Conservation of the Ca\textsuperscript{2+}-permeability through the voltage sensor domain of mammalian CatSper subunit

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ABSTRACT

Cation channel of Spermatozoa (CatSper) is one of the voltage-gated ion channels consisting of voltage sensor domains (VSDs) and pore-gate domains. CatSper is exclusively expressed in spermatozoa and indispensable for Ca\textsuperscript{2+} influx into cytosol. Recently, we have reported that the VSD of ascidian CatSper induces Ca\textsuperscript{2+}-permeable pathways in heterologous expression systems. However, it is not known whether ion permeability through the VSD of CatSper is conserved in mammals. In the present study, electrophysiology and fluorometry in Xenopus oocytes revealed that Ca\textsuperscript{2+}-permeable paths are also formed by expressing the VSD of murine CatSper. We also examined the permeability to monovalent cations other than Na\textsuperscript{+} in the VSD of ascidian CatSper.

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Introduction

Voltage-gated ion channels mediate the passive ion transportation in response to membrane potential changes. In voltage-gated K\textsuperscript{+} channels, the pore-forming subunits consist of six trans-membrane regions which are divided into two functional domains: the voltage sensor domain (VSD; S1 ~ S4) and the pore-gate domain (S5 ~ S6) \cite{1}. The VSD has been considered to exclusively function as the regulator of the pore of voltage-gated ion channels, but multiple studies have reported that direct ion permeation pathways exist in some mutated VSDs and in the voltage-gated proton channel \cite{2–7}. Thus, the VSD of voltage-gated ion channels plays more roles in physiological and pathological situations than as considered in the past several decades.

Cation channel of Spermatozoa (CatSper) belongs to the voltage-gated ion channel superfamily \cite{8}. It has been suggested that the four distinct \( \alpha \) subunits of CatSper (CatSper1, CatSper2, CatSper3 and CatSper4) and several accessory subunits form a CatSper complex \cite{8–13}. Each \( \alpha \) subunit consists of six trans-membrane regions with homologous domains to the VSD and the pore-gate domain of voltage-gated ion channels. Gene knockout of the CatSper \( \alpha \) subunits causes male infertility due to the lack of Ca\textsuperscript{2+} supply into spermatozoa which is essential for hyperactivation of the sperm motility in mice \cite{12,14–17}. Electrophysiological analyses by direct whole-cell recordings from mammalian spermatozoa revealed that the CatSper currents are potentiated by intracellular alkalinization, and monovalent currents are elicited in divalent-free condition \cite{18,19}.

Recently, we have reported that the VSD of the ascidian CatSper3 subunit (CiCS3 VSD) forms Ca\textsuperscript{2+} permeation pathways in the plasma membrane of Xenopus oocytes and cultured mammalian cells \cite{20}. That was the first case that showed evidence for divalent cation permeation through a VSD as well as Ca\textsuperscript{2+} current of CatSper in heterologous expression systems. However, several important issues remain to be addressed. First, mechanisms of ion permeability through the VSD of ascidian CatSper3 are elusive. Detailed information of permeability to monovalent cations is essential for understanding mechanisms of ion permeation through CatSper VSD. Second, it remains unclear whether the Ca\textsuperscript{2+} permeability of VSD is unique to the ascidian CatSper3. In the present study, we studied the permeability of the VSD of the ascidian CatSper3...
to monovalent cations. We also tested if the VSD of CatSper derived from mouse has Ca\(^{2+}\) permeability by heterologous expression in Xenopus oocytes. We found that the VSD of ascidian CatSper3 is permeable to various monovalent cations, such as K\(^{+}\), Cs\(^{+}\) and Li\(^{+}\). On the other hand, we did not obtain any evidence for permeation to protons. Overexpression of the VSD of murine CatSper3 (mCS3 VSD) gave rise to Ca\(^{2+}\) permeation pathways which are activated upon membrane hyperpolarization, suggesting the conservation of Ca\(^{2+}\) permeability between the VSDs of murine and ascidian CatSper subunit.

**Materials and Methods**

**Ethics approval**

Experiments using *Xenopus laevis* and mice were performed in accordance with the guidelines of the Animal Care and Use Committee of Osaka University Medical School.

**cDNA cloning of the voltage sensor domain of murine CatSper3**

Total RNA from a testis of C57BL/6 mouse was extracted with TRIzol Reagent (Thermo Fisher Scientific, Waltham, MA). The cDNA fragment of murine CatSper3 (mCS3; NM_001252187.1) was obtained by RT-PCR and subcloned into a custom modified version of the pcS2+ vector, and sequenced. The primers used for cloning the mCS3 are 5ʹ-cggatccatgcttcatcaccatcaccacaacc-3ʹ and 5ʹ-cggatctctactgttagggctggtc-3ʹ. The cDNA of the voltage sensor domain of murine CatSper3 (mCS3 VSD; M1-T176) was subcloned into the pcS2+ vector by PCR using the following primers; 5ʹ-cggatctccatcaccatcaccacaccacc-3ʹ and 5ʹ-cggatctctactgttagggctggtc-3ʹ.

**Construction of a chimeric protein and mutagenesis**

Constructions of chimeric proteins were done by PCR. A cDNA fragment of iR-pHluorin [21] was kindly gifted by Dr. Hiroyuki Katayama and Dr. Atsushi Miyawaki (RIKEN, Japan). The fusion construct of the VSD of ascidian CatSper3 [20] and iR-pHluorin (CiCS3 VSD-iR-pHluorin) was created with the primers: 5ʹ- ccaagctatggagaaagactccg -3ʹ, 5ʹ- cccgatccatggtgttagggttgagtggtagg -3ʹ, and 5ʹ- cccgatccatggtgttagggttgagtggtagg -3ʹ.

**Two-electrode voltage clamp recordings and fluorometry in oocytes**

Two-electrode voltage clamp (TEVC) recordings and fluorometry were done as described previously [20]. Briefly, *Xenopus* oocytes were defolliculated by treating with type I collagenase (1.0 mg/mL; Sigma-Aldrich, St. Louis, MO) in ND96 solution; 96 mM NaCl, 2 mM KCl, 1.8 mM CaCl\(_2\), 1 mM MgCl\(_2\), 5 mM 4-(2-Hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), 0.1 mg/mL Gentamycin, 5 mM Pyruvate-Na, pH = 7.5 adjusted by NaOH [24]. The defolliculated oocytes were then injected with 50 nL of cRNA solution. The concentration of cRNA synthesized using mMESSAGE mMACHINE transcription kit (Thermo Fisher Scientific) was 1 – 1.5 ng/nL and 0.1 – 0.2 ng/nL for mCS3 VSD and CiCS3 VSD, respectively.

Current recordings were conducted at 2–3 days after cRNA injection by TEVC using an amplifier (OC-725; Warner Instruments, Hamden, CT). Acquired data were digitized by an AD/DA converter (InstruTECH LIH 8 + 8; HEKA Elektronik, Lambrecht/Pfalz, Germany), and analyzed with PatchMaster software (HEKA Elektronik). The
interval between traces was 10 seconds. Current-Voltage (I-V) relationships were obtained by plotting the average current amplitudes at the end of the test pulses against the membrane potential. Glass electrodes had resistance of 0.3–0.5 M ohm after filling with 3 M KCl solution. Perfusion of the bath solution was done using a gravity perfusion system. The composition of the bath solutions are described below.

32 K\(^+\) solution: 32 mM KOH, 5 mM MgCl\(_2\), 140 mM sucrose, 5 mM HEPES, pH 7.5 (pH adjusted by methanesulfonic acid: MeSO\(_3\)H)
96 K\(^+\) solution: 96 mM KOH, 5 mM MgCl\(_2\), 40 mM Sucrose, 5 mM HEPES, pH 7.5
32 Li\(^+\) solution: 32 mM LiOH, 5 mM MgCl\(_2\), 140 mM Sucrose, 5 mM HEPES, pH 7.5
96 Li\(^+\) solution: 96 mM LiOH, 5 mM MgCl\(_2\), 40 mM Sucrose, 5 mM HEPES, pH 7.5
32 Cs\(^+\) solution: 32 mM CsOH, 5 mM MgCl\(_2\), 140 mM Sucrose, 5 mM HEPES, pH 7.5
96 Cs\(^+\) solution: 96 mM CsOH, 5 mM MgCl\(_2\), 40 mM Sucrose, 5 mM HEPES, pH 7.5
32 NMDG solution: 32 mM NMDG, 5 mM MgCl\(_2\), 140 mM sucrose, 5 mM HEPES, pH 7.5
96 NMDG solution: 96 mM NMDG, 5 mM MgCl\(_2\), 40 mM sucrose, 5 mM HEPES, pH 7.5
pH 6 solution: 96 mM NMDG, 5 mM MgCl\(_2\), 40 mM sucrose, 5 mM 2-Morpholinoethanesulfonic acid, pH = 6.0 (MeSO\(_3\)H)

For the fluorometry, macroscopic currents and fluorescence changes were simultaneously recorded using an amplifier (OC-725C) and an inverted microscope (IX70; Olympus, Tokyo, Japan) equipped with a stable 75W xenon lamp (Ushio, Tokyo, Japan). Oocytes were injected with 50 nL of 1 mM Fluo-3 (DOJINDO MOLECULAR TECHNOLOGIES, INC, Kumamoto, Japan) 1–3 hrs before recordings for visualizing changes of intracellular Ca\(^{2+}\) concentration. The GFP-A-BASIC-OMF was used for excitation of Fluo-3 and iR-pHluorin, and an objective lens (× 20, NA 0.70; Olympus) was used to collect fluorescence. A photomultiplier (PMT) module (H5784-02; Hamamatsu Photonics, Shizuoka, Japan) was used for the acquisition of fluorescence. The output signals from the PMT module and the TEVC amplifier were digitized and stored using an A/D converter (1322A; Molecular Devices, Sunnyvale, CA) and pCLAMP8 software (Molecular Devices), respectively. Data were analyzed by Clampfit 10.3 software (Molecular Devices), and the fluorescence signal was digitally filtered at 500 Hz. The interval between traces was 30 seconds. Fluorescence changes (ΔF) were calculated by normalizing with the average signal intensity of 300 ms before the test pulses. Fluorescence-Voltage (F-V) relationships were obtained by plotting average ΔF for 50 ms from the end of the test pulses against membrane potential. The composition of Ca\(^{2+}\) free solution is 96 mM NaCl, 2 mM KCl, 5 mM MgCl\(_2\), 5 mM HEPES, pH 7.5 (NaOH).

Results

The VSD of ascidian CatSper3 is permeable to K\(^+\), Li\(^+\) and Cs\(^+\)

In our previous study, we found that the VSD of ascidian CatSper3 (CiCS3 VSD) is permeable to Na\(^+\) as well as Ca\(^{2+}\) [2, 20]. However, permeability to other monovalent cations has not been examined. We tried to compare the reversal potentials under different bi-ionic conditions but it was unsuccessful since the kinetics of tail currents was too rapid. Permeability to K\(^+\), Li\(^+\) and Cs\(^+\) was examined by comparing the current amplitude in two different concentrations of extracellular cations, 32 mM and 96 mM, in the same cell. All solutions contained Mg\(^{2+}\) because oocytes are unstable without divalent cation (See Methods). Currents recorded in 96 mM solution were larger than those in 32 mM solution in all cases, indicating that the CiCS3 VSD is permeable to K\(^+\), Li\(^+\) and Cs\(^+\) (Figure 1 A-C). In recordings with perfusion from 32 mM NMDG solution to 96 mM NMDG solution, no obvious increase of the current amplitude was observed (Figure 1D), suggesting that the CiCS3 VSD is not permeable to NMDG. The charge carrier of the current recorded in NMDG solutions could be either Mg\(^{2+}\) or protons which are contained in the bath solutions.

To know whether CiCS3 VSD is permeable to protons, pH fluorometry was performed using the pH-sensitive green fluorescence protein, iR-pHluorin [21]. iR-pHluorin fused to the C-terminus of CiCS3 VSD (CiCS3 VSD-iR-pHluorin) was expressed in Xenopus oocytes. A fusion protein of iR-pHluorin and the voltage-
Figure 1. Permeability of CiCS3 VSD to monovalent cations (A-D) Current traces and I-V relationships of the recordings (mean ± SEM) for examining K⁺ (A), Li⁺ (B), Cs⁺ (C) and NMDG (D) permeability. The same cells were recorded in the solutions with two different concentrations of cations. Voltage steps were from +50 mV to −150 mV by 10 mV decrement. Holding potential (Vh) was at −10 mV. The current evoked by a step pulse to −150 mV is colored by red. (E) Test of proton permeability. Currents and fluorescence changes of iR-pHluorin were recorded from an oocyte expressing CiCS3 VSD-iR-pHluorin (left) or iR-pHluorin-CiHv1/VSOP (right) in the pH6 solution. Voltage steps for CiCS3 VSD-iR-pHluorin were from −150 mV to −10 mV. Vₜₜ was −10 mV. The voltage steps for iR-pHluorin-CiHv1/VSOP expressing cells were from −20 mV to +100 mV. Vₜₜ was −80 mV.
gated proton channel cloned from Ciona intestinalis, CiHv1/VSOP [3], which was expressed as a positive control (iR-pHluorin-CiHv1/VSOP), showed clear fluorescence change upon depolarization of the membrane (Figure 1E). However, no fluorescence change was observed upon hyperpolarization in cells expressing the CiCS3 VSD-iR-pHluorin, which showed robust inward current (Figure 1E), suggesting that the proton does not permeate through the CiCS3 VSD. This supports the idea that the currents recorded in NMDG solutions are most likely carried by Mg, but not by protons.

Ca\(^{2+}\) permeable paths are formed by the voltage sensor domain of murine CatSper3

Our previous study on the VSD of ascidian CatSper3 prompted us to test if the VSD of the murine CatSper3 (mCS3 VSD: See Figure 2A) also has ion permeability. 25.3% of amino acids in the trans-membrane regions are identical between the mCS3 VSD and the CiCS3 VSD. When the mCS3 VSD was expressed in Xenopus oocytes, the currents evoked by voltage steps were indistinguishable from those in uninjected cells (Figure 2B). To facilitate the expression of the mCS3 VSD in the plasma membrane, we utilized the N-terminal cytoplasmic region of the ascidian voltage-sensing phosphatase (Ci-VSP) which is known to facilitate the protein translocation to the plasma membrane [22,23]. In oocytes expressing the chimeric protein of mCS3 VSD and N-terminus of Ci-VSP (MT-mCS3 VSD), slowly-activating inward currents were elicited upon membrane hyperpolarization (Figure 2B and C). Although the amount of injected cRNA for MT-mCS3 VSD was 5–10 times larger than that for the CiCS3 VSD [20], amplitudes of the currents in cells expressing the MT-mCS3 VSD (−0.40 ± 0.060 nA, n = 10, at −120 mV) were approximately 4 times smaller than those of the CiCS3 VSD (−1.69 ± 0.29 nA, n = 7, at −120 mV). As in our former study on ascidian CatSper3 [20], rebound currents were evoked upon depolarization to −10 mV from hyperpolarizing voltage steps. This suggests that the calcium-activated chloride channels endogenously expressed in Xenopus oocytes were activated following the membrane hyperpolarization, implying that Ca\(^{2+}\) influx or release of Ca\(^{2+}\) from intracellular stores occurred upon membrane hyperpolarization as already reported for the CiCS3 VSD [20]. Given that positively charged amino acids on the S4 segment are known to play key roles in the activation of the currents in most of voltage-gated ion channels, we examined the effect of a mutation in the S4 segment of the MT-mCS3 VSD. Substitution of the first arginine to a glutamine (R151Q) did not affect the inward currents in oocytes. Upon mutation of Lys-154 (K154Q), the current magnitude was remarkably smaller than that of WT (Fig. S1).

To verify that the ion permeation paths induced by expression of the MT-mCS3 VSD are permeable to Ca\(^{2+}\), we performed simultaneous recordings of macroscopic ionic currents and intracellular fluorescence changes of one of the Ca\(^{2+}\) indicators, Fluo-3. In Xenopus oocytes expressing the MT-mCS3 VSD, inward currents and increases of intracellular fluorescence were observed upon membrane hyperpolarization when the cells were bathed in ND96 solution that contains Ca\(^{2+}\) (Figure 3A left and B). In contrast, such fluorescence changes were not detected from cells bathed in nominally Ca\(^{2+}\) free solution (Figure 3A right and B) though the ionic currents were elicited. Neither ionic current nor fluorescence change was elicited in intact cells bathed in ND96 solution (Figure 3B). These results indicate that Ca\(^{2+}\) influx occurs upon membrane hyperpolarization in oocytes expressing the MT-mCS3 VSD. Notably, the Ca\(^{2+}\) signal was detected from the step pulse to −50 mV that is within the physiological range of the membrane potential.

Discussion

In this study, we addressed the two issues which have remained unclear in our previous study; (1) Permeability of the VSD of ascidian CatSper3 (CiCS3 VSD) to monovalent cations, (2) Conservation of Ca\(^{2+}\) permeability of CatSper3 among species. We showed that the CiCS3 VSD was permeable to various monovalent cations, and the VSD of murine CatSper3 (mCS3 VSD) induced the Ca\(^{2+}\)-permeable pathways in the plasma membrane of Xenopus oocytes. The
mCS3 VSD-induced Ca\(^{2+}\) influx was observed in the physiological range of the membrane potential. Direct current recordings from murine spermatozoa revealed that the CatSper is permeable to monovalent cations, such as Na\(^+\) and Cs\(^+\), as well as Ca\(^{2+}\), [18]. As is consistent with this, we demonstrated that the CiCS3 VSD was permeable to monovalent cations including K\(^+\), Li\(^+\) and Cs\(^+\), indicating that the CiCS3 VSD functions as a non-selective cation channel (Figure 1). However, the CiCS3 VSD was not thought to be permeable to protons as far as examined with a pH-sensitive green fluorescence protein (Figure 1E).

In our recordings, no obvious tail current was observed as in the CiCS3 VSD, and therefore, it was impossible to determine the ion selectivity of the CiCS3 VSD from the reversal potential in different bi-ionic conditions. Similarly, omega currents (or gating pore currents) through the VSDs of Shaker K\(^+\) channel and Nav1.4 channel show no
This suggests that the rapid closing of the ion permeable pathway is a common characteristic among ion-permeable VSDs except Hv1. Further analyses, such as single channel analysis, are required for understanding more detailed biophysical properties of the CiCS3 VSD.

Our previous study revealed that the VSD of ascidian CatSper3 (CiCS3 VSD) forms hyperpolarization-activated Ca-permeable pathways in the plasma membrane of Xenopus oocytes and cultured cells. As is the case with the CiCS3 VSD, inward currents and Ca influx were evoked upon membrane hyperpolarization in the oocytes expressing the mCS3 VSD (Figures 2 and 3). In addition, the inward currents recorded in the nominally Ca free solution indicated that the paths induced by the mCS3 VSD are permeable to other ions, such as Na+ (Figure 3). These common observations suggest that the ion permeability of CatSper3 VSD is an intrinsic trait conserved among species despite the low homology between the primary structure of the mCS3 VSD and the CiCS3 VSD. In voltage-gated Ca channels, negatively charged amino acids in the pore-gate domain are important for the Ca permeation. By analogy to this, negatively charged amino acids in trans-membrane regions may play key roles for the Ca permeability in the CiCS3 VSD and the mCS3 VSD. In fact, although the entire homology is low (only 25.3% amino acid identity), three negatively charged amino acids in the S2 segment are conserved between the ascidian and the murine CatSper3 (Figure 2A).

Figure 3. The ion permeation paths induced by the MT-mCS3 VSD are permeable to Ca^{2+} [2]. (A) Representative current traces and fluorescence changes from oocytes expressing the MT-mCS3 VSD bathed in ND96 (Left) or Ca^{2+} free solution (Right). The voltage steps were from −150 mV to −10 mV by 20 mV increment. Vh was at −10 mV. The traces evoked by a step pulse to −150 mV are colored by red. The recordings in ND96 and Ca^{2+} free solution were from different cells. (B) Fluorescence-Voltage relationships (upper) and Current-Voltage relationships (lower) of the recordings (mean ± SEM).
Facilitation of membrane targeting of the mCS3 VSD by making chimeric protein with the N-terminal cytoplasmic region of Ci-VSP was necessary to detect the ion conductance. The expression level of the mCS3 VSD in the plasma membrane of \textit{Xenopus} oocytes was much lower than that of the CiCS3 VSD which showed robust current without requiring such modification \cite{20}. We also investigated the biophysical properties of VSDs of ascidian CatSper1, CatSper2 and CatSper4, but ionic currents were not recorded reproducibly. That may be because of insufficient expression in \textit{Xenopus} oocytes.

The ionic currents were detectable at more negative voltages than \(-100\) mV in cells expressing the mCS3 VSD (Figure 2), whereas Ca fluorometry indicated that the Ca influx started to be elicited by a step pulse to \(-50\) mV as in ascidian CatSper3 (Figure 3B)\cite{20}. Chávez \textit{et al}. showed that the membrane potential of murine spermatozoa reaches approximately \(-70\) mV during the final maturation of ejaculated spermatozoa called capacitation \cite{27}. Thus, the ion permeation pathways induced by the VSD of mCS3 are capable of conducting Ca in the physiological situation if it is functional in spermatozoa. However, previous studies suggested that the CatSper is activated by depolarization of the membrane \cite{18}. This issue should be addressed by investigating the function of whole CatSper complex in heterologous expression systems. To know whether Ca enters into the cytosol of spermatozoa through the VSD of CatSper\(\alpha\) subunits, it may be worth testing if the infertility of male mice in which spermatozoa lack CatSper complex can be rescued by overexpression of the CatSper VSDs in the sperm membrane.

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\textbf{Disclosure statement}

The authors declare no competing interest.

\textbf{Author contributions}

HA conducted the experiments and analyzed the data. HT and YO designed and supervised the research. HA, HT and YO wrote the paper.

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