Loss of the Na\(^{+}/\)H\(^{+}\) exchanger NHE8 causes male infertility in mice by disrupting acrosome formation

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Mammalian sperm feature a specialized secretory organelle on the anterior part of the sperm nucleus, the acrosome, which is essential for male fertility. It is formed by a fusion of Golgi-derived vesicles. We show here that the predominantly Golgi-resident Na\(^{+}/\)H\(^{+}\) exchanger NHE8 localizes to the developing acrosome of spermatids. Similar to wild-type mice, Nhe8\(^{-/-}\) mice generated Golgi-derived vesicles positive for acrosomal markers and attached to nuclei, but these vesicles failed to form large acrosomal granules and the acrosomal cap. Spermatozoa from Nhe8\(^{-/-}\) mice completely lacked acrosomes, were round-headed, exhibited abnormal mitochondrial distribution, and displayed decreased motility, resulting in selective male infertility. Of note, similar features are also found in globozoospermia, one of the causes of male infertility in humans. Germ cell-specific, but not Sertoli cell-specific Nhe8 disruption recapitulated the globozoospermia phenotype, demonstrating that NHE8’s role in spermiogenesis is germ cell-intrinsic. Our work has uncovered a crucial role of NHE8 in acrosome biogenesis and suggests that some forms of human globozoospermia might be caused by a loss of function of this Na\(^{+}/\)H\(^{+}\) exchanger. It points to NHE8 as a candidate gene for human globozoospermia and a possible drug target for male contraception.

Spermatogenesis is the process in which spermatozoa, the male gametes, are produced. After cycles of mitosis and meiosis, the haploid spermatids undergo drastic morphological changes to transform from a spherical cell into a polarized, elongated spermatozoon in a process called spermiogenesis (1). In this process, a large, mainly Golgi-derived membrane-enclosed structure, the acrosome, is formed on the anterior part of the sperm head. The acrosome plays a crucial role during fertilization. Mice and men in which all spermatozoa lack acrosomes are infertile (2, 3).

The Na\(^{+}/\)H\(^{+}\) exchanger (NHE)\(^{2}\) NHE8 is an integral membrane protein with 12 transmembrane domains as suggested by structures of bacterial homologues (4). NHE8 (SLC9A8) belongs to the SLC9A gene family that comprises nine members in mammals (5). These can be divided into two subfamilies: a plasma membrane subfamily including NHE1–5 and an organellar subfamily comprising NHE6–9. Plasma membrane NHEs are involved in the absorption of salt and water in epithelia and in regulating cytoplasmic pH, osmolarity, and cell volume. Knock-out mouse models have implicated these transporters in disorders like epilepsy and seizures, diabetes, or metabolic acidosis (6–9). Organellar NHEs may regulate luminal ion concentrations, in particular the pH, of organelles in the secretory and endocytic pathways. These organelles display distinct, mostly acidic, luminal pH, which is crucial for organellar function (10–12). The specific pH of each organelle is determined by a balance of V-ATPase-mediated proton pumping and a “proton leak” pathway (13–15). Several studies in yeast (16) and mammalian cells (14, 15, 17–19) demonstrated that overexpression or knockdown of organellar NHEs leads to alkalization or acidification of the organellar lumen, respectively, suggesting that these transporters provide a proton leak. The importance of organellar NHEs is underscored by mutations in NHE6 and NHE9 that lead to neurological disorders such as autism and epilepsy in humans (20, 21).

NHE8 (SLC9A8) is widely expressed in murine tissue, with high mRNA levels found in kidney, liver, skeletal muscle, and testes (17, 22). NHE8 localizes predominantly to mid- to trans-Golgi stacks, where it may alkalinize the luminal pH (17). NHE8 was also reported to be present on multivesicular bodies (23) and somewhat surprisingly at the apical brush-border membrane of epithelial cells of the intestine and renal proximal tubules (22, 24, 25). Nhe8\(^{-/-}\) mice show impaired colonic mucin synthesis and bicarbonate secretion (26, 27), a dysfunction of the retinal pigment epithelium (28, 29), and male infertility (28, 30). In addition to NHE8, two other NHEs are known to affect male fertility in mice. Loss of plasma membrane NHE3 (SLC9A3) leads to increased fluid volume in the epididymis, thereby diminishing sperm concentration (31). Loss of the so-called sperm NHE (sNHE, a member (SLC9C1) of the SLC9C gene family) drastically reduces sperm motility and causes male infertility (32). sNHE, which contains additional voltage-sensor and nucleotide binding domains, localizes to the flagellum of mouse spermatozoa, where it may participate in a signaling complex at the plasma membrane (33).

Here we analyzed the role of NHE8 in male reproductive function. Mice lacking NHE8 globally or specifically only in...
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germ cells produced round-headed spermatozoa lacking the acrosomal cap. This resembles human globozoospermia which likewise is associated with infertility. Instead of acrosomes, Nhe8–/– spermatids displayed isolated, sometimes nucleus-attached Golgi-derived vesicles (GDVs). Hence NHE8 may play a crucial role in the fusion of GDVs to form acrosomes. Our work suggests NHE8 as a candidate gene for human globozoospermia and as a possible drug target for male contraception.

Results
Disruption of Nhe8 causes male infertility in mice
We established Nhe8–/– mice (Fig. 1A) to elucidate physiological roles of NHE8. Wild-type (WT), Nhe8+/–, and Nhe8–/– mice were born at Mendelian ratio (Fig. 1H), grew normally, and all littersmates were superficially indistinguishable. Complete loss of NHE8 protein in Nhe8–/– mice was confirmed in several organs by immunoblot analysis which in the WT showed NHE8 signals of ~60 and 85 kDa as reported (22) (Fig. 1B). NHE8 expression was prominent in kidney, testes, and epididymis, but very low in the female reproductive system (Fig. 1B). Although we frequently obtained offspring from Nhe8–/– females, male Nhe8–/– mice never produced progeny, as reported previously (26, 30). Vaginal plugs observed in breedings with male Nhe8–/– mice indicated normal mating behavior, libido, copulation, and ejaculation. Body (WT, 27.5 ± 1.5 g; Nhe8–/–, 26.4 ± 1.0 g; n = 12) and testes weight (WT, 0.57 ± 0.03% of body weight; Nhe8–/–, 0.59 ± 0.04% of body weight; n = 12) was unchanged in Nhe8–/– mice. Their male reproductive systems appeared macroscopically normal (Fig. 1C), as did their seminiferous tubules (Fig. 1E), the diameters of which were unchanged (Fig. 1J).

Nhe8–/– mice have round-headed spermatozoa and lack acrosomes
Seminiferous tubules of both WT and Nhe8–/– mice contained spermatozoa (Fig. 1F) and the epididymal sperm concentration was not significantly different (Fig. 1J). Contrasting with the typical elongated heads of WT spermatozoa (Fig. 1G, top), Nhe8–/– spermatozoa had round heads, within seminiferous tubules (Fig. 1G, bottom) or the cauda epididymis (Fig. 1D). Their motility was drastically reduced (Fig. 1K). Nhe8–/– spermatozoa lacked acrosomes and showed an abnormal distribution of mitochondria which were found in sperm heads instead of forming a sheath around the initial part of the flagellum. This was revealed by crosses with “green sperm” mice (34) that express dsRed in mitochondria and GFP in acrosomes (Fig. 2A) and by labeling acrosomes with peanut agglutinin (PNA) (35) and mitochondria with MitoTracker® (Fig. 2B) and ascertained by transmission electron microscopic (TEM) images (Fig. 3, A–D). Because the mitochondrial sheath provides ATP for flagellar movement (36), this finding might explain the reduced motility of Nhe8–/– spermatozoa.

The lack of Golgi-derived acrosomes upon ablation of the Golgi-resident NHE8 protein (17) points to a germ cell-intrinsic effect, but also defects in germ-cell nurturing Sertoli cells or hormone-producing Leydig cells may cause infertility (37). X-Gal staining of testes from mice carrying the lacZ reporter gene in the Nhe8 locus (Fig. 1A) revealed Nhe8 transcription in seminiferous tubules, but not in intertubular Leydig cells (Fig. 2C). Immunofluorescence detected NHE8 protein in WT (but not Nhe8–/–) germ cells, where it localized to an acroosome-like structure in close proximity to the nucleus (Fig. 2D). NHE8 perfectly co-localized with the acrosomal marker PNA in all four phases of spermiogenesis (Fig. 2E). Absence of immunoreactivity in Nhe8–/– testes (Fig. 2D) confirmed the specificity of labeling of preacrosomal granules. However, it cannot strictly guarantee that NHE8 is expressed on mature acrosomes because these structures are not present in Nhe8–/– testes. Comparison of spermiogenesis between the genotypes showed that only the Golgi phase appeared normal in Nhe8–/– mice (Fig. 2F). Instead of a single acrosomal granule that grows and flattens in WT mice, many small PNA-positive vesicles were detected on the nuclear surface in Nhe8–/– spermatids (Fig. 2F, cap phase). In the maturation phase, the resulting acrosomeless Nhe8–/– spermatids failed to obtain the typical elongated shape, although nuclei were condensed (Fig. 2F, maturation phase). TEM of WT testis showed the generation of acrosomal granules by fusion of GDVs, their attachment to the nucleus (Fig. 3E), growth, and flattening (Fig. 3F), as well as nuclear elongation (Fig. 3G). By contrast, in Nhe8–/– mice GDVs accumulated in the region between the Golgi and the nucleus (Fig. 3, H and I) but failed to generate larger granules or acrosomal caps. GDVs were more electron-dense in Nhe8–/– cells. Although spermatozoa in the seminiferous tubules of Nhe8–/– mice had condensed chromatin, their nuclei frequently displayed irregular shapes (Fig. 3F). The preacrosomal granule is attached to the nucleus by a cytoskeletal scaffold, the acroplaxome (38). The acroplaxome was visible in TEM as a dense plate in the subacrosomal space in WT spermatids (Fig. 3K) as well as in the space between the nucleus and the GDVs in Nhe8–/– spermatids (Fig. 3, L and M). This indicates that the defects in acroosome biogenesis are probably caused by impaired fusion of GDVs, whereas tethering of these vesicles to the nuclear envelope seems to be unaffected.

Germ cell-specific disruption of Nhe8 causes a phenotype indistinguishable from Nhe8–/– mice
Our data suggest that NHE8 is directly involved in acroosome biogenesis and that disruption of this process causes the infertility of Nhe8–/– mice. To test this hypothesis, Nhe8lox/lox mice with AMH-Cre (39) and Stra8-iCre (40) mice, respectively. Specificity of Cre expression was ascertained by crosses with R26R reporter mice (41) (Figs. 4A and 5A). In the following, we refer to the resulting Sertoli cell- and germ cell-specific knock-out mice as SC-ΔNHE8 and GC-ΔNHE8 mice, respectively. NHE8 protein levels of SC-ΔNHE8 testes were indistinguishable from WT mice, indicating that NHE8 is not, or is only at very low levels, expressed in Sertoli cells (Fig. 4B). As expected, male SC-ΔNHE8 mice were fertile and always produced offspring when mated over a 2-month period (n = 7 SC-ΔNHE8 males). Neither morphology and diameter of seminiferous tubules (Fig. 4, C and E), nor sperm morphology was affected (Fig. 4, E and F). Spermatozoa from SC-ΔNHE8 mice displayed normal acrosomal caps and mitochondrial sheaths (Fig. 4G). Hence NHE8 is dispensable in Sertoli cells for normal
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sperm development and fertility. In contrast, testicular NHE8 protein levels were drastically reduced in GC-DNHE8 mice (Fig. 4B), indicating that in mouse testes NHE8 is predominantly expressed in germ cells. The testicular phenotype of GC-DNHE8 mice appeared identical to that of Nhe8−/− mice, except for the finding that ~5% of spermatozoa from GC-DNHE8 mice appeared normal, which can be attributed to the known ~95% deletion efficiency of Stra8-iCre mice (40). The identical phenotype included male infertility (n = 15 GC-DNHE8 males), normal morphology and diameter of the seminiferous tubules (Fig. 4E), sperm concentration in the caudae epididymis (Fig. 4J), and drastically reduced motility of spermatozoa from Nhe8−/− mice (Fig. 4K, ****, p < 0.0001, unpaired Student’s t test).
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Figure 2. NHE8 expression in testes and germ cells is shown. A, mature green spermatozoa (expressing GFP tagged acrosin in green) and dsRed tagged mitochondria (in red) isolated from the caudae epididymides of WT and Nhe8<sup>−/−</sup> green sperm mice, fixed and counterstained with DAPI (blue) to reveal nuclei. B, immunofluorescent labeling of isolated WT and Nhe8<sup>−/−</sup> spermatozoa. Labeling of the acrosomal cap (PNA, red), the nucleus (DAPI, blue), and the mitochondrial sheath (MitoTracker<sup>®</sup>, green). C, X-Gal staining on sections of Nhe8<sup>−/−</sup> or WT testes showing Nhe8 expression in blue. D, immunofluorescent labeling of NHE8 in seminiferous tubules (surrounded by white lines) of testis from WT and Nhe8<sup>−/−</sup> mice. Magnified regions show the caplike NHE8-positive structures in the WT. Antibody, rbnuNHE8–10. E, co-localization of NHE8 (green) with the acrosomal marker PNA (red) to the developing acrosome in all four phases of spermiogenesis in WT mice. Antibody, rbnuNHE8–10. Nuclei labeled with DAPI. F, PNA labeling (red) contrasts the normal development of the nucleus-apposed acrosome in WT mice to the impaired biogenesis in Nhe8<sup>−/−</sup> mice. Results shown in this figure were observed in at least two WT/KO mouse pairs.

Discussion

Spermatogenesis is the long process that leads from diploid spermatogonia to spermatozoa, highly differentiated haploid motile gametes. The complexity of this differentiation process implies that mutations in many different genes can interfere with spermatogenesis and thereby result in infertility (3). We now showed that disruption of Nhe8 causes a specific infertility syndrome, globozoospermia, which is characterized by round-headed, acrosome-less sperm (42, 43). The acrosome is a unique secretory sperm organelle that is essential for male fertility (2, 3). Golgi-resident NHE8 was needed in a cell-intrinsic manner for the biogenesis of this Golgi-derived organelle. Our work adds NHE8 to the growing list of intracellular ion transport proteins with crucial roles in vesicle trafficking and fusion (44).

Acrosomes are unique large vesicles sandwiched between the plasma membrane and the nucleus at the tip of sperm cells. The acrosome reaction, a highly unusual form of exocytosis, releases lytic enzymes and exposes acrosomal membrane proteins at the outer sperm membrane (3, 45). The release of lytic enzymes is thought to facilitate the dispersion of cumulus cells and the penetration of the zona pellucida, but KO mice showed that many of these enzymes are individually dispensable for fertility (1, 3). By contrast, the immunoglobulin family member Izumo, an acrosome-resident membrane protein that is exposed at the sperm surface after the acrosome reaction, is absolutely required for sperm-oocyte fusion and hence fertility (3, 46). Acrosomes may also play a role in shaping sperm nuclei by their acroplaxome-mediated tight interaction (38). Indeed, the loss of the acrosome is generally associated with abnormally shaped nuclei in globozoospermia (45, 47–49). The loss of a mitochondrial sheath, as observed here, is also found in other genetic forms of globozoospermia (47–49) but the underlying mechanism is unclear.

Acrosomes are mainly generated by fusion of Golgi-derived vesicles but they share some properties, including their strongly acidic luminal pH and many degradative enzymes, with lysosomes (50, 51). The molecular mechanisms underlying the budding, trafficking, and fusion of GDVs are still largely unknown. We detected NHE8 not only on the Golgi, where it has been found in previous studies (17), but also during all phases of spermiogenesis on acrosomal granules and finally on mature
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Figure 3. Defective formation of the acrosome and mitochondrial sheath in Nhe8−/− mice is shown. A and B, TEM images of epididymal spermatozoa from WT (A) and Nhe8−/− (B) mice. x: axoneme; m: mitochondria. Red boxes indicate areas of A and B that are shown in higher magnification in C and D, respectively. E–G, TEM images from WT mice showing different phases of spermiogenesis. E, Golgi phase. F, cap phase. G, maturation phase. H–J, equivalent images from Nhe8−/− mice. H, electron-dense vesicles attached to the nuclear envelop can be found. Acrosomal granules or acrosomal caps were not detected. I, single dense vesicles were found in nuclear notches. J, acrosome-less spermatozoa with irregularly shaped nuclei. g: Golgi apparatus; a: acrosome or acrosomal granule; m: mitochondrial sheath; x: cilia crossections. Nuclei highlighted in red. K–M, magnification of the areas delineated by red lines in E, H, and I, respectively, reveal the acroplaxome (red arrows).

acrosomes. Together with the observation that sperm-specific Nhe8 disruption sufficed to impair acrosome biogenesis, this localization strongly suggests that NHE8 is directly required for acrosome formation. As we observed GDVs in Nhe8−/− spermatids by both PNA labeling and electron microscopy, it is

Figure 4. NHE8 is dispensable in Sertoli cells for normal spermiogenesis. A, X-gal staining on frozen tissue sections from Rosa26R+/−, AMH-Cre mice, and control mice (Rosa26R+/−; AMH-Cre−/−). B, Western blots of testes lysates indicate complete absence of the NHE8 protein in Nhe8−/− mice (lane 2) and drastic reduction in GC−/−NHE8 mice (lane 4) in which Nhe8 is specifically deleted in germ cells using Stra8-iCre mice. Protein levels in Nhe8lox/lox or SC−/−NHE8 mice are equal to Nhe8−/− mice. Protein load, 30 μg per lane. Antibody, rbNHE8-CT. Actin served as loading control. C, tubule diameter was unchanged between genotypes (WT mean, 159 ± 7 μm, n = 51 tubules from two mice; SC−/−NHE8 mean, 165 ± 0.5 μm, n = 90 tubules from two mice; Nhe8lox/lox mean, 167 ± 3 μm, n = 334 tubules from two mice; GC−/−NHE8 mean, 172 ± 2 μm, n = 344 tubules from three mice). D and E, H&E staining on testes sections from (D) WT and (E) SC−/−NHE8. Right panels show a magnification of the areas delineated by broken lines. F, H&E staining of isolated sperm from SC−/−NHE8 mice. G, fluorescent labeling of the acrosome (PNA, red), nucleus (DAPI, blue), and mitochondrial sheath (MitoTracker®, green) on isolated spermatozoa of WT and SC−/−NHE8 mice.
highly likely that NHE8 is needed for its fusion to acrosomes rather than for the detachment from the Golgi. Indeed, NHE8 and the Saccharomyces cerevisiae orthologue Nhx1p, which is closely related to organellar NHEs, have been proposed as regulators of vesicle fusion (23, 52). NHE8 knockdown in HeLa cells led to an increase of intraluminal vesicles in multivesicular bodies, suggesting NHE8 as a regulator of vesiculation or a promoter of the fusion of late multivesicular bodies with lysosomes (23). Finally Hrb−/−, Pick1−/−, and GOPC−/− mice, which display globozoospermia very similar to Nhe8−/− mice, are thought to result from defective GDV fusion (47–49).

How might NHE8 influence the fusion of GDVs? Being a Na+/H+ -exchanger, its loss may change both vesicular pH and Na+ concentration. These changes can secondarily affect other luminal ion concentrations like that of Ca2+, the release of which may be needed for vesicle fusion and trafficking (53–55). Although not quite reaching the pH of lysosomes, the luminal pH of acrosomes (pH ∼5.3) (56, 57) is more acidic than that of the Golgi (estimated pH ∼5.9–6.3, depending on cell type) (58–60). A proton “leak” by NHE8 might tune the acidification of GDVs. Luminal pH can affect vesicular trafficking e.g. in the endosomal/lysosomal pathway (61), where it may affect endosomal carrier formation (62) and the cytosolic binding of proteins involved in trafficking (63). Alternatively or additionally, NHE8 might directly interact with proteins of the fusion machinery, as shown for GOPC and Pick1 (47). Both proteins possess a PDZ domain whereas NHE8 features a PDZ-binding domain. However, preliminary co-immunoprecipitation experiments (data not shown) failed to ascertain the attractive hypothesis that NHE8 binds to one of these proteins.

Although we unambiguously identified a germ cell-intrinsic defect in acrosome formation as underlying the male infertility of Nhe8−/− mice, Xu et al. (30) recently suggested that it is caused by reduced testosterone levels stemming from impaired trafficking of the luteinizing hormone receptor in Leydig cells. The authors state that NHE8 is mainly expressed in Leydig cells (30), without, however, using Nhe8−/− controls as done here. Disconcertingly, they also observed smaller testes and azoosperma. We cannot explain this discrepancy, but differences in genetic background or the disruption of an additional gene in their gene trap approach (30) might possibly be invoked.

The ablation of several genes causes globozoospermia in mice, but so far only three genes are known to underlie globozoospermia in humans, DPY19L2, SPATA16, and PICK1 (64–66). Genetic causes remain unknown in one-third of globozoospermia patients. This study adds SLC9A8 to the list of candidate genes for this disorder. Moreover, it expands the list of ion transporters involved in vesicle trafficking or fusion and highlights the biological importance of vesicular ion homeostasis.

Experimental procedures

Mice

Animal care and experiments conformed to German animal protection laws and were approved by the Berlin authorities (Landesamt für Gesundheit und Soziales Berlin). Sperm of mice heterozygous for the targeted Nhe8<sup>fl</sup> allele (Slc9a8<sup>fl</sup>(<sup>KOMP</sup>)) was ordered via the Knockout Mouse Project (KOMP) (Davis, CA). In vitro fertilization of C57BL/6 female mice was carried out by the transgenic core facility of the Max-Delbrück-Centrum für Molekulare Medizin (MDC) Berlin. Targeted mice were crossed to Cre or FLP recombinase expressing deleter mice to generate different Nhe8 mouse lines (see Fig. 1A) (67, 68). Nhe8<sup>fl</sup> mice were crossed to AMH-Cre mice (B6-Tg(Amhc-cre))<sup>8815</sup>RebJ, obtained from The Jackson Laboratory, no. 017490) for cell-specific deletion of Nhe8 in Sertoli cells or male premeiotic germ cells (39, 40). AMH-Cre
mice and Stra8-iCre mice were crossed to Rosa26R Cre reporter mice (B6.129-Rosa26tm, obtained from Fred Hutchinson Research Center). Green sperm mice (34) were obtained from RIKEN (B6D2-Tg(CAG/Su9-DsRed2,Acr3-EGFP) RBGS002Osbg, no. RBRC03743) and crossed to NHE8 mice. These spermatozoa express NHE8 (indicated by yellow arrows).

**Generation of NHE8 antibodies**

Antibodies to two epitopes of NHE8 (494NAIESEHLSE-LTEEEYEAH512 for rbNHE8–10 and 559EEVRQGPSGSED-DEQELF576 for rbHE8–CT) were raised in rabbits (Pineda Antibody). X-Gal staining was done on frozen sections or on isolated epididymal sperm (see “Immunohistochemistry”). A-1F1 (1:1000), in parallel Probes). Confocal images were taken with a Zeiss LSM 510 META laser scanning microscope and ZEN software (Zeiss). Image processing was done using ZEN software (Zeiss). All stainings were repeated in at least three independent experiments. Spermatids were assigned to different phases of spermiogenesis by their location inside the seminiferous tubule (cells move toward the luminal region during the maturation process) and nuclear morphology. In WT mice, the size and form of the acrosomal cap served as another indicator to assign the correct phase of spermiogenesis.

**Histology**

For histological analyses, mice were perfused with 4% PFA/PBS (w/v) and organs were postfixed overnight at 4 °C. After dehydration in a graded ethanol series, organs were stored in isopropanol/paraffin overnight at 65 °C. After incubation in 100% paraffin for another 6–10 h, organs were embedded in paraffin and 6–μm sections were cut (Cool-cut Microm HM 355 S, Thermo Scientific). Hematoxylin and eosin staining was performed on these sections or on isolated epididymal sperm dried on coverslips. X-Gal staining was done on frozen sections from Nhe8NeoNeo or Nhe8+/− mice (see “Immunohistochemistry”) for 12–48 h at 37 °C in a humidified chamber (staining solution: 2 mM MgCl2, 5 mM KFe(CN)6, 5 mM K4Fe(CN)6, 0.2% Nonidet P-40, 0.1% sodium deoxycholate, 1 mg/ml X-Gal in 0.1 M phosphate buffer, pH 7.4). Images of X-Gal stainings and hematoxylin and eosin stainings were taken with an AxioCam MRa5 (Zeiss) on an Axiophot microscope (Zeiss) with an Achrostigmat 5×/0.12, Plan-Neofluar 10×/0.30, or Plan-Neofluar 40×/0.75 objective using the ZEN software (Zeiss). Image processing was done with Adobe Photoshop.

**Isolation and analysis of mouse spermatozoa**

Mice were killed by cervical dislocation and the caudae epididymides were removed immediately. They were thoroughly cleaned of fat and blood vessels, each transferred to 200 μl of cryoprotective agent (CPA) (180 g/liter raffinose, 30 g/liter skim milk), and ruptured. Sperm was allowed to swim out for 5 min at room temperature. For stainings, sperm in cryoprotective agent were diluted in PBS and smeared on coverslips. Staining was performed as for immunohistochemistry. For investigation of
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the mitochondrial sheath, spermatozoa were incubated with MitoTracker® green (Molecular Probes) at a final concentration of 100 nM for 10 min at 37 °C, centrifuged 1 min at 5000 × g, resuspended in PBS, and smeared on coverslips. Green sperm were stained with chpended in PBS, and smeared on coverslips. Green sperm were

Sperm motility measurements
Isolated sperm were incubated in human tubal fluid (HTF) medium at 37 °C for 5 min in a 1:50 dilution. Sperm motility and concentration were assessed using IVOS sperm analyzer version 12 (Hamilton Thorne Research, Beverly, MA). Spermatozoa were tracked for 30 s and images were captured at a frequency of 60 Hz. The cutoff of motile spermatozoa was 20 μm/s for the path velocity (distance traveled along a smoothed path divided by elapsed time) and 30 μm/s for the linear velocity (straight-line distance between starting and end point divided by elapsed time).

Western blot analyses
Membrane fractions were isolated from mouse tissue. To this end, tissue homogenate was prepared in 20 mM Tris-HCl pH 7.4, 140 mM NaCl, 5 mM EDTA with protease inhibitors (4 mM Pefabloc®, 1Complete EDTA-free protease inhibitor mixture (Roche)) using a glass dounce homogenizer and cleared by centrifugation for 10 min at 1000 × g twice. Membrane fractions were pelleted from the cleared homogenate by ultracentrifugation for 30 min at 270,000 × g, and the pellet was resuspended by sonication in 50 mM Tris pH 6.8, 5 mM EDTA, 2% SDS (w/v) with protease inhibitors. Equal amounts of protein were separated by SDS-polyacrylamide gel electrophoresis and blotted onto nitrocellulose. Blots were reprobed with mouse anti-β-actin (clone AC-74; Sigma, no. A2228) (1:5000) as a loading control.

Transmission electron microscopy
Mice were anesthetized and perfused transcardially (4% formaldehyde, 2.5% glutaraldehyde in 0.1 M phosphate buffer). Testes were isolated and postfixed in the same solution overnight at 4 °C, embedded in 5% agar, and sliced (200 μm thick) with a vibratome. Slices were postfixed in 1% OsO4 and 1.5% potassium hexacyanoferrate, dehydrated in a methanol gradient and propylene oxide, and flat embedded in epoxy resin. After polymerization, seminif-

Statistics
Statistical significance was assessed using the paired or unpaired Student’s t test (Prism, GraphPad Software). Errors are indicated as ± S.E.

Author contributions—K. O. designed, performed, and evaluated experiments and wrote the paper. D. P. designed, performed, and evaluated TEM experiments and wrote the paper. T. J. J. initiated the project, designed and evaluated experiments, and wrote the paper.

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