experiments. The trapping of sperm cells was achieved with an optical tweezer (Ashkin et al., 1986; Fig EV2), using a continuous-wave (cw) diode laser (Lumics Lu0975M500, Germany) at a wavelength of 976 nm (red beam in Fig EV2). The laser beam was expanded by a telescope setup of two lenses (L) to a diameter of 1.7 mm (full width at half maximum) and directed into a 100× oil immersion objective (MO, Plan-Apochromatic 100×/1.40 Oil DIC M27, Zeiss, Germany) with a numerical aperture (NA) of 1.4, allowing for tight focusing of the laser beam onto the head of a sperm swimming inside an observation capillary. To assess the rotation frequency of trapped sperm, the laser light reflected by the sperm head into the objective was directed onto a photomultiplier tube (PMT, H10721-20, Hamamatsu, Japan) by a beam splitter (reflectivity ~4%). A long-pass filter in front of the PMT blocked both the bright-field illumination and ambient light. The PMT signal was sampled with a frequency of 2 kHz, and every 10 points were averaged. The rotation frequency was determined by a fast Fourier transformation in a moving time window of 1.5 s. In parallel to the quantification of the back-reflected laser light, we recorded the trapped sperm with a bright-field microscope. A blue LED, equipped with a 450-nm short-pass filter (SP) and a collimator lens, served as a light source (blue beam in Fig EV2). The bright-field image was reflected by a dichroic mirror (DM), projected by a lens onto the chip of a charged-coupled device (CCD) camera (UI-3140CP-M-GL, IDS, Germany) and recorded with a frame rate of 200 Hz. To measure the rotation frequency of a sperm at different conditions, we trapped sperm inside a microfluidic capillary (dimensions [height × width]: 0.4 × 0.33 μm; μ-Slide III 3in1; Ibidi, Germany) with three separate inlets to establish a continuous, parallel laminar flow of three solutions ((i) control stream with sperm, (ii) barrier stream, and (iii) stimulus stream) with a flow speed of 65 μm/s. The barrier stream was supplemented with fluorescein (1 μM). Fluorescein was excited with the blue LED; fluorescence light was collected through the microscope objective and reflected by two DMs through a long-pass filter onto a second PMT (H10721-210; Hamamatsu, Japan). Using a custom-built mechanical scanning table, the microfluidic capillary was moved in the horizontal plane orthogonal to the flow direction, dragging a trapped sperm within 6.8 s from the control stream through the barrier stream into the stimulus stream. The fluorescein fluorescence, recorded synchronously to the back-reflected laser light and the bright-field images, provided a readout of the position of the trapped sperm inside the flow profile. For control experiments, the control stream with sperm, the barrier stream, and the stimulus stream consisted of HTF++. To study the action of Ca²⁺, the stimulus stream consisted of HTF³Ca²⁺. To study the action of bicarbonate, the control stream with sperm consisted of bicarbonate-free HTF. For paired-plot analysis, the change in frequency was determined after reaching a stable value.

**Rheotaxis assay**

Human sperm in HTF++, in HTF++ containing 100 nM progesterone, or in HTF++ containing 0.2% methylcellulose, or mouse sperm in TYH were observed in shallow microfluidic channels with rectangular cross section of 0.4 × 3.8 mm (Ibidi, Germany) under an inverted microscope (IX73; Olympus, Germany) equipped with a condenser (IX2-LWUCD; Olympus, Germany) and a 10× objective (UPLFLN1X2PH1; Olympus, Germany). The sample was illuminated by a red LED (M660D2; Thorlabs, Germany). Images were collected at ~ 80 to ~ 125 Hz using a sCMOS camera (Zyla 4.2 Plus; Andor, UK). Sperm were exposed to a buffer flow of ~ 13.5 μl/min (human) or ~ 10 μl/min (mouse), respectively, using a syringe pump (World Precision Instruments, USA). Individual human sperm were tracked over 1.86 s in a semi-automatic fashion using a custom-made tracking tool based on the Mtrack2 plugin for ImageJ (NHI, Bethesda, USA). The angle of each track was defined by its start and end point in a two-dimensional Cartesian coordinate system with the flow direction pointing to an angle of 0° and a trajectory straight against the flow pointing to an angle of 180°. Immotile and surface-attached sperm were excluded from analysis. For each experiment, computed trajectory angles were binned into angle intervals of 90° and expressed as fractions of the sperm population (e.g., Fig 4B). Individual mouse sperm were tracked manually for 2 s; the angle of each track was defined by its start and end point in a two-dimensional Cartesian coordinate system with the flow direction pointing to an angle of 0° and a trajectory straight against the flow pointing to an angle of 180°. Like for human sperm, immotile and surface-attached sperm were excluded from analysis. For each experiment, computed trajectory angles were binned into angle intervals of 90° and expressed as fractions of the sperm population (e.g., Fig 4D).

**Measurement of changes in [Ca²⁺]**

Changes in [Ca²⁺] were measured in sperm (in HTF++) loaded with the fluorescent Ca²⁺ indicator Fluo-4-AM at 30°C in 384 multi-well plates in a fluorescence plate reader (Fluostar Omega, BMG Labtech, Ortenberg, Germany) at 30°C as described before (Schiffer et al., 2014; Brenker et al., 2018b). Briefly, sperm were loaded with Fluo-4-AM (5 μM, 20 min) at 37°C in the presence of Pluronic F-127 (0.05% v/v). After incubation, excess dye was removed by centrifugation (700 g, 5 min, room temperature). Sperm were resuspended in HTF at a density of 5 × 10⁶ cells/ml. The wells were filled with 54 μl of the sperm suspension; fluorescence was excited at 480 nm (Fluo-4), and fluorescence emission was recorded at 520 nm. Changes in Fluo-4 fluorescence are depicted as ΔF/F₀ (%), that is, the change in fluorescence (ΔF) relative to the mean basal fluorescence (F₀) before application of buffer or stimuli (6 μl).

**Electrophysiology**

We recorded from sperm in the whole-cell configuration as described before (Strünker et al., 2011). Seals between pipette and sperm were formed either at the cytoplasmic droplet or in the neck
region in standard extracellular solution (HS) containing (in mM): 135 NaCl, 5 KCl, 1 MgSO4, 2 CaCl2, 5 glucose, 1 Na-pyruvate, 10 lactic acid, and 20 HEPES, adjusted to pH 7.4 with NaOH. Monovalent currents were recorded in a sodium-based divalent-free solution (NaDVF) containing (in mM): 140 NaCl, 40 HEPES, and 1 EGTA, adjusted to pH 7.4 with NaOH; the pipette (10–15 MΩ) solution contained (in mM): 130 Cs-aspartate, 5 CsCl, 50 HEPES, and 5 EGTA, adjusted to pH 7.3 with CsOH. Data were not corrected for liquid junction potentials.

**Immunocytochemistry**

Sperm were immobilized on microscope slides and fixed for 10 min with paraformaldehyde in PBS (4%; PBS containing [in mM]: 137 NaCl, 2.7 KCl, 10 Na2HPO4, 1.8 KH2PO4, pH 7.4–7.5). To block unspecific binding sites, sperm were incubated for 1 h with blocking buffer (0.5% Triton-X 100 and 5% ChemiBLOCKER [Millipore, USA] in 0.1 M PBS, pH 7.4). Primary antibodies (anti-CatSper 4, ACC-304, Alomone Labs, Israel; polyclonal antibody raised in rabbits, directed against amino acids 384–402 of CatSper 3) were diluted in blocking buffer and incubated overnight. Fluorescent secondary antibodies were diluted in blocking buffer containing 0.5 mg/ml DAPI (Invitrogen, USA), and pictures were taken with a confocal microscope (FV1000; Olympus, Japan).

**3D-STORM microscopy**

Experiments were performed with a Ti-E microscope (Nikon, Japan) in an imaging buffer (50 mM Tris, pH 8, 10 mM NaCl) with an oxygen scavenging system (0.5 mg/ml glucose oxidase, 40 μg/ml catalase [Roche Applied Science, Germany or Sigma-Aldrich], and 10% [w/v] glucose), and 10 mM 2-aminoethanethiol. Images were acquired with an iXON 897 EMCCD camera (Andor, UK). 10,000–60,000 frames were acquired per data set using a 647-nm excitation laser at 100 mW at the sample plane, unless mentioned otherwise. A 405-nm laser was used to maintain an adequate number of localizations per frame. For 3D STORM acquisition, a cylindrical lens was introduced into the detection path; the “perfect focus system” (Nikon) and a vibration isolation table were used to minimize axial and lateral drifting, respectively. STORM movies were analyzed as described previously using the Nikon software package based on a technology developed by Dr. Xiaowei Zhuang (Huang et al, 2008). Briefly, fluorescence peaks corresponding to individual molecules were identified in each frame and fit, using least-squares fitting or maximum-likelihood estimator fitting, with a two-dimensional Gaussian to determine the (x,y) position of each molecule. For 3D imaging, the ellipticity of the Gaussian was used to assign the z coordinate. Drift correction was applied using cross-correlation. STORM images were rendered with each localization plotted as a Gaussian whose width is weighted by the inverse square root of the number of detected photons for that switching event. Images were filtered to reject molecules with emitted photon number below 500. Molecules with an aspect ratio higher than 1.5 for 2D and 2.5 for 3D datasets were rejected. Moreover, molecules that appeared for more than 10 consecutive frames were rejected. Background noise in STORM images caused by non-specifically bound antibodies, appearing as scattered localizations at low local densities, was removed by a local density filter. Low-density localizations were filtered out by removing a localization if it was surrounded by fewer than 10 localizations in the 80 nm × 80 nm region surrounding the localization.

**Array CGH analysis**

Patients were analyzed by array comparative genomic hybridization (CGH; Agilent platform, Agilent Technologies, Santa Clara, California, USA) using 400k arrays (#G4448A). For details, see Tüttelmann et al (2011).

**Statistical methods**

Unless otherwise indicated, data are displayed as mean ± SD.

**Expanded View** for this article is available online.

**Acknowledgements**

This work was supported by the German Research Foundation (CRU326 to T.S. and F.T.) and the Cells-in-Motion (CiM) Cluster of Excellence, Münster (FF-2016-17 to T.S. and C.F.).

**Author contributions**

CB, CF, and TS conceived the project. CS, SR, CB, SY, HH, DW, FT, AR, UBK, TW, AW, CK, SK, CF, and TS designed research, performed experiments, acquired, and/or interpreted data. CS and TS wrote the manuscript. All authors revised the manuscript critically for important intellectual content and approved the manuscript.

**Conflict of interest**


**References**


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