Extracellular domains of CATSPER_ε are essential for CatSper assembly during development and activity modulation in sperm capacitation

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Abstract

The flagellar-specific Ca²⁺ channel CatSper is a multiprotein complex that is critical for successful fertilization by controlling the sperm Ca²⁺ signaling in space and time. Large extracellular domains (ECDs) of four single-pass transmembrane subunits, CATSPER β , v, δ , and ϵ , form a unique canopy structure over the pore-forming channel. However, the molecular mechanisms of canopy assembly during development and its physiological function in mature sperm remain unknown. Here, using two genetic mouse models and the biochemical isolation of a bioactive CATSPER fragment, we report that CATSPER ECDs are essential for assembling the CatSper canopy, and thus the entire channel complex, and for modulating CatSper function for sperm hyperactivation and fertilization. CATSPERE-deficient males are sterile because their sperm fail to develop hyperactivated motility due to the absence of the entire channel. In transgenic mice overexpressing CATSPERE with truncated ECDs in testicular germ cells, truncated CATSPERE is unable to interact with native CatSper subunits and incorporate into the complex, thus failing to rescue the defective sperm hyperactivation and infertility of Catspere-null males. These findings provide insight into the underlying molecular and developmental mechanisms of CatSper complex assembly and how CatSper channels can be modulated in physiological settings and by therapeutic intervention.

Keyword: CatSper, Sperm, Hyperactivation, Fertility, Channel assembly, Extracellular domains

Introduction

All living organisms enclosed by the plasma membrane communicate with the extracellular environment through membrane-associated proteins on their surface, such as membrane receptors and ion channels. The extracellular domains (ECDs) of these proteins are critical for sensing and transducing external signals into the cells. Typical examples are AMPA and NMDA receptors, where ligand binding to the ECDs triggers intracellular signaling (**Chen et al., 2017; Chou et al., 2024**). ECDs of various ion channels including PIEZO1 and Ca_v1.2, often modulate ion conductance through the pore by undergoing conformational changes, thus participating in coordinating the opening or closing of the pore and influencing its activated or inactivated state (**Gao et al., 2023; Xiao, 2024**). Based on the structural information and functional importance, recent studies have reported various drugs targeting these ECDs to modulate the activity of ion channels and receptors and the resulting downstream events (**Huang et al., 2024**). Notably, diazepam, which is used to treat anxiety and seizures, targets ECDs of GABA_A receptors for its potentiation of GABA response to suppress neuronal activity (**Zhu et al., 2018**).

In mammalian spermatozoa, various membrane proteins are housed in the flagellar membrane for their navigation to successful fertilization in the female reproductive tract (**Wachten et al., 2017; Wang et al., 2021**). Among them, the multiprotein CatSper Ca²⁺ channel complex presents the largest ECDs. Each of the four single transmembrane accessory subunits (CATSPER β , γ , δ , and ε) interacts with one specific pore-forming subunit, forming a huge canopy structure over the heterotetrameric channel and linking the individual holo-complexes extracellularly in zigzag rows (**Lin et al., 2021; Zhao et al., 2022**). CatSper-mediated Ca²⁺ signaling is critical for nearly all events in sperm capacitation (*i.e.*, the collective process of acquiring fertility), including the development of hyperactivated motility and the acrosome reaction, and thus male fertility (**Hwang and Chung, 2023**). However, our understanding of the physiological function of the CatSper canopy and the outcome of its intervention as a drug target remains largely unexplored.

Our previous study showed that CATSPER δ is required to maintain the stability of a pore-forming CATSPER1 subunit in developing male germ cells, suggesting its role in channel assembly (Chung et al., 2011). Indeed, recent advances in structural elucidation of the CatSper holocomplex and *in situ* higher order arrangement further support that the canopy-forming subunits contribute not only to channel complex assembly but also potentially to activity regulation (Lin et al., 2021; Zhao et al., 2022). Because the entire CatSper channel was absent in mature spermatozoa from Catsperd-null males (Chung et al., 2011), the detailed molecular mechanisms of assembly and the function of canopy ECDs in mature sperm remain unknown. This critical hurdle highlights the need for a different approach to further elucidate the physiological function of the canopy and individual TM subunits in mature spermatozoa. In the CatSper holo-complex, CATSPERE, paired with the CATSPER2 subunit (Lin et al., 2021), is located on the wing side of the CatSper zigzag rows (**Zhao et al., 2022**) in the guadrilinear Ca²⁺ signaling nanodomains (Chung et al., 2017; Hwang et al., 2019). Interestingly, among the CatSper genes, human mutations in CATSPERE and its pairing partner CATSPER2 have been reported more frequently (Brown et al., 2018; Young et al., 2024), suggesting that their structural location is potentially vulnerable to cause functional defects in the CatSper channel and thus male infertility.

Here, using genetic and molecular dissection of CATSPER_ε, we show the molecular mechanisms of CATSPER_ε ECDs in channel complex assembly during germ cell development and their physiological function in mature spermatozoa. Genetic ablation of *Catspere* impairs sperm to

develop hyperactivated motility as it leads to the absence of the entire CatSper complex, resulting in male infertility. Overexpression of the canopy roof-truncated CATSPERs prevents its interaction with other CatSper canopy-forming subunits in testicular germ cells. Finally, we demonstrate that treatment of spermatozoa with purified Ig-like domain of CATSPERs ECDs during capacitation attenuates CatSper activity, impairing sperm hyperactivation and fertilization *in vitro*. These findings provide direct insights into the canopy function in CatSper complex assembly and activity regulation during sperm capacitation, and suggest a strategy to target CatSper activity to control sperm fertility.

Results

CATSPER_ε is indispensable for sperm hyperactivation and male fertility

Four single-pass transmembrane (TM) subunits (CATSPER β , y, δ , and ϵ) with their large extracellular domains (ECDs) form the canopy structure of the CatSper channel complex (Lin et al., 2021). The channel complexes are arranged in zigzag nanodomains along the sperm tail with CATSPERs uniquely positioned on the wing side (Zhao et al., 2022; Figure 1A and 1B). However, the detailed roles of CATSPERs and its physiological significance are currently unknown. Like other CATSPER subunits, Catspere is expressed exclusively in the testis (Figure 1C; Figure S1A), and CATSPERE is specifically localized to the flagellar membrane in the principal piece of human and mouse spermatozoa (Chung et al., 2017: Figure 1D and 1E). To understand the physiological significance of CATSPER_ε, we generated Catspere-null (Catspere^{-/-}) mice using CRISPR/Cas9 genome editing (Figure S2A and S2B). By introducing two guide RNAs targeting the 1st and 2nd exons, respectively, we obtained a null allele predicted to produce a frame-shifted protein after the 3rd amino acid. CATSPER_ε is absent in the testis and epididymal spermatozoa of Catspere^{-/-} males (Figure 1E and F; Figure S1B; Figure S2C). Catspere^{-/-} mice show no gross abnormalities in survival, appearance or behavior, and null females are fertile. However, despite normal sperm production (Figure S2D), Catspere^{-/-} null males are completely infertile (Figure 1G and S2E), as are genetic disruptions of genes encoding other TM CatSper subunits (Ren et al., 2001; Quill et al., 2003; Qi et al., 2007; Chung et al., 2011; Huang et al., 2023; Hwang and Chung. 2023). To understand how CATSPERs deficiency leads to male infertility, we examined sperm motility parameters using a Computer-Assisted Semen Analyzer (CASA). Curvilinear velocity (VCL) and lateral head displacement (ALH) are significantly lower in sperm lacking CATSPERE after induction of capacitation (Figure S2F). Flagellar waveform analysis also revealed that Catspere^{-/-} sperm fail to increase the maximum angle of the primary curvature in the midpiece (α -angle) and beat in an overall smaller envelope (Figure 1H and 1I), all indicative of defective sperm hyperactivation. These results demonstrate that CATSPER_E deficiency impairs hyperactivated motility, resulting in male infertility.

Catspere-null sperm lack the entire CatSper channel, resulting in impaired Ca²⁺ signaling

To understand how genetic ablation of *Catspere* impairs sperm to develop hyperactivated motility, we examined whether the CatSper channel is present and functional in *Catspere*^{-/-} sperm (**Figure 2**). All CatSper subunits examined are absent in *Catspere*^{-/-} sperm (**Figure 2A and B**), suggesting the lack of the entire CatSper channel, consistent with the observation in *Catsperd*^{-/-} sperm (**Chung et al., 2011**). *Catspere*^{-/-} sperm prematurely enhance capacitation-mediated tyrosine

phosphorylation compared with wild-type (WT) sperm (**Figure 2C and Figure S2G**), indicating impaired Ca²⁺ influx during capacitation (**Chung et al., 2014**). To directly test for defective Ca²⁺ influx through the CatSper channel, we performed electrophysiological recordings to measure the CatSper current (I_{Catser}) in *Catspere^{-/-}* spermatozoa (**Figure 2D-F**). When elicited under voltage ramp or step protocols, $I_{Catsper}$ is undetectable in *Catspere^{-/-}* spermatozoa, indicating the absence of functional CatSper channels in spermatozoa. These results demonstrate that *Catspere^{-/-}* spermatozoa lack the CatSper channel and therefore fail to transduce Ca²⁺ signals to develop hyperactivated motility during capacitation.

Truncated CATSPERε lacking the extracellular canopy roof does not interfere with the assembly of native full-length CATSPERε into the CatSper complex

The function of the unique large extracellular canopy above the pore-forming channel remains unknown. It could serve as a binding site for natural ligands, as a linker to multimerize the channels into a linear cooperative working unit, and/or as a chaperone to assemble the pore-forming tetramer in the specific counterclockwise order (Lin et al., 2021; Zhao et al., 2022). Notably, sequence comparison analysis revealed that the C-terminal Ig-like domain and the stem region of CATSPER ϵ are highly homologous to those of CATSPER δ (Chung et al., 2017; Figure S3A). In contrast to CATSPER β and γ , which mediate intra- and inter-dimer interactions, respectively, CATSPER ϵ and δ do not participate in the supramolecular interactions in the higher-order arrangement of the CatSper complexes (Zhao et al., 2022). Thus, we hypothesized that this homology at the level of the canopy pole (*i.e.*, the stem and TM region only without the canopy roof) is more indicative of its function in channel assembly. To test this idea at the canopy pole and to further delineate the specific role of the canopy roof, we generated a transgenic mouse line overexpressing the roof-truncated, pole-retaining CATSPER ϵ (*i.e.*, truncated CATSPER ϵ) specifically in the testis using the calmegin promoter (Watanabe et al., 1995; Ikawa et al., 2001; Figure 3A and Figure S3A).

The truncated CATSPERs is expressed in testis and is enriched in the microsomal fraction, indicating its membrane association (Figure 3B and 3C). However, the truncated protein is not detected in epididymal spermatozoa (Figure 3B and 3D). At the same time, the presence of the truncated CATSPERs in testis did not alter the typical guadrilinear localization of the CatSper channel in the principal piece of mature sperm tail (Figure 3D and 3E). These results demonstrate that only the CatSper channel complexes assembled without the truncated CATSPERs are properly trafficked to the flagellar membrane. To test this idea, we first determined whether the truncated CATSPER is complexed with other CatSper subunits. We performed coimmunoprecipitation using testis lysate (Figure 3F). Native full-length CATSPER and other examined CatSper subunits are detected in the CATSPERδ immunocomplex in the testis, independent of the transgene expression encoding the truncated CATSPERE. However, the immunocomplex of the truncated CATSPERs protein with anti-HA antibody does not contain any CatSper subunits in the transgenic animal. This result demonstrates that, contrary to our hypothesis, the canopy pole is not sufficient to assemble the canopy structure but the roof-forming ECDs of CATSPER are also necessary for the assembly of the CatSper channel holo-complex. Without this region, the truncated CATSPER ε is retained in the cell body and progressively depleted in developing spermatids (Figure 3G), supporting its absence in mature spermatozoa (Figure 3B and 3D) and minimal effect on the assembly of native CATSPERs into the CatSper complex (Figure 3F). Accordingly, spermatozoa from the animal carrying both WT and transgenic

alleles (WT; TG+) develop normal CatSper-mediated hyperactivated motility with normal tyrosine phosphorylation after inducing capacitation (**Figure S3B and S3C**). Also, protein levels of the native CatSper subunits are not altered in WT; TG+ spermatozoa (**Figure S3D and S3E**).

The canopy roof-truncated CATSPERε fails to form heterotetrameric auxiliary complex in *Catspere*-knockout mice

In the absence of full-length CATSPER ε , the truncated CATSPER ε could replace the assembly function of native CATSPER ε by incorporation into the CatSper complex, while not being dominantly or equally in the presence of full-length CATSPER ε (**Figure 3**). To test this idea, we crossed the transgenic mice in the background of *Catspere^{-/-}* mice to express only the transgene (*Catspere^{-/-}*; *TG*+; **Figure 4A**). The truncated CATSPER ε proteins are well expressed in the testis but not detected in the epididymal spermatozoa as observed in the spermatozoa of WT; *TG*+ males. CATSPER δ co-immunoprecipitation with testis lysate shows that truncated CATSPER ε is not in complex with any of the native CatSper subunits examined, not even with the neighboring CATSPER δ or γ (**Figure 4B**). This result demonstrates that the ECDs corresponding to the canopy roof of CATSPER ε are indispensable for the assembly of all four auxiliary subunits into a fourfold complex via its interaction with CATSPER δ and γ .

Next, to understand how the CATSPER ϵ truncation affects the assembly of the CatSper canopy and the entire CatSper complex, we analyzed the AlphaFold predicted interactions of truncated CATSPER ϵ with δ or γ (**Figure 4C-E**). In the predicted models, the intact CATSPER ϵ ECDs interact normally with both δ and γ , and the predicted ϵ - δ and ϵ - γ dimers fit well into the atomic model of the CatSper canopy (CATSPER γ - β - δ - ϵ from PDB:7EEB) (**Figure 4D and E**). However, the truncated CATSPER ϵ ECDs (*i.e.*, C-terminal Ig-like domain and stem region without TM domain) are predicted to interact abnormally with these neighboring subunits, and the aberrant dimers do not fit the reported structure. Thus, the truncated roofless partial ECDs are likely to have altered their interaction with other canopy-forming auxiliary subunits, resulting in unsuccessful canopy assembly and canopy interactions with the pore-forming channel (CATSPER1-4-3-2).

The truncated CATSPER ϵ fails to rescue impaired sperm hyperactivation by CATSPER ϵ deficiency

Truncation of the canopy roof alters the interaction of CATSPER ε with δ and γ , impairing ε incorporation into the canopy and thus into the entire CatSper channel complex (**Figure 3 and 4**). In addition, the truncated CATSPER ε appears to have little effect on the assembly and trafficking of the CatSper channel complex in the presence of native full-length CATSPER ε in sperm from both WT and *Catspere*^{+/-} males (**Figure S3, S4A-S4E**). Thus, we predict that *Catspere*^{-/-;} *TG*+ spermatozoa will lack a functional CatSper channel and will not rescue the physiological defects of *Catspere*^{-/-;} animals.

Indeed, all CatSper subunits examined are absent in *Catspere^{-/-}*; *TG*+ spermatozoa (**Figure 5A and 5B**, **Figure S4F**). Capacitated *Catspere^{-/-}*; *TG*+ spermatozoa aberrantly potentiate tyrosine phosphorylation (**Figure S4G**), indicating impaired CatSper-mediated Ca²⁺ signaling. Electrophysiological recording shows that $I_{Catsper}$ is not detectable in *Catspere^{-/-}*; *TG*+ spermatozoa (**Figure 5C-E**), which is not significantly different from that recorded in *Catspere^{-/-}* spermatozoa (**Figure 2D-F**). Although *Catspere^{-/-}*; *TG*+ males produce a comparable number of spermatozoa as heterozygous males (**Figure S4H**), they are infertile (**Figure 5F and S4I**) with defective sperm

hyperactivation (**Figure 5G and S4J**) like *Catspere*^{-/-} males (**Figure 1**). All these results demonstrate that *Catspere*^{-/-}; *TG*+ sperm lack the CatSper channel and that truncated CATSPER ϵ cannot rescue male infertility caused by CATSPER ϵ deficiency.

Ig-like domain of CATSPER ϵ is critical in modulating CatSper activity and sperm hyperactivation

Our results using two *Catspere* genetic models provide strong evidence that the CATSPERε ECD regions that form the canopy roof are critical for their interaction with neighboring subunits to assemble the canopy and thus the CatSper channel holo-complex in developing male germ cells (**Figures 3, 4, and 5**). However, it remains to be determined whether and to what extent the canopy functions physiologically in mature spermatozoa, where the channel complexes are arranged in a higher order (**Zhao et al., 2022**).

The Ig-like domain, which is present in all four canopy-forming subunits, mediates extensive interactions between them at the center of canopy roof, whereas the stem and transmembrane domains do not (**Lin et al., 2021**). Consistently, our modeling also predicts that among the individual ECDs of CATSPER ϵ (*i.e.*, NTD, BPD, and Ig-like), only the Ig-like domain is predicted to interact with the δ and γ ECDs at the canopy roof (**Figure S5A**), suggesting its importance in stabilizing the canopy structure. Furthermore, our preliminary solubility test of the serially deleted recombinant CATSPER ϵ mutant proteins showed that the Ig-like domain (CATSPER ϵ ^{Ig-like}; **Figure 6A**) is more soluble than other parts of CATSPER ϵ (**Figure S5B and S5C**). Therefore, we purified recombinant mCherry-tagged CATSPER ϵ ^{Ig-like} expressed in CHO cells (**Figure 6B**) to test whether it could affect sperm physiology when administered exogenously.

We found that CATSPER $\epsilon^{Ig-like}$ remained stably bound to live spermatozoa in a CatSperdependent manner when the proteins were added to sperm incubation under capacitating conditions (**Figure 6C and D**). These results demonstrate that the soluble CATSPER $\epsilon^{Ig-like}$ fragments can directly interact or intercalate with the canopy roof of the native CatSper channel complex. Recombinant CATSPER $\epsilon^{Ig-like}$ treatment reduces motility parameters associated with sperm hyperactivation dose-dependently without affecting total motility (**Figure 6E**), suggesting reduced CatSper activity. Therefore, we performed a spectrofluometeric sperm membrane potential (Em) assay on sperm loaded with DiSC₃(5) (**Luque et al., 2023**). CATSPER $\epsilon^{Ig-like}$ application during sperm capacitation blocks the previously reported CatSper-specific, EGTAinduced Em depolarization by more than 70% (**Figure 6F and G**). Under the same conditions, CATSPER $\epsilon^{Ig-like}$ treated spermatozoa inhibited fertilization rate of COC eggs *in vitro* by ~ 50% (**Figure 6H**). Thus, we isolate the Ig-like domain of CATSPER ϵ as a region important for modulating CatSper function. Our results strongly support that altering protein-protein interactions at the level of the CatSper canopy roof can prevent capacitation-induced opening of the CatSper pore-forming channels, thereby compromising sperm fertilization ability.

Discussion

CATSPER_E ECDs are essential for assembly of the CatSper canopy and pore-forming channel

CatSper is a multiprotein channel complex composed of more than 10 TM subunits and 3 cytosolic components, arranged in higher-order zigzag rows (Hwang and Chung, 2023). All reported TM subunit deficiencies in mice result in the absence of a functional CatSper channel in mature spermatozoa, leading to impaired male fertility due to the defective hyperactivation (Ren et al., 2001; Quill et al., 2003; Qi et al., 2007; Chung et al., 2011; Huang et al., 2023) despite the presence of all other CatSper subunits in the testis except the target protein (Chung et al., 2017; Hwang et al., 2019; Hwang et al., 2022; Huang et al., 2023). Thus, assembly defects at various steps from the assembly of individual holo-complexes (composed of canopy and pore-forming channels) to their higher-order zigzag arrangement can be attributed to the absence of functional CatSper channels in mature sperm from these TM knockout males (Chung et al., 2011; Hwang et al., 2022; Huang et al., 2023). We have previously shown that CATSPERθ deficiency allows transient assembly of single holo-complexes, but results in defective interaction within CatSper dimers (Huang et al., 2023). The defective intra-dimer interaction prevents CatSper holocomplexes from exiting the cell body and trafficking to flagella of developing male germ cells (Huang et al., 2023). In contrast, the absence of either pore-forming or canopy-forming CatSper TM subunits fails to assemble transient CatSper holo-complexes in the testis (Chung et al., 2011; Hwang et al., 2022). For example, CATSPER1 deficiency leads to incomplete organization of the pore-forming complex, whereas canopy-forming subunits are complexed in male germ cells (Chung et al., 2011; Hwang et al., 2022). Deficiency of CATSPERδ or ε not only results in incomplete canopy assembly, but also reduces the protein levels of the pore-forming subunits, presumably due to their altered stability (Chung et al., 2011; Figure 4). Thus, a deficiency of either pore-forming or canopy-forming TM subunits likely impairs the assembly of the CatSper holo-complexes during spermatogenesis, resulting in the absence of a functional channel in epididymal spermatozoa. However, considering the earlier mRNA expression of canopy-forming subunits than pore-forming subunits during development and their different interaction patterns and protein levels in the testis (Chung et al., 2011; Chung et al., 2017; Hwang et al., 2019), a heterotetrameric canopy structure should form first, which presumably serves as a scaffold for 1:1 pairing with the pore-forming channels. Canopy roof-truncated CATSPER cannot be incorporated into the CatSper complex (Figure 4), highlighting the importance of the protein interactions between individual canopy TM subunits for functional CatSper channel assembly. Therefore, ECD-mediated canopy assembly may be another checkpoint to ensure a complete CatSper holo-complex for flagellar trafficking and functional expression in mature sperm.

Canopy structure is involved in modulating CatSper activity during sperm capacitation

Ion channel ECDs are important for the regulation of channel activity (**Wang et al., 2019**; **Nadezhdin et al., 2021**). Recent advances in cryo-EM techniques have revealed that ECDs undergo conformational changes during gating processes. For example, the ECD of GLIC, a proton-activated prokaryotic channel, is compacted and rotated counterclockwise by protonation, resulting in a widening of the pore region in the TM domains to evoke ion conductance (**Bharambe et al., 2024**). The protonated ECD of another pH-gating channel, TWIK1, closes the channel by sealing the top of the selectivity filter and suppresses K⁺ conductance (**Turney et al., 2022**). The ligand-mediated conformational changes of the ECDs and the associated gating processes have also been reported for several neurotransmitter receptors. AMPA subunits are twisted by ligand binding, resulting in conformational changes of the gating ring at the ECDs and subsequent AMPA activation (**Chen et al., 2017**). Ligand binding to other heteropentameric cys-loop family receptors, such as NMDA and glycine receptors, also induces rotation of the ECDs to propagate pore

opening and channel activation (Liu and Wang, 2023; Chou et al., 2024). All these results show that channel activation is strongly associated with structural changes of ECDs. The current atomic structure of the CatSper holo-complex is a closed conformation resolved using the purified complex from testis and epididymis (Lin et al., 2021). Structural analyses revealed that the single TM domain of each canopy-forming subunit interacts with the S2 segment of the paired poreforming subunit. In addition, their Ig-like domains interact with each other and stabilize the canopy structure at the roof just above the pore. Thus, the canopy structure connected to those of the neighboring channels might prevent the pore from opening and keep the channel in a closed conformation in non-capacitated sperm. Interestingly, exogenous addition of purified recombinant CATSPERE Ig-like domain inhibits CatSper function in sperm incubated under capacitating conditions (Figure 6). This result suggests that the recombinant Ig-like domain interferes with the interaction of the native CATSPERE ECD with those of the adjacent canopy-forming subunits, CATSPERy and δ , and impairs the coordinated conformational changes of the canopy structure and/or supramolecular interaction of the zigzag arrangement during sperm capacitation. Direct structural analysis of the canopy structure in capacitated sperm will clarify this possibility and contribute to a better structural understanding of CatSper gating mechanisms during capacitation.

CATSPERs Ig-like domain is a promising, defined drug target to intervene CatSper function

Worldwide, more than 200 million pregnancies are unintended each year (Nickels and Yan, 2024). Strikingly, more than 50 % of unintended pregnancies eventually end in abortion, resulting in various health and social burdens for mothers and family members (Horvath and Schreiber, 2017; Bearak et al., 2020). Because of the negative consequences of unintended pregnancy, there is a growing awareness and demand for improved and/or diversified contraceptive methods. Especially, men currently have very limited options such as condoms and vasectomy. To this end, the recently reported plakogloblin-SPEM1 interaction inhibitor triptonide (Chang et al., 2021) and soluble adenylyl cyclase inhibitor TDI-11861 (Balbach et al., 2023) are promising candidates for reversible male contraceptives that have been screened to target proteins specifically involved in spermiogenesis and sperm function, respectively.

CatSper is another good target validated for male fertility because of its specific expression and function in regulating sperm motility and fertility (**Mariani et al., 2023; Lee and Hwang, 2024**). However, the multi-component organization and structural complexity of CatSper has hindered the development of CatSper-targeted contraceptives. Interestingly, the large volume of ECDs and the zigzag arrangement of the CatSper channels with a 180° rotation (**Zhao et al., 2022**) suggest a uniquely high accessibility of the wing side of CATSPER ϵ to potential biologics (*see also* the related, accompanying manuscript by **Clouser et al., 2024**). Here we show that treatment with a recombinant CATSPER ϵ Ig-like domain, which would disrupt the protein-protein interaction between CATSPER ϵ and its neighbors CATSPER γ and δ within the holo-complex, inhibits CatSper and thus reduces sperm fertility (**Figure 6**). Thus, small biologics and/or small molecules that specifically bind to CATSPER ϵ ECDs could be potential strategies to intervene in CatSper function for the development of novel non-hormonal contraceptives.

Materials and methods

Animals

Catsper1 and Catsperd-null mice (**Ren et al., 2001; Chung et al., 2011**) are maintained on a C57BL/6 background. WT C57BL/6 and CD1 mice were purchased from Charles River Laboratory. Mice were managed in accordance with the guideline approved by Yale Institute Animal Care and Use Committee (IACUC, #20079).

Generation of Catspere-null and Catspere-TG Mice and Genotyping

Catspere-null mice were generated on a C57BL/6 background using CRISPR/Cas9 system. Two guide RNAs. 5'-CGCCATGTTAGCCCGGCAGGTGG-3' and 5'-AGAAGAACTGCAGCCTCCAGTGG-3', in px330 vector were injected into the fertilized eggs collected from super-ovulated C57BL/6 females after mating with males. The developing 2-Cell embryos were transplanted into pseudopregnant CD1 females and the founders' toe were biopsied for genotyping. Truncation of the target region was examined by gDNA PCR using F2 (5'-ACTGCCCTCGTTAGCTTTTTGTCA-3') and R1 (5'-CCTCCTTGGGCAGTTGTAGTTCA-3') primer pair. PCR products were sanger sequenced and female founders carrying the truncated allele were backcrossed with the WT males to examine germline transmission of the allele. The Catspere-null mice line with 63 kb-gDNA deletion were maintained and genotyping were performed by primer pairs of F2 - R1 for the Catspere-null allele and F1 (5'-GCATACTAATTGCTTGGTCAAAAAC-3') - R1 for WT allele.

Transgenic mice were generated by introducing transgene encoding the truncated CATSPERε (*Catspere-TG*) in *pClgn-mCherry* vector. Cloned *pClgn-Catspere-TG-mCherry* was linearized by restriction enzyme digestion and electroporated into the fertilized eggs collected from C57BL/6. The embryos were transplanted pseudopregnant CD1 females. Founders' toes were clipped for gDNA extraction. Extracted gDNA were subjected for PCR with forward (5'-AAGATTTACAGGCAGTTTATTATTGAG-3') and reverse (5'-GTCGGAGGGGAAGTTGGT-3') primers. *Catspere-null; Catspere-TG* mice were generated by mating *Catspere*-TG male with WT background and *Catspere*-null females.

Human samples (BioBank)

Human materials for the present study were collected in accordance with the principles of the Declaration of Helsinki. The regional medical research ethics committee of the Capital Region of Copenhagen approved the use of human tissues stored in the biobank at the Department of Growth and Reproduction for gene expression studies (permit number H-16019636). All patients have given informed consent to donate their residual tissue for research.

Cell lines

HEK293T cells derived from embryonic kidney (ATCC) were cultured in DMEM (GIBCO)

supplemented with 10% FBS (Thermofisher) and 1x Pen/Strep antibiotics (GIBCO) at 37 $^{\circ}$ C, 5% CO₂ condition. The cultured cells were used to test solubility of the recombinant proteins.

Bacterial strains

NEB 10- β bacterial strain (NEB) was used for molecular cloning.

Mouse Sperm Preparation

Epididymal sperm from adult male mice were collected from caudal epididymis by swim-out in either M2 medium (EMD Millipore) or HEPES saline (HS medium; 135 NaCl, 5 KCl, 1 MgSO4, 2 CaCl₂, 20 HEPES, 5 Glucose, 10 Lactic acid, and 1 Na pyruvate) adjusted to pH 7.4). Collected epididymal sperm were incubated in human tubular fluid (HTF) medium (EMD Millipore) at 2×10⁶ cells/ml concentration under the 37 °C, 5% CO₂ condition for 90 min to induce capacitation.

Mouse Testicular Cells Preparation

Dissociated mouse testicular cells were prepared as previously (**Hwang et al., 2021b; Hwang et al., 2022**). Briefly, seminiferous tubules in adult testes were dissociated mechanically after removing tunica albuginea. The tubules were washed with cold PBS, chopped, and minced to dissociate testicular cells. The minced tubules in PBS were filtered with 40 μ m-mesh cell strainers to remove debris. Filtered testicular cells were subjected to immunostaining.

Human Testis Section Preparation

Human testis tissues were excised from the normal, tumor-free periphery of orchiectomy specimens from four adult patients with testicular germ cell cancer. Only the specimens with tumors typical of young adults, i.e., derived from germ cell neoplasia *in situ* (GCNIS), either seminoma or nonseminoma, were included. Excised specimens were immediately fixed in GR-fixative (7.4% formaldehyde, 4% acetic acid, 2% methanol, 0.57% sodium phosphate, dibasic and 0.11% potassium phosphate, monobasic) for at least 16 hours. Samples were then dehydrated and embedded in paraffin.

Antibodies and Reagents

Rabbit polyclonal anti-mouse CATSPER1 (**Ren et al., 2001**), 3, 4 (**Qi et al., 2007**), β (**Liu et al., 2007**), δ (**Chung et al., 2011**), EFCAB9 (**Hwang et al., 2019**), and anti-human CATSPER ϵ (α -h ϵ -31; **Chung et al., 2017**) were described previously.

Custom, in-house antibodies generated in this study.

Rabbit polyclonal anti-mouse CATSPER ϵ (α -m ϵ -331; see also **Nand et al., 2023** and related manuscript by **Clouser et al., 2024**) was generated by immunizing rabbits with KLH-conjugated peptide (DGTVYLRTEDEFTKLDES; Open Biosystems). Rabbit polyclonal anti-mouse CATSPER γ antibody (α -m γ -957) was generated by immunizing rabbits with the KLH-conjugated peptide (RDYTEEEIFRYNSPLDTTNSLI; BioMatik). From the collected sera, the antibodies are affinity purification using AminoLink coupling resin (Pierce). A mouse monoclonal anti-CATSPER1 (clone 6D3) was established from screening hybridomas generated from mice immunized with purified N-terminus of mouse CATSPER1 (1-150 aa; GenScript) followed by Protein G affinity purification.

Commercially available antibodies.

Mouse monoclonal antibodies against HA (clone HA-7, H3663), acetylated tubulin (clone 6-11B-1, 7451), protein tyrosine phosphorylation (clone 4G10, 05-321), and Calmodulin (05-173) were from Sigma-Aldrich. Mouse monoclonal anti-Na⁺/K⁺ ATPase was from SantaCruz (clone H-3, sc-48345). Mouse monoclonal anti-mCherry was from Novus (Novus NBP1-96752). Rabbit monoclonal anti-HA (clone C29F4, 3724) was from Cell Signaling Technology. Goat anti-mouse (115-035-003) and anti-rabbit (111-035-144) IgG conjugated with HRP were from Jackson ImmunoResearch. Veriblot for immunoblotting of the coIP was from AbCam (ab131366). Alexa 568-conjugated goat anti-mouse (A-11031) and anti-rabbit (A-11036) IgG (H+L), and Alexa 647-conjugated WGA (W32466) were from Invitrogen. Hoechst dye was from TOCRIS (5117).

All other chemicals and reagents were from Sigma Aldrich unless indicated.

Molecular Cloning

A construct to generate *Catspere-TG* transgenic mice was cloned. Mouse *Catspere* which is sequence-homologue to the mouse *Catsperd* was subcloned into the *pClgn-mCherry* vector, gifted by Dr. Masahito Ikawa using NEBuilder® HiFi DNA Assembly Kit (NEB). Sequences encoding the native signaling peptide and HA tag are placed at the 5' region of the *Catspere-TG* sequences. Constructs encoding fragments of the CATSPERε extracellular domains (ECDs) were subcloned into phCMV3 vector using NEBuilder® HiFi DNA Assembly Kit (NEB).

Protein Extraction and Immunoblotting

Epididymal Sperm

Whole sperm proteins were extracted as described previously (Hwang et al., 2022). Collected sperm from cauda epididymis were washed with PBS followed by lysis using 2X LDS buffer for 10 minutes with vortaxing at room temperature (RT). The lysates were centrifuged at 4 °C, 18,000 x g, and supernatants were collected and denatured by incubation with 50 mM DTT at 75 °C for 10 minutes. The denatured sperm proteins were used for SDS-PAGE and immunoblotting. Primary antibodies used are: rabbit polyclonal anti-mouse CATSPER β , CATSPER δ , CATSPER1, CATSPER3, CATSPER4, and EFCAB9 at 1 µg/ml, anti-mouse CATSPER ϵ (1.6 µg/ml), rabbit

monoclonal anti-HA (1:2,000), and mouse monoclonal anti-pTyr (1:2,000) and AcTUB (1:20,000).

Testis

Testis proteins were obtained as previously (Hwang et al., 2022). Testes were homogenized in 0.32M sucrose buffer using dounce homogenizer and centrifuged at 4 °C, 1,000 x g for 15 minutes to discard nucleus and debris. Collected supernatant were either lysed to 1X LDS buffer or subjected to centrifugation at 4 °C, 100,000 rpm for 1 hour to separate microsome and cytosolic fraction. Separated fractions were lysed to 2X LDS volume-equivalently. Lysed testes samples were denatured by heating at 75 °C for 2 or 10 minutes with 50 mM DTT supplement, followed by SDS-PAGE and immunoblotting. Primary antibodies for the immunoblotting were: rabbit polyclonal anti-CATSPERε (2.4 µg/ml) and CATSPER1 (1 µg/ml), rabbit monoclonal anti-HA (1:2,000), and mouse monoclonal anti-Na⁺/K⁺ ATPase (1:2,000), CaM (1:2,000), and AcTub (1:20,000). HRP-conjugated goat-anti mouse or rabbit antibodies were used for secondary antibody with 1:20,000 concentration according to the host species of the primary antibody.

Transfected mammalian cells

Cultured 293T cells were transfected with HA-tagged recombinant CATSPER ϵ ECDs using linear polyethylenimine (40 kDa, Polyscience). Cells were harvested after 36 - 48 hours from transfection followed by washing with PBS two times. The transfected cells were solubilized using 1% Triton X-100 in PBS for 4 hours at 4 °C by gentle rocking and centrifuged at 4 °C, 18,000 x g for 1 hour to pellet insoluble fraction. Soluble proteins were mixed to 1x LDS buffer and insoluble pellet fractions were lysed using 2X LDS buffer volume-equivalently. Proteins in LDS sample buffer were denatured by boiling at 75 °C for 2 minutes with 50 μ M dithiothreitol. Denatured samples were subjected to SDS-PAGE and immunoblotted using mouse anti-HA (1:2,000) to probe recombinant CATSPER ϵ ECD proteins.

Testis Co-Immunoprecipitation

Proteins in testes microsome were solubilized using PBS containing 1% Triton X-100 and cOmplete Mini, EDTA-free Protease Inhibitor Cocktail (Roche) at 4 °C for 3 hours with gentle rocking. The lysates were centrifuged at 4 °C, 18,000 x g for 1 hour and supernatants with solubilized protein were collected. Solubilized microsome proteins were incubated with SureBeadsTM Protein A Magenetic Beads (Bio-Rad) conjugated with rabbit polyclonal anti-CATSPER δ (0.5 µg) or mouse monoclonal anti-HA (0.5 µg) at 4 °C, overnight. The incubated magnetic beads were washed with 1% Triton X-100 in PBS and the immunocomplexes were eluted with 2X LDS buffer containing 50 mM DTT, followed by heating at 75 °C for 10 minutes. The elutes were used for SDS-PAGE and immunoblotting. Used Primary antibodies are: rabbit polyclonal anti-CATSPER δ (2 µg/ml), CATSPER ϵ (2.4 µg/ml), and rabbit monoclonal anti-HA (1:1,000). Veriblot (Abcam) was used for secondary antibody with 1:200-1:500 concentration.

Immunocytochemistry

Fixed cell immunocytochemistry

Collected epididymal sperm and testicular germ cells were immunostained as previously with minor modification (Hwang et al., 2021b; Hwang et al., 2022). Briefly, epididymal sperm and dissociated testicular cells were washed with PBS and centrifuged at 700 x g for 5 minutes or at 250 x g for 3 minutes, respectively, to attach on the glass coverslips. For testicular cells, glass coverslips were coated with poly-D-lysine before centrifugation. Attached epididymal sperm and testicular cells were fixed with 4% PFA for 10 minutes at RT and washed three times with PBS. Fixed cells were permeablized by 0.1% Triton X-100 in PBS for 10 minutes followed by blocked with 10% normal goat serum in PBS for 1 hour at RT. Blocked samples on coverslips were incubated with primary antibodies at 4 °C overnight. Primary antibodies used for the immunocytochemistry were: rabbit polyclonal anti-CATSPER1 (10 µg/ml), rabbit monoclonal anti-HA (1:200), mouse monoclonal anti-pTyr (1:100). The samples were washed with 0.1% Triton X-100 in PBS one time and PBS two times followed by incubation with goat anti-mouse or rabbit IgG conjugated with Alexa 568 (1:1,000) in blocking solution for 1 hour at RT. Testicular cells were stained with WGA conjugated with Alexa 647 (0.1 µg/ml) together with secondary antibody. Immunostained coverslips were mounted with Vectashield (Vector Laboratory) and imaged using Plan-Apochrombat 63X/1.40 oil objective lens equipped in Zeiss LSM710 Elyra P1(Invitrogen). Hoechst was used for counter staining.

Live cell Immunocytochemistry

Prepared sperm were incubated in M2 medium supplemented with 20 μ g/mL of the anti-mouse CATSPER ϵ at 37°C for 90 minutes. Sperm incubated with the primary antibody were washed with PBS followed by incubation with goat anti-rabbit IgG conjugated with Alexa 568 (1:1000) and Hoechst (1:500) in HS medium at 37 °C for 30 minutes. Sperm were washed with PBS and placed on the glass coverslip coated with poly D-lysine for 15 minutes followed by centrifugation at 700 x g for 5 minutes. Sperm were fixed with 4% PFA for 10 minutes at RT and washed with PBS three times. Prepared coverslips were mounted with ProLong Gold (Invitrogen) and imaged by a Plan-Apochrombat 63X/1.40 oil objective lens equipped in Zeiss LSM710 Elyra P1 (Invitrogen).

Immunohistochemistry

Immunohistochemistry was conducted by a standard indirect peroxidase method as previously (**Nielsen et al., 2019**). Briefly, 4 μ m tissue sections were subjected to heat-induced antigen retrieval using medical decloaking chamber (Biocare, Concord, CA, USA) in 0.01M citrate buffer (pH 7.4) at 110 °C for 30 min. Endogenous peroxidase was blocked with 1% H₂O₂ in methanol for 30 min and the sections were blocked using 0.5% skimmed milk in Tris-buffered saline (TBS) for 30 min at RT. Sections were incubated with the anti-human CATSPER ϵ antibody (**Chung et al., 2017**) diluted 1:75 in TBS overnight at 4 °C and for 1 hour at RT in a humidified chamber. The slides incubated with the primary antibody were washed with TBS followed by incubation with anti-rabbit ImmPRESS HRP (peroxidase) antibody (Vector laboratories, CA, USA) at RT for 30

min. After incubation with secondary antibody, slides were washed with TBS and visualized using ImmPACT DAB peroxidase (HRP) substrate (Vector Laboratories). The sections were subsequently counterstained with Mayer's hematoxylin and mounted with Aquatex® mounting medium (Merck KGaA, Germany).

Recombinant CATSPERε^{lg-like} proteins.

Ig-like domain of the CATSPERε (CATSPERε^{Ig-like}) tagged with signaling peptide sequences of mouse IgG heavy chain at N-terminus and mCherry and 6xHis at C-terminus was obtained by CHO express service (GenScript). Briefly, the recombinant protein was expressed in CHO cells and purified using HisTrap[™] FF Crude (GE Health). The purified CATSPERε^{Ig-like} was subjected to SDS-PAGE followed by gel staining using Imperial[™] Protein Stain (Thermo Scientific[™]) and immunoblotting with anti-mCherry to examine its purity and specificity.

Binding test of the Recombinant CATSPERs^{Ig-like} to Epididymal Sperm

Immunoblotting after sperm incubation with CATSPERE^{lg-like}

Collected epididymal sperm were incubated with the purified CATSPER $\epsilon^{lg-like}$ and subjected to immunoblotting and live cell immunocytochemistry to test its binding to the epididymal sperm. For immunoblotting, sperm were capacitated in HTF medium supplemented with or without 20 µg/ml of the recombinant CATSPER $\epsilon^{lg-like}$ for 90 minutes. The capacitated sperm were washed three times with PBS to remove unbound proteins and subjected to immunoblotting as described above. Bound recombinant protein was detected by anti-mCherry (1:1,000).

Live cell immunocytochemistry

Collected epididymal sperm were incubated with the purified CATSPER*e*^{Ig-like} in capacitating conditions for 90 minutes. The sperm were washed one time with PBS followed by an hour incubation with anti-mCherry (1:1000) supplemented in mixture of 0.4% BSA-containing HEPESbuffered HTF (H-HTF; Chung et al., 2017) and immunoenhancer (Pierce) in 1:1 ratio at 37 °C. The sperm were washed two times with PBS and incubated with goat anti-mouse IgG conjugated with Alexa 568 (1:1,000) and Hoechst (1:500) in H-HTF for 30 minutes. After the incubation, sperm were washed with PBS two times and prepared on the coverslips and fixed with 4% PFA in PBS followed by washing with PBS three times for the imaging as described above.

Structural Illumination Microscopy

Mouse epididymal sperm on coverslips were immunostained as described above and subjected to 3D structural illumination microscopy (SIM) imaging. 3D SIM imaging was performed with Zeiss LSM710 Elyra P1 using alpha Plan-APO 63X/1.40 oil objective lens (Carl Zeiss). Z-stack images was acquired with 100 nm intervals and each section was taken using images were taken using a laser at 561 nm wavelength and 5 grid rotations with a 51 nm SIM grating period. Raw images

were processed and rendered using Zen 2012 SP2 software (Carl Zeiss).

Histology Analyses

Histological analyses of the mouse tissues were performed as previously (Hwang et al., 2021b). Briefly, collected testes and epididymis were rinsed with PBS and fixed with 4% PFA in PBS at 4 °C overnight. The fixed tissues were washed with PBS and serially dehydrated by incubation in ethanol to 100%. The tissues were embedded into paraffin followed by sectioned, and the tissue sections were deparaffinized to stain with hematoxylin and eosin (H/E). The stained sections were observed using Nikon E200 microscope under 10x phase contrast objective (CFI Plan Achro 10X/0.25 pH1 WF, Nikon) and imaged by the equipped ac1300–200 μ m CMOS camera (Basler AG).

Mating Test

WT female mice were caged with $Catspere^{+/-}$, $Catspere^{+/-}$; TG+, or $Catspere^{-/-}$; TG+ males and monitored for two months to record pregnancy and litter size.

Motility Analysis

Flagella Waveform Analyses

Flagellar waveform analyses were carried out as described before (Hwang et al., 2021a; Wang et al., 2022). Briefly, non-capacitated or capacitated epididymal sperm ($2-3 \times 10^5$ cells) were transferred to the 37°C HS and H-HTF medium, in fibronectin-coated Delta T chamber (Bioptechs). Planar flagellar movements of the head-tethered sperm were recorded for 2 seconds with 200 frame per seconds (fps) using pco.edge sCMOS camera equipped in Axio observer Z1 microscope (Carl Zeiss). Recorded images stacks were subjected to FIJI software (Schindelin et al., 2012) to measure beating frequency and α -angle of sperm tail (Chung et al., 2011), and to generate overlaid images to trace waveform of sperm flagella as previously described (Chung et al., 2017).

Computer Assisted Semen Analysis

Computer-assisted semen analysis (CASA) was carried out as previous studies (Wang et al., 2022; Hwang et al., 2022). Aliquot of non-capacitated and capacitated sperm $(3.0 \times 10^6 \text{ cells/ml})$, which were incubated with or without recombinant CATSPER $\epsilon^{\text{Ig-like}}$ as above, were placed in slide chamber (CellVision) and motility was examined on a 37°C stage of a Nikon E200 microscope under 10X phase contrast objective (CFI Plan Achro 10X/0.25 Ph1 BM, Nikon) equipped in Nikon E200 microscope. Images were recorded at 50 fps using CMOS video camera (Basler acA1300-200um, Basler AG, Ahrensburg, Germany) and analyzed by sperm Class Analyzer (v6.3; Microptic, Barcelona, Spain). Over 200 motile sperm were analyzed for each trial, at least 3 biological replicates were performed for each experimental group.

In Vitro Fertilization (IVF)

4-8 weeks old female CD1 mice were injected with 10 IU of pregnant mare serum gonadotropin (PMSG, ILEX Life Science) followed by 10 IU of human chorionic gonadotropin (hCG, EMD Milipore) injection after 48 hours. 16 hours after the hCG injection, female mice were sacrificed to collect cumulus-oocyte complexes (COCs) that were placed in fresh TYH medium (**Toyoda and Chang, 1974**) covered with filtered embryo-grade mineral oil (Sigma Aldrich). Epididymal sperm were collected from WT males by swim out into TYH medium followed by capacitation in TYH medium with or without 20 µg/ml of purified CATSPERe^{Ig-like} protein for 90 minutes at 37 °C, 5% CO₂ condition. Collected COCs were incubated for 1 hour at 37°C, 5% CO₂ condition and supplemented with 4 x 10⁵ cells of capacitated sperm. They were incubated together for 6 hours at 37°C, 5% CO₂ condition. After the incubation, oocytes were washed with TYH three times and transferred to fresh TYH medium. Cleaved 2-cell embryos after one day incubation were considered as fertilized eggs.

Electrophysiology

Whole sperm patch clamping was performed as described before (**Hwang et al., 2022**). Sperm were collected from corpus epididymis and washed twice. Washed sperm were resuspended in HS medium followed by adhesion on a culture dish (Corning). Gigaohm seals were formed at the cytoplasmic droplet of motile sperm (Kirichok et al., 2006). Cells were broken through by voltage stimulation (400-600 mV, 5 ms) and suction. Whole-cell CatSper currents were recorded in the divalent-free bath solution composed of (in mM): 150 Na gluconate, 20 HEPES, and 5 Na₃HEDTA with pH 7.4 or HS medium, pH 7.4. Intrapipette solution for recording consisted of 135 CsMeSO₃, 5 CsCl, 10 EGTA, and 10 HEPES adjusted to pH 7.2. Data were sampled at 10 Hz and filtered at 1 kHz. The obtained current data were analyzed with Clampfit (v10.4, Axon, Gilze, Netherlands) and plotted with Grapher (v8.8, Golden Software, Inc., Gloden, Colorado).

Chelation-Based Spectrofluorometric Sperm Em assay

Prepared WT and *Catsper1*-null sperm were capacitated in HTF with or without 20 µg/ml of CATSPER $\epsilon^{lg-like}$ at 2.0 x 10⁶ cells/ml concentration for 90 minutes. Incubated sperm were washed with PBS one time and resuspended into prewarmed H-HTF. 2.0 x 10⁵ cells in 100 µL of H-HTF were loaded with 10 µL of 10 µM DISC3(5) dye suspended in DMSO (American Bio, AB00435-00500). 1 x 10⁵ cells were loaded in each well of a 96-well plate and incubated for 2 minutes at 37°C. After 6 seconds of orbital shaking, fluorescence was measured for 3 minutes using Tecan Infinite 200 Pro operated by i-control software. 3.5 mM of EGTA was used to chelate free Ca²⁺ in H-HTF medium, and fluorescence levels were measured for 2 additional minutes after adding EGTA. All relative fluorescence units (RFU) were background subtracted. RFU changes (Δ RFU) were calculated by subtracting mean fluorescence intensities of EGTA-treated sperm to those of the same sperm prior to EGTA addition. Each value is taken 1 minute before and after adding

EGTA respectively. Δ RFU in WT sperm capacitated without recombinant CATSPER $\epsilon^{lg-like}$ were set to 1 for normalization.

Tissue and Testicular Expression analyses

Transcript data of the CatSper subunits in adult mouse and human tissues were obtained from Mouse ENCODE transcriptome data (**Yue et al., 2014**) and HPA RNA-seq normal tissues data set (**Fagerberg et al., 2014**), respectively, at NCBI (<u>https://www.ncbi.nlm.nih.gov/gene/</u>). Enriched tissue expression was calculated by normalizing transcript level in each tissue with sum of the transcript levels in entire tissues. Enriched tissue expression levels of the CatSper subunits are shown as heatmap using GraphPad Prism 8.0 (GraphPad Software Inc., San Diego, CA).

Structural modeling and data analyses

A structure for mouse CatSper complex was from RCSC Protein Data Bank, PDB (<u>https://www.rcsb.org/</u>; 7EEB). Protein structures were rendered and visualized by iCn3D at NCBI (<u>https://www.ncbi.nlm.nih.gov/Structure</u>; **Wang et al., 2020**). Dimer and trimer structures for ECD domains of the CATSPER auxiliary subunits were predicted using AlphaFold-Multimer under Google Colab notebook (**Evans et al., 2021**). UCSF ChimeraX (**Meng et al., 2023**) was used to visualize atomic model of the CatSper channel (PDB: 7EEB) and to fit the predicted structures to the reference structure using matchmaker tool incorporated in ChimeraX.

Statistical Analyses

Statistical analyses were performed by the Student's t-test and one-way ANOVA with Dunnett's test. Differences were considered significant at *p<0.05, **p<0.01, and ***p<0.001.

Acknowledgement

We thank Masahito Ikawa at Osaka University for sharing *pClgn-mCherry* vector and Katerina Politi at department of pathology, Yale School of Medicine for the use of Tecan plate reader (Infinite 200 Pro). We also appreciate Caro Wiesehoefer for helping initial optimization of using the fluorescence plate reader, Mira Raju and Nathan Lang for helping initial characterization of waveform analysis of WT;*TG*+ sperm, and Sofia Boeg Winge at department of growth and reproduction, Rigshospitalet, Copenhagen University Hospital for supporting human histology sample preparation. This study was supported by MCI Sokal award, NIH R01HD096745, Blavatnik Innovation Fund to J.J.C. J.Y.H. was a recipient of MCI postdoctoral fellowship.

Competing Interest

The authors declare no competing interests.

Author contributions

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Figure Legends



Figure 1. The transmembrane CATSPER ϵ subunit is indispensable for sperm hyperactivation and male fertility. (A-B) Molecular position of CATSPER ϵ within the atomic model and cryo-ET map of the mouse CatSper channel complex. Shown are top-down views of CATSPER ϵ (blue) in the single holo CatSper complex (A, PDB: 7EEB), which was fitted into the cryo-ET map of CatSper (B, EMDB: 24210). The epitope used to raise α -CATSPER ϵ (α -m ϵ -331) that recognize an ECD region (Nand et al., 2023; related manuscript by Clouser et al., 2024) is colored in green and the lg-like domain is colored in navy. (C) *Catspere* mRNA expression in mouse tissues. *Catspere* (blue bars) is specifically expressed in adult testis. (D) Immunostained CATSPER ϵ in human testis. Human CATSPER ϵ antibody (α -h ϵ -31 (Chung et al., 2017) was used. Shown in magnified inset (*bottom*) are testicular sperm cells. (E) Detection of CATSPER ϵ in live epididymal sperm from heterozygous (+/-; *top*) and knockout (-/-; *bottom*) mice. Shown are

merged images of fluorescence and corresponding DIC field images. DNA is stained with Hoechst. (F) Immunoblotting of CATSPER ϵ in epididymal sperm from *Catspere*^{+/-} and *Catspere*^{-/-} (-/-) mice. Acetylated tubulin (AcTUB) is a loading control. (G) Litter sizes from fertile females mated with *Catspere*^{+/-} or *Catspere*^{-/-} males. Individual litter sizes are marked. (H) Flagellar waveform of the epididymal sperm from *Catspere*^{+/-} and *Catspere*^{-/-} mice before (*left*) and after (*right*) inducing capacitation *in vitro*. Flagellar movement of the head-tethered sperm were recorded at 200 fps speed. The tail movements for two beating cycles are overlaid to show the waveform color-coded in time. (I) Maximum angles of the primary flagellar curvature (α -angle) of the epididymal sperm. α -angles of *Catspere*^{+/-} (+/-, gray bars) and *Catspere*^{-/-} (-/-, blue bars) sperm were measured before (0 min; +/-, 117.0° ± 2.9°; -/-, 107.6° ± 6.7°) and after (90 min; +/-, 126.2° ± 4.6°; -/-, 59.6° ± 12.8°) inducing capacitation (cap). ns, non-significant; ***p<0.001. Data are represented as mean ± SEM (G and I). *See also* Figures S1 and S2.



Figure 2. Catspere-null spermatozoa lack functional CatSper channel. (A) Immunoblotting of CatSper subunits in WT (+/+) and Catspere-null (-/-) epididymal sperm. Acetylated tubulin (AcTUB) is a loading control. (B) Confocal images of immunostained CATSPER1 in Catspere^{+/-} (top) and Catspere-/- (middle and bottom) epididymal sperm. Fluorescence and corresponding DIC images are shown for *CatSpere^{-/-}* sperm (*bottom*). Hoechst is used for counter staining DNA. (C) Immunoblotting of protein tyrosine phosphorylation (pY) in WT (+/+) and Catspere-null (-/-) sperm. Time-course development of pY was examined for 90 min incubation of cauda epididymal sperm under capacitation (Cap) conditions. (D and E) Representative traces of CATSPER current (*I_{CatSper}*) of corpus epididymal sperm from *Catspere^{+/-}* and *Catspere^{-/-}* males. *I_{CatSper}* traces were elicited by voltage-ramp protocol from -100 mV to +100 mV range with 0 mV holding potential (D) or by step protocol from -120 mV to +140 mV ranges with 20 mV increments (E). A cartoon in panel D represents pH and ion composition (mM) in pipette solution and used bath solution for I_{CatSper} measurement by voltage-ramp and step protocols. Divalent free (DVF) solution or HEPESbuffered saline (HS) were used for bath solutions. (F) Inward I_{CATSPER} measured from Catspere^{+/-} (gray, n=5) and Catspere^{-/-} (blue, n=3) corpus sperm at -100 mV. Circles indicate the current from individual spermatozoa. Data are shown to mean ± SEM. See also Figure S2.



Figure 3. Truncation of the extracellular domain fails CATSPERε to localize at the sperm flagella. (A) A schematic diagram for *Catspere* transgene (*Catspere*^{TG}) encoding extracellular domain (ECD) truncated CATSPERε in mice. The transgene encodes the CATSPERδhomologous region of CATSPERε fused to its native signal peptide and HA at N-terminus and mCherry at C-terminus (~61 kDa, *bottom*), respectively. *Clgn* promoter is for transgene expression specific to testicular germ cells (*top*). (B) Immunoblotting of the truncated CATSPERε in testis and epididymal sperm from *Catspere*^{TG+} (*TG*+) mice. Asterisks (*) indicate truncated CATSPERε proteins. Acetylated tubulin (AcTUB) is used as a loading control. (C) Immunoblotting of the truncated CATSPERε in cytosol and microsome fraction of testis from WT; *TG*+ males. Na⁺/K⁺ ATPase and Calmodulin (CaM) serve as controls for microsome and cytosol fraction, respectively, and for loading. (D) Confocal images of immunostained CATSPER1 and HA in epididymal sperm from WT (*top*) and WT with the transgene (WT; *TG*+, *bottom*) male mice. Merged are fluorescence images of sperm immunostained with α-HA and corresponding DIC images (*right*). (E) 3D structural illumination microscopy images of immunostained CATSPER1 in sperm from WT (*top*) and WT; *TG*+ (*bottom*) mice. Colors in *x-y* projection indicate *z*-depth

information (*left*). *y-z* projection represents cross-section images (*right*). DNA was counter-stained with Hoechst. (F) Co-immunoprecipitation of native CATSPER subunits and truncated CATSPER ϵ in WT, *Catsperd*-null (*d-/-*), and WT; *Catspere*^{TG} (*TG*+) testis. Native CATSPER subunits – β , δ , ϵ , and 1 – and truncated CATSPER ϵ are examined in HA and CATSPER δ immunocomplexes reciprocally. (G) Confocal images of truncated CATSPER ϵ immunostained with α -HA in developing spermatids from WT (*left*) and WT; *TG*+ (*right*) males. Shown are fluorescence (*top*) and corresponding DIC images merged with fluorescence for DNA (*bottom*). Wheat germ agglutinin (WGA) and Hoechst were used to stain sugar residues and DNA, respectively. The truncated CATSPER ϵ is probed by α -HA (B, C, D, F, and G). *See also* Figure S3.



Figure 4. Protein-protein interaction of CATSPERε canopy with neighboring ancillary subunits is essential to assemble the entire CatSper channel complex. (A) Immunoblotting of truncated CATSPERε in testis and sperm from WT (+/+) and *Catspere*-null (-/-) males with the transgene (*TG*). Acetylated tubulin (AcTUB) is a loading control. Asterisks (*) indicate the truncated CATSPERε expressed from the transgene. (B) Co-immunoprecipitation of the truncated CATSPERε and CATSPER subunits in testis from WT and *Catspere*-null (e-/-) mice carrying the transgene (*TG*). α-HA was used to detect the truncated CATSPERε (A and B). *Catsperd*-null testis (*d*-/-) is used as negative control. (C) Organization of CATSPERε, γ, and δ extracellular domains (ECDs). Domain boundaries of each subunit are labeled. A gray sash in CATSPERε indicates ECD region remaining in the truncated CATSPERε. (D and E) Predicted interaction of the truncated CATSPERε and CATSPERε and CATSPERδ (D) but not with CATSPERγ (E). Shown are AlphaFold

Multimer-predicted interaction of the whole (*left*) or truncated (*right*) CATSPER ε ECD with whole ECDs of CATSPER δ (D) or CATSPER γ (E). Modeled structures for interacting two ECDs are fitted to the reference canopy structure of the CATSPER channel colored in gray (PDB: 7EEB). Black lines marked in top view structures (*top*) indicate the orientation for the corresponding side views (*bottom*).



Figure 5. ECD-truncated CATSPER ϵ fails to rescue impaired sperm hyperactivation and male infertility in *Catspere*-null mice. (A) Confocal images of immunostained CATSPER1 and HA in epididymal sperm from WT, *Catspere*^{+/-}, and *Catspere*^{-/-} mice crossed with the transgenic (*TG*) mice. Shown are merged views of fluorescence and corresponding DIC images. Hoechst is used to counterstain DNA. (B) Immunoblotting of CATSPER subunits in epididymal sperm from WT and *Catspere*-null (*Catspere*-/-) males with the transgene (*TG*). Acetylated tubulin (AcTUB) is used as loading control. (C and D) CatSper current (*I*_{CATSPER}) from sperm from *Catspere*-null mice expressing the transgene (*Catspere*^{-/-}; *TG*+) and WT sperm. Shown are representative *I*_{CATSPER} traces elicited by ramp (C, -100 mV to +100 mV) and step (D, -120 mV to +140 mV, 20 mM increment) protocols. Holding potential for voltage-ramp protocol and step protocol was 0 mV

and -120 mV, respectively. A cartoon in panel C represents ion composition (mM) and pH in bath and pipette solution. (E) Comparison of the inward $I_{CATSPER}$ of corpus WT sperm (gray bar, N=3) and *Catspere*-null sperm with the transgene (*Catspere*^{-/-}; *TG*+, blue bar, N=3) recorded at -100 mV. Circles indicate $I_{CATSPER}$ measured from individual sperm. Data are represented to mean ± SEM. (F) Pregnancy rate of WT females mated with *Catspere*^{+/-} and *Catspere*^{-/-} males carrying the transgene (*Cse*^{+/-}; *TG*+ and *Cse*^{-/-}; *TG*+, respectively). (G) Flagellar waveform of sperm from *Catspere*^{+/-} and *Catspere*^{-/-} mice expressing the transgene (*TG*+). Tail movements of headtethered spermatozoa were recorded at 200 fps before (non-capacitated, 0 min; *left*) and after (capacitated, 90 min; *right*) incubation under capacitating conditions. Images for two beating cycles are overlaid and color-coded in time. *See also* Figure S4.



Figure 6. Administration of recombinant CATSPER ϵ Ig-like domain protein attenuates CATSPER function and sperm fertility *in vitro*. (A) Location of CATSPER ϵ Ig-like domain in the canopy structure. *Top*, shown left is the atomic model of CATSPER ϵ , γ , δ , and β that form the canopy (from PDB: 7EEB). Shown right is top view of the canopy structure. Ig-like domain of CATSPER ϵ (CATSPER ϵ ^{Ig-like}) is highlighted in light purple. *Bottom,* a diagram of recombinant CATSPER ϵ ^{Ig-like} protein fused with mCherry and 6xHis tag. The signal peptide of mouse IgG heavy chain (gray) was added for secretion of the recombinant protein expressed in CHO cells. (B) Purified recombinant CATSPER ϵ Ig-like domain. Shown are Coomassie blue stained (*left*) and

immunoblotted (*right*) recombinant CATSPER $\varepsilon^{lg-like}$ for purity and specificity validation. CATSPERelg-like was probed by anti-mCherry antibody. (C and D) CATSPER-specific CATSPERE^{lg-like} binding to epididymal sperm. Sperm cells were incubated with recombinant CATSPER $\epsilon^{lg-like}$ ($\epsilon^{lg-like}$) proteins under capacitating conditions and the protein bound to the sperm surface are detected by immunoblotting (C) or immunostaining (D) by anti-mCherry after washing unbound proteins. Detection of recombinant CATSPERE^{Ig-like} protein added to Catsper+/- (left) and Catsper^{-/-} (right) sperm. Immunofluorescence images and corresponding DIC images are shown (D). CATSPERelg-like was detected by anti-mCherry. (E) CASA analysis of sperm motility. Total motility, hyperactivated motility, curvilinear velocity (VCL), and amplitude of lateral head (ALH) were measured from sperm after incubating under capacitating conditions with or without CATSPERE^{lg-like}. **p<0.01 and ***p<0.001. Data are represented as mean ± SD. (F and G) CATSPER activity assay by chelation-based Em changes. Live WT (left) and Catsper1-null (right) sperm capacitated with (gray) or without (blue) CATSPER $\epsilon^{Ig-like}$ are loaded with ~1 uM DiSC₃(5) dye for Em imaging (F). Relative fluorescent unit changes (ΔRFU) after EGTA treatment are normalized and the change in WT sperm not incubated with CATSPER^{g-like} was set to 1 (G). **p<0.01. N=3. Data are represented as mean ± SEM. (H) 2-cell cleavage rate of in vitro fertilized eggs with WT sperm capacitated with (gray) or without (blue) recombinant CATSPERE^{Ig-like}. ***p<0.001. N=3. Data are represented as mean ± SD. CSε^{lg-like}, CATSPERε^{lg-like} (E, F, G, and H).