

The sodium/proton exchanger SLC9C1 (sNHE) is essential for human sperm motility and fertility

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1 The sodium/proton exchanger SLC9C1 (sNHE) is essential for human sperm motility and

2 fertility.

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- 21
- 22 **<u>Running Title</u>** : *SLC9C1* mutation in human asthenozoospermia
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28 Abstract

29 Asthenozoospermia, defined by the absence or reduction of sperm motility, constitutes the most frequent cause of human male infertility. This pathological condition is caused by 30 31 morphological and/or functional defects of the sperm flagellum, which preclude proper sperm progression. While in the last decade many causal genes were identified for asthenozoospermia 32 associated with severe sperm flagellar defects, the causes of purely functional 33 asthenozoospermia are still poorly defined. We describe here the case of an infertile man, 34 displaying asthenozoospermia without major morphological flagellar anomalies and carrying a 35 homozygous splicing mutation in SLC9C1 (sNHE), which we identified by whole-exome 36 37 sequencing. SLC9C1 encodes a sperm-specific sodium/proton exchanger, which in mouse regulates pH homeostasis and interacts with the soluble Adenylyl Cyclase (sAC), a key 38 regulator of the signalling pathways involved in sperm motility and capacitation. We 39 demonstrate by means of RT-PCR, immunodetection and immunofluorescence assays on 40 patient's semen samples that the homozygous splicing mutation (c.2748+2T>C) leads to in-41 42 frame exon skipping resulting in a deletion in the cyclic nucleotide-binding domain of the protein. Our work shows that in human, similar to mouse, SLC9C1 is required for sperm 43 motility. Overall, we establish a homozygous truncating mutation in SLC9C1 as a novel cause 44 of human asthenozoospermia and infertility. 45

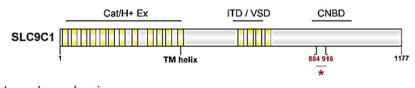
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47 <u>Key words</u>: male infertility; asthenozoospermia; gene mutation; whole-exome sequencing;
48 SLC9C1 - sNHE; ion channel.

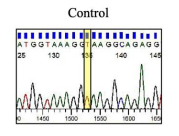
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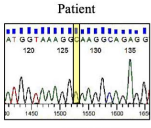
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SLC9C1 (sNHE): Sperm specific Sodium/Proton Exchanger

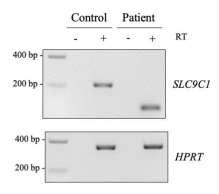


Cat/H+ Ex: Ion exchange domain ITD/VSD: Voltage sensing domain CNBD: Cyclic Nucleotide Binding Domain

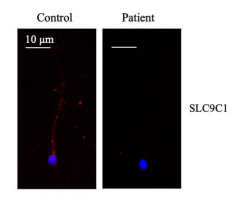




c.2748+2T>C



In-frame exon skipping



In-frame protein truncation **p. del 884-916**

54 Introduction

55 Asthenozoospermia, defined by the absence or reduction of sperm motility, is observed in nearly 80% of male infertility cases, either alone or in association with other sperm defects¹. 56 57 This pathophysiological condition is induced by structural defects of the sperm flagellum and/or functional alterations impairing flagellar beating and sperm progression. Among the structural 58 defects, a remarkable condition described as Multiple Morphological Abnormalities of sperm 59 Flagella (MMAF) results from defective assembly of the flagellum and leads to the presence in 60 the ejaculate of spermatozoa with short, absent, coiled, bent or irregular tails. This phenotype 61 of asthenoteratozoospermia was previously identified as 'dysplasia of the fibrous sheath', 'short 62 tails' or 'stump tails'²⁻⁴, and in the last decade, numerous mutations causing this phenotype 63 were identified (AK7, ARMC2, MAATS1, CEP135, CFAP43-44-65-69-70-251, DNAH1-2-6-64 8-17, ORICH2, SPEF2, TTC21A, TTC29)^{5,6}. Besides MMAF, some asthenoteratozoospermic 65 patients with milder morphological defects such as defective annulus and abnormal 66 mitochondrial sheath were also described to carry mutations (TEKT2, SEPT12 or 67 *SLC26A8*)^{1,7,8}. Importantly, asthenozoospermia is also detected in the complete absence of 68 morphological defects, suggesting the involvement of functional defects in activation and/or 69 regulation of flagellar beating. To date, such cases of functional asthenozoospermia remain 70 71 poorly characterized, and only a few mutations were identified in genes encoding for proteins with enzymatic properties (GALNTL5), seminal component (SEMG1) and ion channels 72 (CATSPER1-2- ε , SLC26A3, VDAC2)^{1,9-12}. Overall, the knowledge obtained by studying those 73 sperm pathological phenotypes provides evidence that, in addition to proper morphology and 74 structure, sperm fertilization potential relies on proper activation of the signalling pathways 75 76 regulating motility, metabolism and energy production.

Sperm functional maturation occurs after spermiation during the transit within theepididymis, where sperm cells acquire the ability to move forward, and ultimately within the

female genital tract, where they acquire complete fertilization potential through a process 79 known as capacitation¹³. Numerous ion channels and transporters have been identified at the 80 surface of sperm cells and conduct complex fluxes between the male and female genital tract 81 milieus and the sperm cytoplasm¹⁴. These ion exchanges activate several biochemical and 82 electrophysiological changes that are essential for sperm motility and fertilization potential¹⁵. 83 Among the changes occurring within the female genital tract (i.e. capacitation), were described 84 alkalinization of the cytoplasm, membrane hyperpolarization and protein phosphorylation 85 cascades induced by the activation of soluble adenylate cyclase (sAC) and Protein Kinase A 86 (PKA), which specifically target flagellar components required for sperm fertilization 87 potential¹⁶. 88

In the present study, by performing Whole-Exome Sequencing (WES) on a patient presenting a very low progressive motility but no detectable ultrastructural defects of the axoneme, we identified a homozygous pathogenic mutation in *SLC9C1*. This gene, also called *sNHE*, encodes a sperm-specific member of the Na⁺/H⁺ exchanger family involved in the regulation of pH and cell volume in a wide range of tissues¹⁷. We demonstrate here that *SLC9C1* is critical for human sperm motility and that mutations in *SLC9C1* are a novel cause of human asthenozoospermia.

96 Materials & methods

97 Subject and biological samples

The study was approved by the Comité de Protection des Personnes CPP IIe de France III (record number CPP02748) and performed in accordance with ethical guidelines (Declaration of Helsinki). All patient and control individuals included in the study gave their informed consent prior to their inclusion in the study. Semen samples were obtained by masturbation after 2-7 days of sexual abstinence. Semen parameters were evaluated according to the World Health Organization (WHO) guidelines¹⁸ and David's classification¹⁹.

104

105 WES analysis and Sanger sequencing

106 WES analysis was performed on genomic DNA extracted from a blood sample, as previously described²⁰. In brief, the enrichment of coding regions together with intron/exon boundaries 107 was performed with Exon V5 kit (Agilent Technologies, Wokingham, UK). Sequencing was 108 performed with Illumina HiSeq 2000 at the Genoscope (Evry, France). Exome-seq data were 109 analysed using a bioinformatics pipeline developed in-house as previously described²¹. The 110 homozygous mutation in SLC9C1 identified by WES was confirmed by Sanger sequencing 111 using ABI 3130XL and SeqScape software (Applied Biosystems; Foster City, CA, USA). 112 F: Primer sequences (5'-3'): TCCCAACCACTTCTAAAATGTTGT, R: 113 114 TGCATTTATAAATAACACTGCCTGGT.

115

116 Transmission electron microscopy analysis

Sperm cells were collected from fresh ejaculate and washed with M2 medium (Sigma-Aldrich
Co. Ltd; Irvine, UK) by centrifugation at 300g/RT for 10 minutes. Sperm cells were then fixed

and embedded for semi-thin sections as previously described²². All sections were examined
with a JEOL 1011 electron microscope (Jeol Ltd; Tokyo, Japan). Images were acquired with
Digital Micrograph software coupled to a Gatan Erlangshen CCD camera.

122

123 **RT-PCR analysis**

Total RNA from control and patient spermatozoa (800-1000ng) was extracted using
NucleoSpin RNA kit (Macherey-Nagel; Düren, Germany) and used for reverse transcription
with High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Thermo Fisher
Scientific; Waltham, MA, USA) as previously described²². Amplicons were gel purified and
sequenced (Eurofins Scientifics, France). Results were analysed with BioEdit software (Ibis
Therapeutics; Carlsbad, CA, USA).

130 Primer sequences (5'-3'): *SLC9C1*-F: TTATCAGGCCTCTTACTGTTG, *SLC9C1*131 R: TGATCAATCCCTAAACCTGGC; *HPRT*-F: CCTGCTGGATTACATTAAAGCACTG,
132 *HPRT*-R: GTCAAGGGCATATCCAACAACAAAC.

133

134 Immunofluorescence assay

Slides were prepared by spreading 10 µL of fresh semen sample onto a Superfrost Plus slide (Menzel Glasbearbeitungswerk, GmbH & Co. KG; Braunschweig, Germany), followed by fixation 10 minutes in PBS 4% paraformaldehyde. Immunostaining was performed as previously described²², using commercial antibodies, which specificity was not validated.

Primary antibodies : SLC9C1 rabbit polyclonal antibody raised against the central region
(Invitrogen PA5-104160; 1:100 dilution); SLC9C1 rabbit polyclonal antibody raised against
the C-terminal region (Sigma SAB2106664; 1:100 dilution); mouse monoclonal anti-α-Tubulin

Sigma T9026 (1:500 dilution). Secondary antibodies: goat polyclonal anti-rabbit IgG Alexa
Fluor 568 (Invitrogen A-11029; 1:500 dilution) and goat polyclonal anti-mouse IgG Alexa
Fluor 488 (Invitrogen A-28175; 1:500 dilution).

145

146 Western blot analysis

147 10 million sperm cells from control individual and patient were washed in M2 medium and next 148 in PBS. Sperm cells were then subjected to sonication (20s on, 60s off, 15 cycles) in RIPA buffer [150 mM NaCl, 1% NP40 (Sigma-Aldrich Co.; St. Louis, MO, USA), 0.5% DOC, 0.1% 149 150 SDS, 25 mM TrisHCL, cOmplete Protease Inhibitor Cocktail (Roche, Basel, Switzerland)]. Samples were kept on ice for 30 minutes and centrifuged 13000g at 4°C for 15 minutes. The 151 supernatant was collected and denatured in Laemmli sample buffer, prior to SDS-PAGE [8% 152 acrylamide/bisacrylamide (40%, 37.5:1)] and transfer to nitrocellulose membranes. After a 153 blocking step in PBS 0.1% Tween, 5% BSA or milk for SLC9C1 and β-Tubulin detection, 154 155 respectively, incubation was performed with primary and secondary antibodies. Primary antibodies: rabbit polyclonal anti-SLC9C1 Invitrogen (PA5-104160; 1:500 dilution) and mouse 156 monoclonal anti-β-Tubulin clone AA2 (Sigma 05-661; 1:2500 dilution). Secondary antibodies: 157 158 rabbit polyclonal anti-mouse Ig coupled to Horse Radish Peroxydase (HRP) (Dako; 1:1000 dilution), swine polyclonal anti-rabbit Ig coupled to HRP (Dako; 1:1000 dilution). 159

161 **<u>Results</u>**

162 The patient presented in this study first consulted for primary infertility in 2016 at the age of 31, after two years of natural conception failure, and was followed at the reproductive 163 164 biology centre of the Cochin hospital (Paris, France). Apart from a non-evolutive chronic hepatitis B diagnosed in 2008, no urogenital disease, neither lifestyle factors nor exposures 165 potentially deleterious for fertility were reported. The couple benefited from intra-marital ICSI 166 performed with fresh ejaculated sperm cells but frozen oocytes because the man failed to 167 produce a semen sample on the day of oocytes pick-up of the first ICSI attempt. Seven out of 168 nine of the thawed oocytes were injected and 3 zygotes were observed 18 hours after ICSI. 169 170 After a prolonged embryo culture, 3 embryos with a good quality, according to the Istanbul consensus 2011 classification²³, were obtained. Embryo transfer scheduled on day 5 was 171 postponed due to a high endometrial thickness; two of the embryos were frozen at day 5 and 172 the third one at day 6. To date, only one thawed embryo was transferred but the resulting 173 pregnancy ended with a miscarriage after 5 weeks and 3 days. 174

The successive semen analyses performed over 3 years on the patient revealed a severe 175 alteration of sperm progressive motility, which never exceeded 5% (normal value > 32%), 176 177 contrasting with normal sperm counts and vitality rates (Table 1). The mean percentage of typical spermatozoa was slightly decreased but the patient did not present any sperm 178 abnormalities indicative of a MMAF phenotype (the percentage of spermatozoa with short, 179 absent and irregular calibre of flagella being under the lower reference values¹⁹) (Table 1). 180 However, semen analyses indicated a significant proportion of spermatozoa with an excess 181 residual cytoplasm and disorganised midpiece with occurrences of plasma membrane lysis 182 (Figure 1A). In addition, flagellar bending was noticed together with some head anomalies (thin 183 head) (Table 1). Acrosomal staining performed with fluorescent peanut agglutinin (PNA) on 184 sperm sample from the patient indicated an intact acrosome on most of the spermatozoa; hence 185

the rate of PNA negative sperm cells, indicative of premature acrosomal reaction, was 8.8% 186 and 5.4% for control and patient, respectively. The procedure of sperm selection with density 187 gradient centrifugation slightly increased the fraction of progressive spermatozoa (5-25%) but 188 also induced an increase of the number of isolated sperm heads. We found that the percentage 189 of isolated sperm heads remained very low in control samples defined by normal sperm 190 parameters, including morphology according to WHO¹⁹ (1% and 3% before and after selection, 191 respectively), while it significantly increased in patient sample after density gradient 192 centrifugation (2% and 33% before and after selection, respectively). Importantly, we noted 193 that flagellar bending and breaking points on sperm from the patient were not limited to the 194 195 connective piece but were also present along the flagella (Supplementary Figure S1). Overall, 196 these observations suggested an important fragility of the sperm tail, amplified by the selection procedure. 197

We performed Transmission Electron Microscopy (TEM) analysis to precisely examine 198 the sperm ultrastructure of the patient. In line with the observations by optic microscopy, we 199 200 detected the presence of midpiece defects, in particular, an excess of residual cytoplasm around 201 the mitochondrial sheath and plasma membrane fragility illustrated by plasma membrane lysis (Figure 1B). Importantly, no alteration of the axonemal structure was observed in patient's 202 203 sperm as quantification indicated a percentage of normal transversal sections (9+2 conformation) close to that recorded in semen from control individuals (Patient: 84%; Control 204 205 mean: 84.5%, n=4). Overall, the patient presented in this study was classified as a case of asthenozoospermia with a severe reduction of the proportion of progressive sperm, membrane 206 and flagellar fragility but no major defects of the sperm flagellum ultra-structure and assembly. 207

We thus subsequently performed WES analysis of this patient in search for genetic variants potentially involved in the infertility phenotype. After processing and filtering the data, we identified homozygous variants with high predicted impact on the encoded proteins in three

genes: c.2748+2T>C in SLC9C1, c.3583C>T in ABCB5 and c.9del in ZNF891. Among those, 211 212 the variant c.2748+2T>C in SLC9C1 appeared as the best candidate considering the pattern and abundance of expression of the three genes and the frequency of the variants in the 213 population according to public databases (Supplementary Table 1). Indeed, in contrast to the 214 215 two other candidates, SLC9C1 was shown to be preferentially expressed in the testis (Illumina Body Map; The Human Protein Atlas database; Reprogenomics Viewer) and the encoded 216 protein was previously detected in human sperm proteome²⁴ (Supplementary Table 1). The 217 homozygous variant c.2748+2T>C identified in SLC9C1 impairs a consensus sequence of the 218 splice donor site located downstream of exon 22 and was associated with a low allele frequency 219 in the gnomAD database (4.6x10⁻⁵, chr3:112180562 A>G (GRCh38.p12), rs2007949663-220 221 111899409-A-G). The presence of this homozygous variant in the patient was also consistent with the familial history of infertility and the known consanguinity of his parents (cousins). The 222 proband originated from West Africa (Guinea Conakry); he has two brothers, one of them 223 known to be infertile, and two fertile sisters (Figure 2A). In addition, the proband has one cousin 224 who also displays male infertility; as it was not possible to define if the cousin was from 225 maternal or paternal side, this individual is not included in the genealogic tree (Figure 2A). 226 227 Unfortunately, we could not obtain to biological samples from the family members. In 228 conclusion, considering all the above arguments, we retained the c.2748+2T>C variant identified in SLC9C1 by WES, which we also confirmed by Sanger sequencing on DNA sample 229 from the proband (Figure 2B). 230

SLC9C1, also known as *sNHE* or *SLC9A10*, is located on chromosome 3 and comprises
29 exons (Figure 2C). Alternative splicing produces five transcript variants, three of which
encoding proteins, and among them transcript *SLC9C1-201* encodes for the longer isoform of
135 kDa (Figure 2C). The full SLC9C1 protein isoform comprises the ion exchange domain
(Cat/H⁺ ex), which is characteristic of SLC9 family members and is generally formed by ten to

twelve transmembrane (TM) helices (Figure 2C). The protein also exhibits a unique C-terminal tail with four additional TM helices showing similarity to a Voltage-Sensing Domain (ITD/VSD) and a consensus sequence for a Cyclic Nucleotide-Binding Domain (CNBD), which is composed of several α -helices and β -sheets and protrudes towards the cytoplasmic side^{25,26}. These two latter features are typical of ions channels, which, unlike solute carriers, are not activated by ion gradients but display a more sophisticated regulatory mechanism dependent on the cell status¹⁴.

To investigate the molecular consequence of the mutation identified in *SLC9C1*, we 243 performed RT-PCR analysis on semen sample from control and patient individuals using 244 245 forward and reverse primers localised in exons 21 and 23, respectively. The PCR amplicon obtained for control individual was observed at the expected size of 217 bp while product 246 amplified from the patient sample was considerably smaller (Figure 2D). The experimental 247 normalization, operated by analysis of the housekeeping gene HPRT, did not reveal any 248 difference in SLC9C1 transcripts amount between the patient and control individual. 249 250 Sequencing of SLC9C1 RT-PCR amplicons indicated the total absence of exon 22 in the 251 transcripts from the patient (Figure 2E). This exon encodes for a region included in the Cyclic Nucleotide-Binding Domain (Figure 2C) and comprises 99 bp. Such exon skipping would 252 therefore cause a 33 amino acids in-frame deletion (corresponding to amino acids 884-916). 253 These results confirm the deleterious impact of the identified variant. 254

We next performed immunofluorescence assays with two antibodies raised against different epitopes of the SLC9C1 protein: a so-called 'Central antibody', which recognizes amino-acids 600-750 of the protein, spanning the VSD, upstream from the deletion and a socalled 'C-Terminal antibody' targeting amino-acids 882-932, in the CNBD and mainly coinciding with the deleted sequence. We first observed that both antibodies detected the SLC9C1 protein along the sperm flagellum with a signal intensity decreasing towards the tail end. Importantly, in consistence with RT-PCR data which predicted the deletion of amino acids
884-916, immunofluorescence assays performed with the 'Central antibody' marked SLC9C1
protein in spermatozoa from both control and patient samples (Figure 3A), whereas the 'CTerminal antibody' whose epitope largely overlaps with the deleted region only marked the
control sperm (Figure 3A).

Lastly, we performed western blot immunodetection from control and patient sperm protein extracts. Using the 'Central antibody' we detected a band at the expected molecular weight of the SLC9C1 longer isoform (135 kDa) in both control and patient sperm samples (Figure 3B). Technical limits of the gel electrophoresis did not permit to evidence the small size difference between the normal and the deleted protein but this data firmly confirm the presence of the SLC9C1 truncated protein in sperm cells from the studied patient.

272 **Discussion**

273 In conclusion, we characterize an infertile patient displaying functional asthenozoospermia caused by a homozygous splice mutation in SLC9C1, which generates a 33 274 275 amino-acid deletion within the Cyclic Nucleotide-Binding Domain (CNBD). This protein domain is considered to be involved in the interaction and reciprocal modulation between 276 SLC9C1 (sNHE) and the soluble Adenylate Cyclase (sAC) in mouse and sea urchin sperm^{26,27}, 277 the latter being a key element for the regulation of sperm motility and capacitation. 278

In mice, the disruption of *Nhe-1* did not impact male fertility²⁸ while disruption of 279 Slc9c1 (sNhe) as well as Nha1 and Nha2 (single and double conditional KO mice) all resulted 280 in a phenotype of null sperm motility. Interestingly, these phenotypes were significantly 281 rescued by sperm incubation with cAMP analogues, confirming that Na⁺/H⁺ family members 282 are required for the activation of sperm motility and capacitation through regulation of the 283 cAMP-PKA signalling pathway^{25,29}. SLC9C1 (sNHE) was previously described to only localise 284 to the principal piece of the mouse sperm flagellum²⁵. Zhang et al.³⁶ proposed the same 285 localisation in human sperm cells, although the staining was weak and quite heterogeneous. 286 Our immunostaining assays carried out with two different antibodies suggest a distinct flagellar 287 profile in human sperm. Hence, while we detected a prominent localisation of sNHE in the 288 principal piece of murine spermatozoa (Supplementary Figure S2), in human sperm, SLC9C1 289 was detected along the whole length of the flagellum. 290

Regarding, the molecular mechanisms of action, it is well established that intracellular pH regulation is fundamental at different steps of sperm maturation and capacitation³⁰, and Na⁺/H⁺ exchangers (NHE) play a remarkable role in sperm alkalinisation ³¹. SLC9C1 (sNHE) was shown to be required for pH homeostasis and suggested to promote CatSper activation following Slo3-mediated sperm hyperpolarization during mouse sperm capacitation^{25,32}. In humans, to date no information is available regarding sNHE possible implication in sperm pH homeostasis. Some elements have pointed a role of the proton channel $Hv1^{33,34}$, although so far, no mutation in Hv1 were reported in human asthenozoospermia. Our study therefore sheds light on the essential role of sNHE and a likely similar function in pH homeostasis in humans. Importantly, our findings are in line with a recent comparative genomic analysis, which identified the molecular trio sNHE-sAC-CatSper as an evolutionary conserved machinery for the regulation of sperm flagellar beating in Metazoa³⁵.

The involvement of *SLC9C1* in human sperm physiology has been poorly investigated, 303 and so far only weak evidence was provided regarding its dysfunction in human 304 asthenozoospermia 17,36 , 37 . The mutation in *SLC9C1* we described here constitutes a solid 305 demonstration of SLC9C1 involvement in the regulation of human sperm motility and 306 fertilization potential. We show that this mutation has a drastic impact on sperm progression; 307 however sperm motility was not totally abrogated as observed in *Slc9c1* KO mouse model²⁵. 308 This difference might be explained by the nature of the mutations in those two models; the KO 309 310 mouse model totally lacks the protein while the patient carries a truncated protein, which could 311 preserve partial functionality. In addition, while in patient sperm we observed flagellar angulation, excess residual cytoplasm and overall a flagellar fragility upon sperm selection, 312 such defects were not reported in the KO mouse model. These observations in patient sperm 313 are compatible with a dysregulation of the ion equilibrium and overall cell volume regulation, 314 which were previously described to depend on channels and aquaporins activity³⁸. 315

Overall, this article provides a new evidence of the physiological importance of ion exchange regulation and pH homeostasis in the control of sperm motility and fertilization potential. The first and most remarkable evidence was provided for the CATSPER channel, for which mutations (*CATSPER 1-2-* ε subunits) were identified in several asthenoteratozoospermic individuals with altered fertilizing capacity and/or sperm with coiled flagella^{12,39}. More

recently, pathogenic variants affecting members of the SLC26 family of anion exchangers, 321 namely SLC26A3 and SLC26A8 (TAT1), were also shown to impair the functionality of the 322 Cystic Fibrosis Transmembrane conductance Regulator (CFTR) channel, causing 323 asthenozoospermia, associated with defective capacitation, midpiece and annulus 324 disorganization in the case of $SLC26A8^{11,40,41}$. The voltage-dependent anion channels VDAC2325 and 3 were also identified with pathogenic defects leading to idiopathic asthenozoospermia ⁴² 326 ⁹. The specific location of all those ion channels/transporters at the plasma membrane of sperm 327 cells, constitutes an interesting cellular background for potential therapeutic strategies to 328 improve sperm motility in asthenozoospermic men. 329

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336

337 <u>Author's roles</u>

338 AT designed the study. ED and CP recruited the patient, performed clinical analysis and characterization. AC and PS carried out the ICSI procedure for the patient and his partner. 339 NTM, CC and PR performed exome and bioinformatics analysis. EC did experimental work 340 (RT-PCR, Sequencing, Immunofluorescence, Immunoblotting, MO analysis), 341 data quantification and analysis. AT did TEM analysis. MW, LS and PL contributed in setting up 342 the experimental conditions. EC and AT performed data illustrations. EC and AT analysed the 343 data and wrote the manuscript. MW, AC, PL, CC, CA, PR, NTM, CP and ED did critical 344 reading of the manuscript. 345

346

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354 Data Availability Statement

- 355 The data that support the findings of this study are available from the corresponding author,
- 356 [AT], upon reasonable request.
- 357
- 358

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	General Semen Characteristics					Flagellar defects					Midpiece defects	Head defects		Acrosome defects		
	Volume (ml)	Sperm concentration (10 ⁶ /mL)	Total Motility	Progressive Motility	Vitality	Typical forms	Absent	Short	Irregular	Coiled	Bent	Excess residual cytoplasm	Tapered	Thin	Post- acrosomal	Acrosomal
	3.9	117.5	25	5	49	26	1	1	1	5	19	14	0	3	53	35
	3.4	52	30	5	62	23	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
	3.5	74	35	5	82	10	3	3	0	5	10	1	3	36	22	78
	2.4	73	15	0	80	15	0	2	0	3	6	10	2	25	45	80
	3.8	103	25	5	67	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
mean	3.4	83.9	26	4	68	18.5	1.3	2	0.3	4.3	11.6	8.3	1.7	21.3	40	64.3
Reference limits ^a	1.5	15	40	32	58	23	5	1	2	17	13	4	3	14	42	60
	(1.4-1.7)	(12-16)	(38-42)	(31-34)	(55-63)	(20-26)	(4-6)	(0-2)	(1-3)	(15-19)	(11-15)	(3-5)	(2-4)	(12-16)	(39-45)	(57-63)

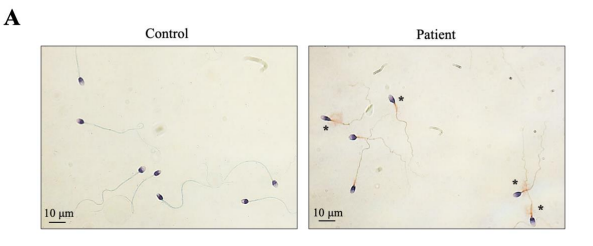
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Table 1 – Semen parameters and sperm morphological defects (flagellum, head and acrosome) of the patient carrying a mutation in *SLC9C1* gene
 Five semen evaluations were performed for the patient over 3 years. Values are expressed in percent, unless specified otherwise. ND: not
 determined.

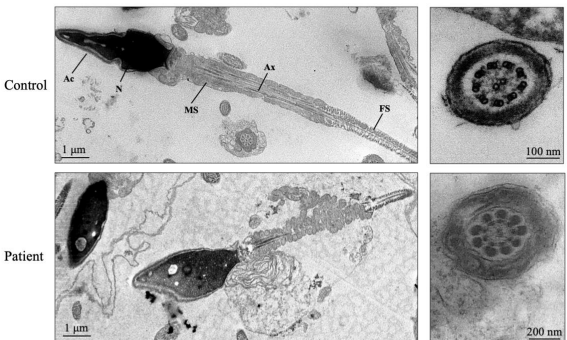
^a Lower or upper reference limits (5th centiles and their 95% confidence intervals) according to the World Health Organization (WHO) standards
 and the distribution range of morphologically abnormal spermatozoa observed in fertile individuals^{18,19}; in bold characters: abnormal values.

<u>Figures</u>

Figure 1







493 Figure 1 – Morphological and ultrastructural anomalies of spermatozoa from the patient.

(A) Morphology of spermatozoa from the patient (right) compared to spermatozoa from a 494 control individual (left). Semen analysis showed a fraction of spermatozoa carrying 495 496 disorganised midpiece with residual cytoplasm and plasma membrane lysis (*). Scale bars represent 10 µm. (B) Transmission Electron Microscopy analysis of semen sample from a 497 control individual (top) and from the patient (bottom), showing flagellar longitudinal and 498 499 transversal sections. Spermatozoa from the patient showed residual cytoplasm associated with plasma membrane lysis and irregular mitochondrial sheath. The transversal sections of 500 501 spermatozoa from the patient showed normal ultra-structure with the (9+2) pattern. Scale bars 502 represent 1µm, 100 and 200 nm. Ac, acrosome; N, nucleus; MS, mitochondrial sheath; Ax, axoneme; FS, fibrous sheath. 503

Figure 2

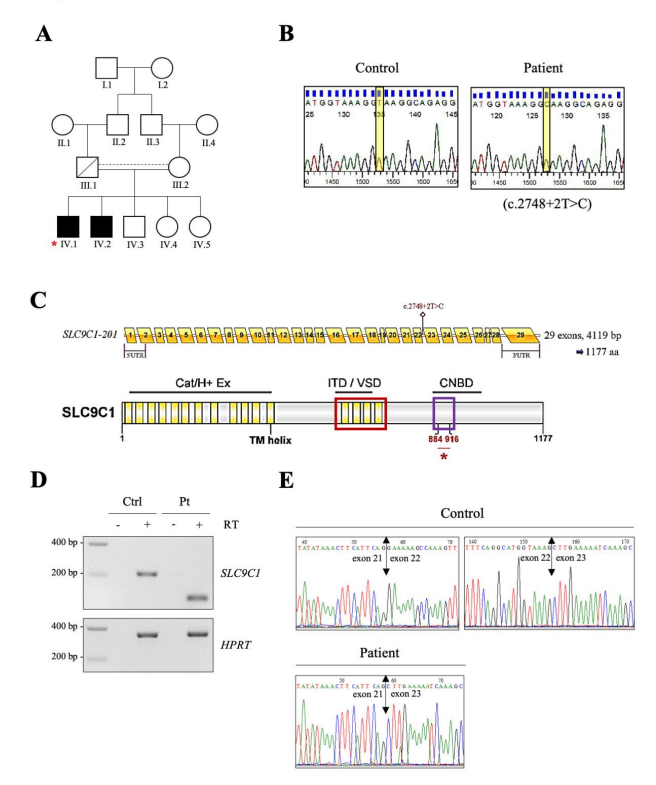
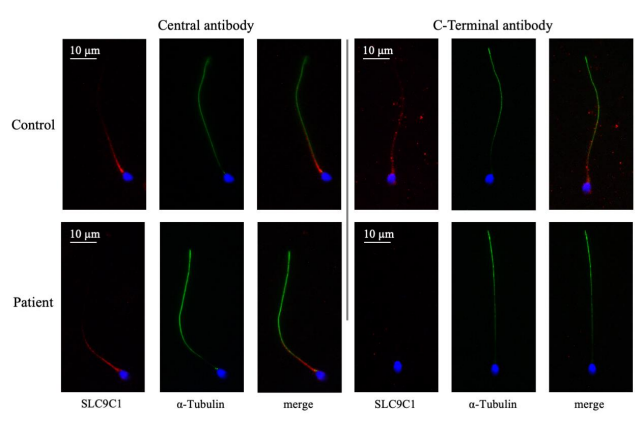


Figure 2 – Pedigree of the patient, description of the *c.2748+2T>C* mutation in *SLC9C1*and analysis of the transcript in sperm from the patient.

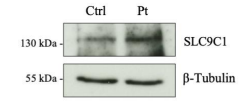
508 (A) Pedigree of the infertile patient with functional asthenozoospermia. The proband (IV.1) is 509 indicated by the red asterisk. Consanguinity between parents (III.1 and III.2) is indicated by the dashed line. One of the brothers of the proband (IV.2) is also known to be infertile; the second 510 one (IV.3) has not manifested desire of parenthood yet. The two sisters of the proband (IV.4 -511 512 IV.5) are fertile and had children. A cousin not illustrated in the pedigree was also reported by the proband to be infertile. (B) Electropherograms of SLC9C1 Sanger sequencing for patient 513 and control individual. The c.2748+2T>C SLC9C1 mutation identified by WES was confirmed 514 515 by Sanger sequencing. (C) (top) Linear structure of SLC9C1 main transcript (SLC9C1-201) showing the exons (yellow boxes), according to the gene description available from Ensembl 516 database. The localization of the mutation is pointed by a stick [figure obtained with IBS tool 517 of DOG software⁴³]. (bottom) Linear structure of SLC9C1 protein (SLC9C1-201 isoform). The 518 yellow boxes represent transmembrane (TM) helices and the horizontal bars indicate all 519 520 functional domains: Cat/H⁺ Ex, Cation/H⁺ Exchanger domain; ITD / VSD, Ion Transport-like 521 Domain / Voltage-Sensing Domain; CNBD, Cyclic Nucleotide-Binding Domain. Red and purple boxes highlight the epitopes of the two antibodies used in the study, the 'Central 522 523 antibody' and the 'C-Terminal antibody' respectively. The region between amino-acids 884 and 916 is encoded by exon 22 [figure obtained with DOG software⁴⁴]. (D) RT-PCR analysis 524 of a semen sample from the patient carrying the c.2748+2T>C mutation. The expected 525 amplicon size is 217 bp for SLC9C1 transcript and 352 bp for HPRT. RNA samples treated in 526 527 absence of Reverse Transcriptase (- RT) constituted negative experimental controls. (E) 528 Electropherogram of SLC9C1 transcript sequencing for a control individual and the patient. In the sample from the patient, sequencing indicates the absence of exon 22 due to the 529 c.2748+2T>C mutation, which abrogates intron 22 splice donor site. 530

Figure 3





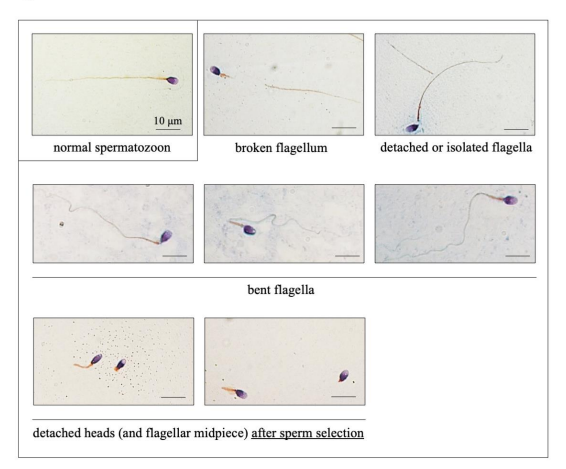
B



532 Figure 3 – Analysis of SLC9C1 protein in sperm from the patient carrying the 533 *c.2748+2T>C* mutation.

(A) SLC9C1 protein detection (red) by immunofluorescence assay in sperm cells from control 534 535 individual (top) and patient (bottom). Two different antibodies were used: one antibody directed against the central region of SLC9C1 and a second antibody directed against the C-Terminal 536 region and including exon 22 coding region. Co-staining with α-Tubulin antibody (green) was 537 538 also performed. The protein localises at the midpiece and principal piece of sperm flagella, with a decreasing gradient towards the end of the tail. The Central antibody allowed to confirm 539 presence of SLC9C1 in patient spermatozoa, while absence of signal of the C-Terminal 540 541 antibody confirmed an abnormal protein lacking part of the CNBD. Scale bars represent 10µm. (B) Immunoblotting detection of SLC9C1 protein (135 kDa) in sperm lysate from control 542 543 individual (left) and patient (right). β-Tubulin (55 kDa) was used for normalization.

Figure S1



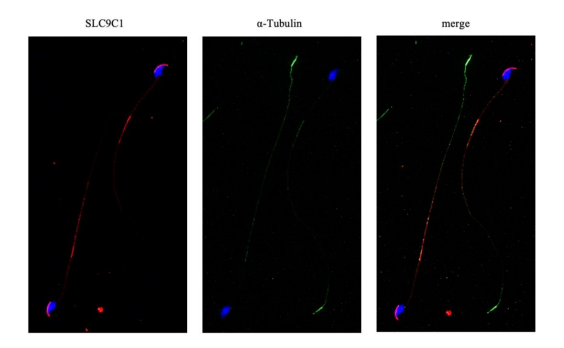
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Figure S1 – Details of the flagellar morphological anomalies observed in sperm from the patient.

Catalogue of the sperm flagellar defects observed in semen from the patient carrying the c.2748+2T>C mutation, compared to a normal spermatozoon (top, left corner). Various flagellar angulation and break points were observed at the level of the connecting piece or along the flagellum, suggesting a fragility of the sperm tail. This phenotype was even more evident after sperm selection with density gradient centrifugation. Scale bar represents 10 μ m.

554

Figure S2



556

557 Figure S2 – SLC9C1 detection in mouse spermatozoa.

Immunofluorescence detection of SLC9C1 protein on mouse spermatozoa using the antibody
directed against the central region of SLC9C1, co-staining with α-Tubulin (green). SLC9C1
(red) preferentially localises at the principal piece of the flagella. Scale bar represents 10µm.

Gene	Testis/all tissue gene expression ratio	Variant	Frequency (gnomAD database)	Nature of the variant	Predicted effect at protein level	Detection in sperm proteome (Wang <i>et al.</i> , 2013)
SLC9C1	12.71	c.2748+2T>C	4.576 10-5	Splice donor variant	-	+
ABCB5	2.37	c.3583C>T	0.0008645	Stop gained	p.Gln1195Ter	-
ZNF891	1.71	c.9del	0.003403	Frameshift variant	p.Met4TrpfsTer9	-

Supplementary Table 1 – Homozygous variants with high pathogenic impact proposed for the patient by bioinformatic analysis of WES data.