Simultaneous knockout of Slo3 and CatSper1 abolishes all alkalization- and voltage-activated current in mouse spermatozoa

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters.

Citation

Published Version
doi:10.1085/jgp.201311011

Accessed
July 14, 2018 8:00:43 AM EDT

Citable Link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:12064351

Terms of Use
This article was downloaded from Harvard University's DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA

(Article begins on next page)
Simultaneous knockout of Slo3 and CatSper1 abolishes all alkalinization- and voltage-activated current in mouse spermatozoa

Xu-Hui Zeng,1 Betsy Navarro,2,3 Xiao-Ming Xia,4 David E. Clapham,2,3,5 and Christopher J. Lingle4

1Institute of Life Science, Nanchang University, Nanchang, Jiangxi 330031, China
2Howard Hughes Medical Institute and 3Department of Cardiology, Boston Children’s Hospital, Boston, MA 02115
4Department of Anesthesiology, Washington University in St. Louis School of Medicine, St. Louis, MO 63110
5Department of Neurobiology, Harvard Medical School, Boston, MA 02115

During passage through the female reproductive tract, mammalian sperm undergo a maturation process termed capacitation that renders sperm competent to produce fertilization. Capacitation involves a sequence of changes in biochemical and electrical properties, the onset of a hyperactivated swimming behavior, and development of the ability to undergo successful fusion and penetration with an egg. In mouse sperm, the development of hyperactivated motility is dependent on cytosolic alkalinization that then results in an increase in cytosolic Ca2+. The elevation of Ca2+ is thought to be primarily driven by the concerted interplay of two alkalinization-activated currents, a K+ current (KSPER) composed of pore-forming subunits encoded by the Kcnu1 gene (also termed Slo3) and a Ca2+ current arising from a family of CATSPER subunits. After deletion of any of four CATSPER subunit genes (CATSPER1–4), the major remaining current in mouse sperm is alkalinization-activated KSPER current. After genetic deletion of the Slo3 gene, KSPER current is abolished, but there remains a small voltage-activated K+ current hypothesized to reflect monovalent flux through CATSPER. Here, we address two questions. First, does the residual outward K+ current present in the Slo3−/− sperm arise from CATSPER? Second, can any additional membrane K+ currents be detected in mouse sperm by patch-clamp methods other than CATSPER and KSPER? Here, using mice bred to lack both Slo3 and CATSPER1 subunits, we show conclusively that the voltage-activated outward current present in Slo3−/− sperm is abolished when CATSPER is also deleted. Any leak currents that may play a role in setting the resting membrane potential in noncapacitated sperm are likely smaller than the pipette leak current and thus cannot be resolved within the limitation of the patch-clamp technique. Together, KSPER and CATSPER appear to be the sole ion channels present in mouse sperm that regulate membrane potential and Ca2+ influx in response to alkalinization.

INTRODUCTION

After ejaculation, mammalian sperm must undergo a maturational process termed capacitation to acquire competence to fertilize an egg (Darszon et al., 2007). Capacitation entails an extensive spectrum of fundamental changes in sperm properties, including biochemical, electrical, and motile (Visconti and Kopf, 1998; Visconti et al., 1998; Darszon et al., 2011). One important and visually obvious step in sperm maturation is the acquisition of a hyperactivated motility, which is thought both to facilitate movement of sperm through the female reproductive tract and, subsequently, to enable successful penetration by sperm of protective layers surrounding an egg (Suarez, 2008). Although the full set of endogenous signals that lead to capacitation and hyperactivation is not entirely understood (Visconti et al., 2002; Fraser et al., 2006), an important step in the process by which sperm acquire hyperactivated motility is cytosolic alkalinization (Ho and Suarez, 2001; Suarez, 2008). Alkalinization gradually arises, in part, from relocation of sperm from the acidic environment of the epididymis to a more alkaline environment of the female reproductive tract (Kirchok and Lishko, 2011), with proton reequilibration, at least in humans, perhaps being mediated by proton channels (Lishko et al., 2010). Importantly, alkalinization is associated with an increase in cytosolic Ca2+ (Wennemuth et al., 2003; Darszon et al., 2005). Together, the rise in pH and [Ca2+], are the cytosolic signals essential for initiation of hyperactivation (Suarez, 2008).

An extensive body of early work has supported the view that activation of a combination of voltage-dependent Ca2+ channels and K+ channels played a role in controlling sperm membrane potential and [Ca2+], (Darszon et al., 1999). However, despite an extensive list of proposed sperm channel candidates (Darszon et al., 1999, 2006), the molecular identity of specific channels that might mediate these effects remained elusive until the last 10 years. With the advent of direct patch-clamp recording from sperm and specific genetic
Kirichok et al., 2006) mediates influx of Ca\(^{2+}\) necessary for activation of CATSPER during alkalization (Ren et al., 2001; Zeng et al., 2011). KSPER and CATSPER are thought to work in concert during alkalization, with KSPER activation helping to maintain a negative sperm membrane potential (\(V_m\)) sufficient to promote influx of Ca\(^{2+}\) through CATSPER channels (Navarro et al., 2007). KSPER activation is the primary or only determinant of \(V_m\) of mouse spermatozoa during alkalization (Navarro et al., 2007; Zeng et al., 2011), serving to drive sperm \(V_m\) more negative than \(-40\) mV. Simultaneous with the KSPER-mediated hyperpolarization, activation of CATSPER during alkalization (Ren et al., 2001; Kirichok et al., 2006) mediates influx of Ca\(^{2+}\) necessary for activation of Ca\(^{2+}\)-dependent processes required for sperm hyperactivation. CATSPER is thought to be assembled from a set of four distinct CATSPER subunits (CATSPER1–4; Carlson et al., 2003; Quill et al., 2003; Jin et al., 2005) and at least three accessory subunits (\(\beta\), \(\gamma\), and \(\delta\); Liu et al., 2007; Wang et al., 2009; Chung et al., 2011). Genetic deletion of any of the four CATSPER subunits (Qi et al., 2007) or the \(\delta\) subunit (Chung et al., 2011) abolishes sperm hyperactivated motility and results in male infertility. In fact, the initial demonstration of KSPER current in mouse sperm benefited from the use of CATSPER-null sperm (Navarro et al., 2007). KSPER activation increases over the range of pH 6.0–8.0, allowing it to play a major hyperpolarizing role during alkalization. The identity of the critical pore-forming subunit of KSPER was recently established, with the demonstration that genetic KO in mice of the pH-regulated SLO3 K\(^+\) channel (Santí et al., 2010; Zeng et al., 2011) reduces or abolishes KSPER and results in infertile male mice. Supporting the idea that KSPER encoded by the SLO3 gene plays a predominant role in defining sperm \(V_m\) during capacitation, a SLO3-dependent progressive increase in hyperpolarization measured by voltage-sensitive dyes during exposure to capacitating conditions has been observed in mouse sperm (Chávez et al., 2013).

Together, CATSPER and KSPER appear to be the primary participants in regulation of sperm \(V_m\) and Ca\(^{2+}\) influx during alkalization (Kirichok et al., 2006; Navarro et al., 2007; Zeng et al., 2011). However, it is possible that other unidentified currents may be present either in resting conditions or after alkalization. In fact, after genetic deletion of SLO3, although most outward K\(^+\) current was abolished, voltage steps above \(-100\) mV result in a slowly activating increase in outward current (Zeng et al., 2011). This current was proposed to arise from monovalent cation efflux through CATSPER channels, but this proposal requires confirmation. Here, by generating animals in which both Slo3 and CatSper1 genes have been disrupted, we explicitly tested whether the residual K\(^+\) current in Slo3 \(^{-/-}\) sperm cells can be explained by CATSPER and whether additional currents can be identified in mouse sperm cells. In the absence of both CATSPER and KSPER, we were unable to detect any residual K\(^+\) current, either under resting conditions at pH 6.0 or during alkalization at pH 8.0. Although other factors or ligands may result in activation of ion channels not observed here, it is remarkable the extent to which only two ion channels may account for all of the changes in \(V_m\) and Ca\(^{2+}\) elevation that are associated with the hyperactivation process.

**MATERIALS AND METHODS**

**Generation of double KO (dKO) mice**

Both Slo3 \(^{-/-}\) and CatSper1 \(^{-/-}\) male mice are infertile, whereas females are reproductively normal. Generation of the dKO mice required, first, generation of Slo3 \(^{-/-}\), CatSper1 \(^{-/-}\) males, followed by breeding of the heterozygous males with Slo3 \(^{-/-}\), CatSper1 \(^{-/-}\) females. Such a mating is expected to result in full dKOs in \(-25\%\) of pups. Genotypes of all animals were confirmed by PCR. All animal husbandry and experimental procedures were approved by and performed in accordance with guidelines of the Animal Studies Committee of the University School of Medicine.

**Electrophysiology**

Before whole-cell access, for the dKO cells studied here, about half had pipette seal resistances between 5 and 10 GΩ, with the others >10 GΩ. Thus, pipette leak conductance (\(G_L\)) is typically \(-0.2\) nS, but on average \(-0.1\) nS. In all whole-cell recordings, cytosolic access was gained through the sperm cytosolic droplet (Kirichok et al., 2006; Navarro et al., 2007; Qi et al., 2007). HEPES-buffered saline (HS) was used for sperm swim-out from corpus epididymis and recordings: 135 mM NaCl, 5 mM KCl, 1 mM MgSO\(_4\), 2 mM CaCl\(_2\), 20 mM HEPES, 5 mM glucose, 10 mM lactic acid, and 1 mM Na pyruvate, pH 7.4 with NaOH. For most voltage-clamp experiments, cells were bathed in a high K\(^+\) HS saline: 160 mM KOH, 10 mM HEPES, 150 mM methanesulfonic acid (MES), and 2 mM Ca(MES)\(_2\), adjusted to pH 7.4 with MES. The usual high K\(^+\) pipette solution contained 155 mM KOH, 5 mM KCl, 10 mM BAPTA, 20 mM HEPES, and 115 mM MES with pH adjusted to 6.0, 7.0, or 8.0 with KOH or MES. For current clamp recordings in which NH\(_4\)Cl was used to alter cytosolic pH, the following pipette solution was used: 144 mM KOH, 5 mM KCl, 5 mM NaCl, 3 mM MgATP, 0.5 mM Na\(_2\)GTP, 1 mM BAPTA, 5 mM HEPES, and 140 mM MES with pH adjusted to either 6.0 or 8.0 with MES or KOH. For testing for the presence of Ca\(^{2+}\)-activated K\(^+\) currents, a K-MES cytosolic solution with 10 µM Ca\(^{2+}\) contained 140 mM K-MES, 20 mM KOH, 10 mM HEPES (titrated to pH 7.0), and 5 mM HEDTA titrated with Ca-MES to obtain 10 µM Ca\(^{2+}\) as defined by a Ca\(^{2+}\)-sensitive electrode calibrated with a commercial set of Ca\(^{2+}\) standards (World Precision Instruments). Solutions were applied directly via a local perfusion system allowing switching between different test solutions. Solution exchange time with this system is typically \(-1\) s. Current waveforms were analyzed with Clampfit (Molecular Devices). Points and error
bars on figures correspond to mean ± SEM. All experiments were conducted at room temperature (22–25°C). Chemicals were obtained from Sigma-Aldrich.

**RESULTS**

Simultaneous deletion of Slo3 and CatSper1 abolishes all voltage- and alkalization-activated current

Whole-cell recordings from the cytoplasmic droplet were used to compare K⁺ currents in WT, Slo3⁻/⁻, and Slo3⁻/⁻ CatSper1⁻/⁻ (dKO) mouse spermatozoa. With symmetrical K⁺ solutions and a pipette solution of pH 8.0, WT sperm exhibit a pronounced voltage-dependent K⁺ conductance (Fig. 1A), consistent with the previously described KSPER current (Navarro et al., 2007). In Slo3⁻/⁻ sperm, total K⁺ current was markedly reduced (Fig. 1A), with some small residual voltage-dependent K⁺ current that becomes particularly apparent at potentials >100 mV (Fig. 1B; Zeng et al., 2011). It has been previously proposed that the residual K⁺ current in Slo3⁻/⁻ sperm arises from monovalent cation efflux through CATSPER channels (Zeng et al., 2011). Consistent with this suggestion, voltage ramp protocols (Fig. 1A) and steps (Fig. 1B) applied to dKO sperm failed to activate any voltage-dependent K⁺ current despite a pipette pH of 8.0. For a set of Slo3⁻/⁻ sperm and dKO sperm studied with the voltage step protocol, the K⁺ conductance at potentials negative to 0 mV was <0.2 nS for both Slo3⁻/⁻ and dKO sperm (Fig. 1C). For the dKO sperm, this conductance corresponds to a mean resistance of 6.7 GΩ (Fig. 1C, dashed red line). In contrast, at positive potentials, net conductance markedly differed between the two. We are unable to exclude the possibility that there may be a very small increase in conductance in the dKO sperm at potentials >100 mV. However, command steps to such potentials may increase conductance from compromised recording stability. Given an ~0.1–0.2-nS conductance (G_L) of a 5–10-GΩ pipette seal, we were unable to detect any residual K⁺ conductance over the range of voltages from −100 to 100 mV in the dKO sperm that was separable from that expected from G_L. For the three genotypes, net current at 100 mV
was compared (Fig. 1 D). The results indicate that the alkalization-activated current, still present at very positive potentials in the \( \text{Slo3}^{-/-} \) sperm, is absent in the dKO sperm. Thus, the alkalization-activated current in the \( \text{Slo3}^{-/-} \) sperm arises from monovalent cation flux through CATSPER channels. Furthermore, after deletion of both SLO3 and CATSPER subunits, no remaining alkalization-activated K⁺ current was detected. Despite the absence of alkalization-activated K⁺ current in the dKO sperm, application of 100 µM of extracellular ATP to dKO sperm elicited a desensitizing, reversible, rectifying conductance (Fig. 1 E) in the dKO sperm, consistent with the previous observation of a P2X2 current in mouse sperm (Navarro et al., 2011).

The small residual conductance in the dKO sperm is certainly dominated by GL but might also include some small contribution of membrane ion channels, which might be important in intact sperm when KSPER and CATSPER are not activated. As an additional test of the nature of any conductance that might still be present in dKO sperm with pipette pH 8.0, we used quinidine, a rather general ion channel blocker which has been shown to block SLO3 currents (>90% inhibition at 100 µM quinidine at >100 mV; Tang et al., 2010; Zeng et al., 2011) as well as CATSPER (Zeng et al., 2011). In addition, quinidine also inhibits two pore leak channels (e.g., TASK2: 65% inhibition at 100 µM [Reyes et al., 1998]; and TASK3: 37% block by 100 µM [Kim et al., 2000]) that have also been proposed as possible sperm ion channels (Barfield et al., 2005), in addition to Kv channels (Fedida, 1997; Wang et al., 2003). Thus, because of the general effectiveness of quinidine on a host of cation channels, any sensitivity of the residual currents in the dKO sperm to quinidine might suggest the presence of additional types of channels. In WT sperm, 100 µM quinidine substantially inhibits both outward and inward currents recorded in symmetric K⁺ solutions with pipette pH of 8.0 (Fig. 2, A and B). This reflects inhibition of both KSPER and CATSPER currents (Zeng et al., 2011). Note that the residual conductance in WT sperm at negative potentials after application of 100 µM quinidine approaches that of the dKO sperm (Fig. 2, B and D). When 100 µM quinidine was applied to dKO sperm at pH 8.0 (Figs. 1 D and 2, C and D), the residual current was little affected. The effects of 100 µM quinidine on conductances measured from −100 to 100 mV were determined for six dKO sperm (Fig. 2 E). Individual sperm in some cases exhibited small decreases in current during application of quinidine (e.g., Fig. 2 C). Although the differences measured over the set of cells was not significant, the mean conductance in the presence of quinidine (Fig. 2 C) corresponds to a decrease of 12.7 ± 4.9% compared with control levels. We conclude that quinidine-sensitive ion channels make minor contributions, if any, to the residual conductance of dKO sperm.

Figure 2. Residual current in dKO sperm is insensitive to 100 µM quinidine. (A) Currents were activated in a WT sperm with the indicated voltage protocol with symmetrical K⁺ gradients and pH of 8.0. (left) Control saline. (right) 100 µM quinidine. Red traces correspond to 100-mV step. (B) In another WT sperm, the ±100-mV voltage ramp was used to activate currents before, during, and after application of 100 µM quinidine (red), highlighting effects of quinidine at more negative voltages. (C) Currents were activated in a dKO sperm as in A without (left) and with (right) 100 µM quinidine. Red traces correspond to 100 µM quinidine. Red traces correspond to 100 µM quinidine. Red traces correspond to 100 µM quinidine. (D) Ramp-activated currents in a dKO spermatozoa were monitored before, during (red), and after 100 µM quinidine. (A–D) The dashed lines represent 0 current level. (E) Mean conductances were calculated at the indicated voltages for six dKO sperm without (black) and with (red) 100 µM quinidine. There were no significant differences between groups. Error bars indicate SEM.
As a final evaluation of this issue, we examined ramp-activated currents over the range of −90 to 90 mV at high gain (Fig. 3). Such traces were filtered at 2 kHz, and a linear leak current (Fig. 3, A and B) defined by the slope of the currents over the range of −30 to 30 mV was subtracted from the records. According to this procedure, asymmetries around the 0 current level would be potentially indicative of additional voltage-dependent conductances. Close inspection of raw traces suggested that in the dKO sperm, some sperm exhibited a small nonlinear increase in current at both the most negative and most positive voltages (Fig. 3, A and B). After filtering and subtraction, currents from Slo3−/− sperm at pH 8.0 (Fig. 3 C) show the upward curvature associated with monovalent flux through CATPSER channels, while also showing a small asymmetric increase in current variance at the most negative potentials. For dKO sperm at pH 8.0 (Fig. 3 D), there remained some small asymmetric current at both positive and negative potentials, with the currents at negative potentials indistinguishable from Slo3−/− sperm and from those in dKO sperm at pH 6.0 (Fig. 3 E). Whatever the origins of this asymmetric increase in current, it is not only insensitive to pH but also to 100 µM quinidine (Fig. 3 F). We suspect that the small asymmetric currents at very negative and positive potentials may reflect recording instabilities (i.e., unstable seal conductance).

The resting potential of dKO sperm measured by patch-clamp is insensitive to changes in cytosolic pH. Activation of KSPER is sufficient to drive Vm changes during cytosolic alkalization in mouse spermatozoa (Navarro et al., 2007; Zeng et al., 2011). In WT sperm with physiological cation gradients, NH4Cl-mediated alkalization typically results in a pronounced hyperpolarization (Fig. 4 A), whereas in Slo3−/− sperm, alkalization produces a small depolarization presumably because of CATPSER activation (Zeng et al., 2011). In current clamp experiments with physiologically normal Na+/K+ cation gradients, we measured membrane potentials in dKO sperm during NH4Cl application (Fig. 4 B). With a pipette pH of 6.0, Vm in dKO sperm was indistinguishable from 0 mV (Fig. 4 C) and NH4Cl induced no obvious changes in potential. Likewise, with an intracellular (pipette) solution of pH 8.0, the dKO sperm Vm was indistinguishable from 0 mV (Fig. 4 D).
similar to earlier results for the \( \text{Slo3}^{-/-} \) KO sperm (Zeng et al., 2011). These results further support the idea that there are no alalkalization-activated \( K^+ \) conductances other than KSPER available to influence mouse sperm membrane potential during sperm alkalization. This conclusion must be tempered by the fact that any residual conductance, in order to influence \( V_m \) under our experimental conditions, would have to be an appreciable fraction of \( G_L \).

**Elevated cytosolic \( Ca^{2+} \) does not activate additional conductances in dKO mouse sperm**

The absence of KSPER and CATSPER currents in the dKO sperm provide a useful background environment to test for the presence of other conductances. Given the elevations in cytosolic \( Ca^{2+} \) that occur during CATSPER activation (Carlson et al., 2003; Wennemuth et al., 2003), we tested for the presence of known \( Ca^{2+} \)-activated \( K^+ \) currents. We used a 10 \( \mu M \) \( Ca^{2+} \) pipette solution with symmetrical \( K^+ \) to assess whether there might be any contributions of \( Ca^{2+} \)-activated \( K^+ \) conductances (Fig. 5, A and B). Such ionic conditions would be expected to be sufficient to result in activation of all known forms of \( Ca^{2+} \)-activated \( K^+ \) currents (Vergara et al., 1998), including the \( Ca^{2+} \)- and voltage-dependent BK-type \( K^+ \) channels, the voltage-independent small conductance SK-type \( Ca^{2+} \)-dependent channels, and intermediate conductance IK-type \( K^+ \) channels. For four dKO sperm, the measured currents over the range of \(-100 \) to \( 100 \) mV were indistinguishable from currents recorded from dKO sperm with low cytosolic \( Ca^{2+} \) and \( pH \) 8.0 (Fig. 5 B). Based on the inability of 10 \( \mu M \) \( Ca^{2+} \) and positive voltages to increase \( K^+ \) current in the dKO sperm, it is unlikely that any known \( Ca^{2+} \)-dependent \( K^+ \) currents are expressed in mouse sperm.

**DISCUSSION**

Previous work on mouse sperm established that the alkalization-activated KSPER current accounts for most of the \( K^+ \) current in CATSPER-deficient corpus epididymal sperm (Navarro et al., 2007). More recently, alkalization-activated \( K^+ \) current in testicular sperm was shown to be reduced after \( \text{Slo3}^{-/-} \) (Santi et al., 2010), and, independently, it was shown the \( \text{Slo3}^{-/-} \) abolished all KSPER current in mouse corpus epididymal sperm (Zeng et al., 2011). Yet, at very positive potentials, a pharmacologically distinct, slowly activated outward current was present in the \( \text{Slo3}^{-/-} \) spermatozoa (Zeng et al., 2011). Although we attributed this current to monovalent \( K^+ \) efflux through CATSPER that persists despite the presence of 2 \( mM \) extracellular \( Ca^{2+} \), this result appeared to differ from an earlier result showing that, in cells with 140 \( mM \) \( Cs^+ \) pipette solution, 2 \( mM \) extracellular \( Ca^{2+} \) largely prevented any outward current at positive potentials through CATSPER (Kirichok et al., 2006). Our present results now show unambiguously that the residual outward current flux present in \( \text{Slo3}^{-/-} \) sperm can only reflect \( K^+ \) efflux through CATSPER channels. We would suggest that differences in the effectiveness of 2 \( mM \) \( Ca^{2+} \) in reducing monovalent cation efflux probably arise from the differential relative permeabilities of \( K^+ \) versus \( Cs^+ \) and their ability to compete with \( Ca^{2+} \) that occupies CATSPER pore sites. This would be generally consistent with differences in relative permeabilities of \( K^+ \) and \( Cs^+ \) in classical studies of selectivity of \( Ca^{2+} \) selective channels (Hess et al., 1986). Despite the ability of \( K^+ \) to permeate CATSPER channels in the presence of extracellular \( Ca^{2+} \) as shown by the present experiments, it should be
noted that at potentials negative to 0 mV with physiological ionic gradients, K+ efflux will most likely be negligible. However, rigorous estimates of the selectivity of CATSPER channels to different ions will be required to fully address this topic.

Here we now assess the implications of these results for two situations, during alkalization and then under conditions of low cytosolic pH. The present results further support the view that, during alkalization, KSPER and CATSPER are the primary and perhaps the only two ionic currents present in the mouse corpus epididymal sperm principal piece. After deletion of both CATSPER and KSPER, the sperm membrane potential is indistinguishable from 0 mV at pH 8.0. The patch-clamp method as used here is unable to identify any other current other than KSPER and CATSPER that is activated by voltage and alkalization, and any residual current at pH 8.0 in the dKO sperm cannot be distinguished within the background of pipette leak conductance. Any minor K+ currents hidden in the pipette leak conductance would essentially be irrelevant to sperm function and physiology as long as CATSPER and KSPER remain activated by alkalization.

The situation under so-called resting conditions of lower pH is less clear. Although we have made measurements at pH 6.0, what resting conditions actually are when sperm first enter the female reproductive tract are not entirely clear. As with the results at pH 8.0, our recordings both in voltage-clamp and current-clamp are dominated by the pipette leak conductance. However, for the pH 6.0 condition, the possibility that there may be currents buried in the background pipette leak conductance is potentially more critical to sperm physiology. In a cell of high input impedance, even a mean conductance is potentially more critical to sperm function and fertility. To date, of ion transport proteins that have been proposed to be expressed in sperm, only green fluorescent protein (GFP)-expressing SLO3 ('Slo3') sperm have reported Vm of about $-7 \text{ mV}$ (Navarro et al., 2007) and $-13 \text{ mV}$ (Zeng et al., 2011). Here, for a set of WT sperm monitored under identical conditions, Vm was about $-18 \text{ mV}$ (Fig. 4 C), with notable variability in individual measurements. The variability in estimates is probably not surprising because both differences in Gli among cells and variation in the SLO3 current density among cells might be expected to contribute to such differences. Both in previous work (Zeng et al., 2011) and here, Vm at pH 6.0 was somewhat depolarized in Slo3$^{-/-}$ sperm relative to WT sperm. This supports the view that some weak activation of KSPER at pH 6.0 can influence Vm. However, recent results estimating Vm with potential-sensitive dyes in WT and Slo3$^{-/-}$ sperm suggest both that SLO3 contributes little to Vm in mouse sperm under noncapacitated conditions and that such sperm have a measurable negative membrane potential arising from other, as yet unidentified K+ channels (Chávez et al., 2013). The present work suggests that if such currents are present in mouse sperm, they are invisible to the patch-clamp technique. To confirm the existence of such currents, molecular deletion of the relevant subunits will probably be required.

Despite the abundance of channels that have been proposed to be present in sperm (Barfield et al., 2005; Darszon et al., 2007), very few transporter/channel proteins have met the test of being indispensable to sperm function. To date, of ion transport proteins that have been proposed to be expressed in sperm, only genetic deletion of CATSPER-related subunits (Quill et al., 2003; Carlson et al., 2005; Jin et al., 2007; Qi et al., 2007; Chung et al., 2011), the SLO3 subunit (Santi et al., 2010; Zeng et al., 2011), a Ca-ATPase (PMCA4; Okunade et al., 2004; Schuh et al., 2004), a sperm-specific Na,K-ATPase a4 isoform (Jimenez et al., 2011), and a sperm-specific Na-H+ exchanger (NHE5; Wang et al., 2003) have resulted in clear loss of mouse fertility in a fashion related to sperm function and not as the result of a developmental defect. It is notable that, of these five examples, four involve sperm-specific proteins, whereas PMCA4 has a broader tissue distribution. Given the pace of genetic deletions of various categories of ion channels, the absence of additional cases of loss of male fertility seems telling. Of course, if a channel/transporter important to sperm function was also more broadly expressed, the genetic deletion approach might result in deleterious consequences that preclude tests of sperm function and fertility.

In sum, under the conditions of the present experiments, we were unable to observe any conductance above the background of Gli other than the cation fluxes.
arising from KSPER and CATSPER, either at pH 6.0 or pH 8.0. Furthermore, our tests with elevated pipette Ca2+ suggest that any Ca2+ elevation that would be expected to occur as a consequence of CATSPER activation would be unlikely to activate secondary K+ conductances, unless such conductances also require additional regulatory cofactors. Based on the observations presented here, once alkalization is initiated in mouse sperm, the critical ionic currents that control sperm Vm and the elevation of Ca2+ can be accounted for by only two channels, KSPER, arising from the expression of SLO3 subunits, and CATSPER. Because CATSPER is essentially the only current component present after genetic KO of the SLO3 subunit, the Slo3+/− sperm may serve as a useful tool for investigation of a relatively uncontaminated CATSPER current, at least until successful heterologous expression of CATSPER channels is accomplished.

We thank the Department of Anesthesiology, Washington University in St. Louis School of Medicine, for their support of this project. We thank Drs. Chengtao Yang and Vivian Gonzalez-Perez for comments on the manuscript.

This work was also supported in part by National Institutes of Health (NIH) grant GM081748 to C.J. Lingle, Natural Science Foundation of China grants 31070767 and 31171116 to X.-H. Zeng, and NIH grant U01 HD045857 to D.E. Clapham.

Angus C. Nairn served as editor.

Submitted: 18 April 2013
Accepted: 2 August 2013

REFERENCES


