## MscS-Like10 is a stretch-activated ion channel from *Arabidopsis thaliana* with a preference for anions

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Like many other organisms, plants are capable of sensing and responding to mechanical stimuli such as touch, osmotic pressure, and gravity. One mechanism for the perception of force is the activation of mechanosensitive (or stretch-activated) ion channels, and a number of mechanosensitive channel activities have been described in plant membranes. Based on their homology to the bacterial mechanosensitive channel MscS, the 10 MscS-Like (MSL) proteins of Arabidopsis thaliana have been hypothesized to form mechanosensitive channels in plant cell and organelle membranes. However, definitive proof that MSLs form mechanosensitive channels has been lacking. Here we used single-channel patch clamp electrophysiology to show that MSL10 is capable of providing a MS channel activity when heterologously expressed in Xenopus laevis oocytes. This channel had a conductance of  $\sim 100$  pS, consistent with the hypothesis that it underlies an activity previously observed in the plasma membrane of plant root cells. We found that MSL10 formed a channel with a moderate preference for anions, which was modulated by strongly positive and negative membrane potentials, and was reversibly inhibited by gadolinium, a known inhibitor of mechanosensitive channels. MSL10 demonstrated asymmetric activation/ inactivation kinetics, with the channel closing at substantially lower tensions than channel opening. The electrophysiological characterization of MSL10 reported here provides insight into the evolution of structure and function of this important family of proteins.

The perception of mechanical stimuli like gravity, touch, or osmotic pressure is essential to normal plant growth and development and is further implicated in biotic and abiotic stress responses (1). One of the best-studied strategies for perceiving force involves membrane-embedded channels that are gated by tension, known as mechanosensitive (MS) channels (2). Numerous MS channel activities (>17 to date) have been described in the membranes of diverse tissues from a variety of plant species (summarized in ref. 3, also refs. 4 and 5). Many of these observed MS channel activities differ in their conductance, ion selectivity, and/or sensitivity to the direction of activation pressure, suggesting that multiple classes of mechanosensitive channels are present in plant cells.

No mechanosensitive ion channel activity discovered in plant membranes has yet been definitively identified at the molecular level, but two families of proteins serve as strong candidates. The first is the Mid1-Complementing Activity (MCA) family, members of which are required for root response to touch in the model plant Arabidopsis thaliana, induce Ca<sup>2+</sup> uptake in rice and Arabidopsis cells (6, 7), and are associated with increased current in response to hypotonic stimulation of *Xenopus* oocytes (8). The second family of candidates for plant MS channels is the MscS-Like (MSL) family, first identified based on modest homology to the wellcharacterized bacterial MS channel MscS from Escherichia coli (3, 9). MscS is a largely nonselective stretch-activated channel that is gated directly by membrane tension, generating a 1-nanoSiemen (nS) conductance (reviewed in refs. 10 and 11). The primary function of MscS is to provide a conduit for the release of osmolytes from the bacterium in response to extreme hypoosmotic stress (12, 13).

There are 10 MscS homologs in *Arabidopsis*, three with predicted or observed localization to organellar membranes and seven with predicted or observed localization to the plasma and vacuolar membranes (3, 5, 14). Reverse genetic analyses have demonstrated that two plastid-localized homologs, MSL2 and MSL3, are required for the proper size, shape, division, and hypoosmotic volume control of plastids (14–16). In addition, two plasma membranelocalized homologs, MSL9 and MSL10, are genetically required for the predominant MS channel activity in the plasma membrane of *Arabidopsis* root cell protoplasts, providing support for the hypothesis that MSL proteins form MS channels (5). A MscS homolog from the unicellular green alga *Chlamydomonas reinhardtii*, MSC1, has MS channel activity when expressed in giant *E. coli* spheroplasts (17).

However, it has been proposed that the contribution of MSL9 and MSL10 to MS channel activity in root protoplasts may be indirect (1), and a more rigorous test by expression and electrophysiological characterization in a heterologous system is needed. Indeed, it seems likely that MSLs do not form MS channels, because the plasma membrane-localized MscS homologs from plants are highly divergent from MscS with respect to topology and show limited sequence similarity, even within the conserved domain (Fig. S1). Some bacterial MscS homologs do not appear to provide MS channel activities, further suggesting that a subset of the family has evolved to perform diverse functions (reviewed in ref. 18). We were thus motivated to develop a system for the electrophysiological investigation of MSL proteins to determine whether eukaryotic plasma membrane-localized MscS homologs indeed form MS channels and, if so, to compare their electrophysiological behavior with that of other MscS homologs. Here, we describe the heterologous expression and characterization of Arabidopsis MSL10 in Xenopus laevis oocytes.

## Results

MSL10 Forms a ~100-pS Mechanosensitive Channel in the Plasma Membrane of Xenopus Oocytes. We chose to characterize MSL10 in X. laevis oocytes, an established system for the expression and electrophysiological characterization of heterologous ion channels, including those from plants (19). The endogenous MS channels of Xenopus (20) were effectively inactivated upon excision of the membrane patch, as reported (21). Xenopus oocytes produced strong GFP signal at their periphery by 48 h after injection with MSL10-GFP cRNA, indicating that the MSL10 protein is efficiently translated and localized in or near the plasma membrane (Fig. 1A).

Inside-out patches excised from oocytes injected with cRNA encoding either MSL10-GFP or untagged MSL10 reproducibly exhibited channel activity in response to membrane stretch in symmetric ND96 buffer (Fig. 1*B*), although we consistently observed higher activity when untagged MSL10 was used. MSL10 channel activity depended on an increase in membrane tension,

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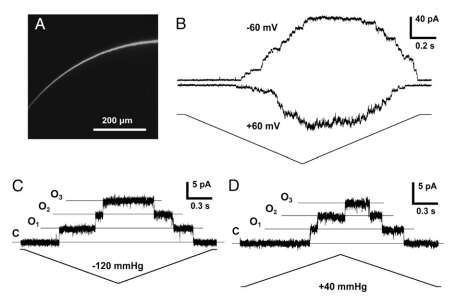


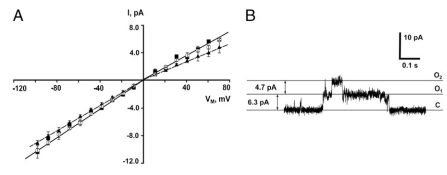
Fig. 1. MSL10 expressed in *X. laevis* oocytes produces a channel activity upon membrane stretch. (A) Confocal image of a portion of an oocyte 5 d after injection with MSL10-GFP cRNA. (*B*) A representative trace illustrating the activation of MSL10 channels at both negative and positive membrane potentials in the same patch. Pipette bubble number (BN) 6, -70 mmHg. (*C*) Single-channel opening events induced by negative pipette pressure. Membrane potential -50 mV, pipette BN 4. (*D*) Single-channel opening events induced by positive pipette pressure. Membrane potential -40 mV, pipette BN 5. Symmetric ND96 buffer was used in *B*-D.

but we observed that the same tension resulted in different current amplitudes at opposite membrane potentials in the same patch (representative traces are shown in Fig. 1*B*). Under negative membrane potentials, MSL10 single-channel events were easy to observe because of low noise and their stable behavior, whereas under positive membrane potentials, noise and flickery behavior were dominant. The ratio of peak current amplitude at negative to positive membrane potentials was  $1.24 \pm 0.15$  (n = 7 oocytes). The results at negative membrane potentials are most likely to represent the behavior of the MSL10 *in planta*, because the transmembrane potential of *Arabidopsis* root cells has been measured at approximately -180 mV (for example, see ref. 22). MSL10 singlechannel openings were readily detected in response to membrane stretch generated by both negative (Fig. 1*C*) and positive (Fig. 1*D*) pipette pressures.

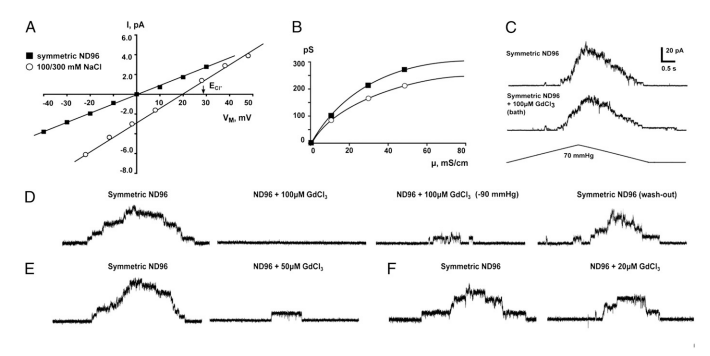
The current-voltage (I/V) curves for MSL10 and MSL10-GFP indicated that the single-channel conductances of both MSL10 and MSL10-GFP were  $103 \pm 3$  pS in excised patches, measured as the slope of the I/V curve within the range of 0 to -60 mV (Fig. 24). We did not use data from potentials lower than -60 mV in our calculation of conductance because of the presence of

conducting substates (an example of which is shown in Fig. 2*B*). A conductance of  $103 \pm 3$  pS is in good agreement with an MSL10dependent activity present in *Arabidopsis* root protoplasts, which was measured at 137 pS under slightly different ionic conditions (5). Although the I/V curve for MSL10 was linear at positive and negative membrane potentials, the slopes were slightly different under the two conditions, with a single-channel conductance of  $80 \pm 2$  pS at positive membrane potentials. This slight current rectification may explain the 1.24-fold difference in current under positive and negative potentials described above and shown in Fig. 1*B*.

**MSL10 Exhibits a Moderate Preference for Anions.** The reversal potential of MSL10 under asymmetric 100/300 mM salt was +19 mV (Fig. 3*A*), and the theoretical reversal potential of Cl<sup>-</sup> ion, derived from the Nernst equation under a threefold gradient of ion concentration, is +28 mV. The Goldman–Hodgkin–Katz equation gives a ratio of Cl<sup>-</sup> to Na<sup>+</sup> permeability ( $P_{Cl}$ : $P_{Na}$ ) of 5.9 for MSL10. We also measured the conductance of MSL10 when Na<sup>+</sup> was replaced with TEA<sup>+</sup>, a large ion with an estimated diameter of 8 Å (23) often used as a pore blocker of potassium



**Fig. 2.** MSL10 and MSL10-GFP single-channel conductance. (*A*) The current–voltage curves for untagged MSL10 (open circles) and MSL10-GFP (filled squares) in symmetric ND96 buffer, and for untagged MSL10 in symmetric 98 mM TEA-Cl (filled triangles) (n = 5 oocytes for each protein). Solid and dashed lines represent linear fits for the channels in ND96 and TEA-Cl buffers, respectively. (*B*) A typical trace illustrating the appearance of conductive substates at more negative membrane potentials, in this case -60 mV. Pipette BN 5, symmetric ND96 buffer.

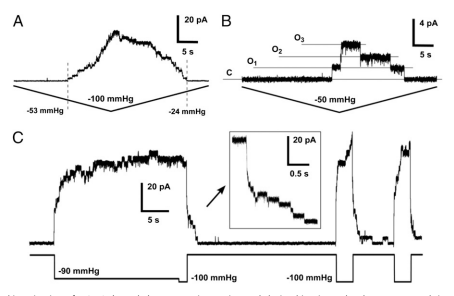


**Fig. 3.** Ion selectivity and  $Gd^{3+}$  inhibition of MSL10. (A) Current–voltage curves for MSL10 in symmetric ND96 (96 mM NaCl, filled squares) and in asymmetric 100 mM/300 mM NaCl buffer (open circles).  $E_{CL}$ , reversal potential for  $CL^-$  ions. (B) Single-channel conductance under increasing [NaCl] at negative (filled squares) and positive (open circles) membrane potentials. Buffers containing 4 mM MgCl<sub>2</sub> and 5 mM Hepes supplemented with 100, 300, or 500 mM NaCl were used, membrane potential –30 mV. (C) Representative traces showing MSL10 channel activity in the same outside-out patch (pipette BN 5, membrane potential for P Representative traces showing MSL10 channel activity in the same patch before and after perfusion with 100  $\mu$ M GdCl<sub>3</sub> (*D*), or 20  $\mu$ M GdCl<sub>3</sub> (*F*). In *D*, a trace from the same patch after washout is shown at right. Membrane potential –40 mV, BN 5, in symmetric ND96 supplemented with the indicated amounts of GdCl<sub>3</sub> from the bath side. Pressure applied to the pipette was –60 mmHg in all cases except for the third trace in *D*, where the pressure was –90 mmHg.

channels (24, 25). Because Hille's equation (26) also predicts an approximate pore diameter of 8 Å for MSL10 (assuming the pore to be a uniform cylinder 5 nm in length), TEA<sup>+</sup> is not likely to permeate the MSL10 channel pore, nor does TEA<sup>+</sup> serve to block MSL10 (Fig. 24). The single-channel conductance of MSL10 in 100 mM symmetric TEA-Cl was ~84% of that measured in symmetric ND96 (96 mM NaCl). This result is consistent with the P<sub>Cl</sub>:P<sub>Na</sub> ratio of 5.9 for MSL10 calculated from Fig. 3A, which predicts that 83% of the MSL10 current in ND96 is provided by Cl<sup>-</sup>. The MSL10-dependent ion channel activity previously characterized in root protoplasts showed no change in conductance when CaCl<sub>2</sub> in the bath solution was replaced with TEA-Cl, whereas current was abolished when Cl- was replaced with MES<sup>-</sup>—indicating that neither Ca<sup>2+</sup> nor TEA<sup>+</sup> can permeate the channel (5). MSL10 single-channel conductance showed saturation at relatively low ionic strength at both positive and negative membrane potentials (Fig. 3B). In summary, MSL10 forms a channel with a moderate preference for anions, passing ~6 chloride ions for every sodium ion.

MSL10 Is Reversibly Inhibited by  $Gd^{3+}$  lons in Inside-Out Patches. Gadolinium ions are commonly used to inhibit K<sup>+</sup>, Ca<sup>2+</sup>, and metazoan MS channels (27) and have also been demonstrated to inhibit the activity of MS channels in plants (28–30) and bacteria (31). Inhibition of MSL10 activity was observed after excised inside-out patches were perfused in a bath containing 100 µM GdCl<sub>3</sub> (Fig. 3D). This inhibition was reversible, and MSL10 activity was recovered upon Gd<sup>3+</sup> washout. Less effective inhibition was observed in patches perfused with 50 or 20 µM GdCl<sub>3</sub> (Fig. 3 *E* and *F*), and an identical regime on outside-out patches did not significantly inhibit MSL10 activity (Fig. 3*C*). MSL10 Gating Kinetics and Inactivation. As shown in Fig. 1 B-D, all MSL10 traces-regardless of pipette size, transmembrane potential, or amount of applied pressure-exhibited a peak tension-induced current that was delayed compared with the peak of applied pressure. We used relatively fast ramp speeds ( $\sim 1$  s) in our initial characterization of MSL10, in accordance with previous studies of MscS (e.g., ref. 32) and to reduce artifacts associated with changes in patch structure during recordings (33). Substantially slower ramp speeds (~25 s) still produced the observed asymmetric pressuredependence of channel opening and closing and did not depend on the number of activated channels (Fig. 4 A and B). Increasing the Mg<sup>2+</sup> concentration of bath and pipette solutions, which has been shown to improve membrane-glass adhesion and facilitate gigaseal formation (34), did not alter the slow closing behavior of MSL10 (Fig. S2 A and B). Finally, we sequentially applied pressure ramps of different lengths (1 s, 5 s, and 25 s) to the same patch and compared the pressure at which the first channels opened or the last channels closed at each ramp speed (Fig. S3). The threshold pressure required to open MSL10 channels was reduced with slower ramp speeds, dropping  $1.42 \pm 0.17$ -fold between 1-s and 5-s ramps and  $2.19 \pm 0.34$ -fold between 1-s and 25-s ramps (n =7 patches). In contrast, the ramp pressure at which all MSL10 channels had closed was always below the opening threshold pressure, regardless of the pressure ramp speed. In more than half of these experiments, the last MSL10 channel closed under zero applied pressure. We were not able to use the midpoint gating tension in our characterization of MSL10, because the oocyte membrane routinely ruptured before current saturation regardless of pressure ramp length, although this was not observed with MscS under the same conditions (Fig. S4 A and B).

To gain further insight into MSL10 gating kinetics and adaptive behavior, we applied tension to the membrane in multiple sharp steps, as reported for MscS (35). Under these conditions, MSL10



**Fig. 4.** Gating kinetics and inactivation of MSL10. (*A* and *B*) Asymmetric opening and closing kinetics under slow ramp speeds in patches with many (*A*) and few (*B*) channels. Dashed lines in *A* indicate the first channel opening and last channel closing events. Both traces are 50 s long, membrane potential –20 mV, pipette BN 4.5, symmetric ND96 buffer. (*C*) Slow gating and absence of inactivation under sustained tension. The length of the whole trace is 45 s. Pipette BN 5, symmetric ND96 buffer, and membrane potential –40 mV.

displayed very slow opening and closing kinetics compared with MscS, even after multiple cycles (Fig. 4*C*). Occasionally a fraction of the channels stayed in the open state and did not close after pressure release at higher potentials (Fig. S2*B*). These data indicate that, under a variety of experimental conditions, the MSL10 channel closes at a much lower tension than is required for it to open, and in some conditions can remain open even in the absence of applied pressure. Unlike MscS, which enters a tension-unresponsive state after sustained stimulus (12, 32), we did not detect inactivation of MSL10 even after 10–20 s of sustained tension (Fig. 4*C* and Fig. S2*B*).

## Discussion

The presence of multiple diverse MS ion channel activities in the plasma and vacuolar membranes of land plants has been well documented over the past 20 y (summarized in ref. 3), and two candidate gene families have been identified in the model plant A. thaliana (reviewed in ref. 1). Here, we used single-channel patch clamp electrophysiology to provide direct evidence that a member of one of these families, MSL10, provides a stretchactivated channel activity when heterologously expressed in X. laevis oocytes. MSL10 is likely to represent the activity described in the root protoplasts of a msl9-1;msl10-1 double mutant transiently expressing MSL10, an anion-preferring activity with a conductance of  $\sim 137$  pS at -182 mV (5). Although we were unable to measure MSL10 channel conductance in oocytes at such high potentials, this value is close to the conductance of oocyteexpressed MSL10 under our conditions (103 pS at 0 to -60 mV). In addition to the activity in roots that we can now assign to MSL10, a nonselective channel activity described in Arabidopsis, leaf mesophyll cells (36) also shows a preference for anions ( $P_{CI}:P_{K}$ ratio of 1.9), and may represent the activity of MSL10 or another MscS homolog.

Our characterization of MSL10 channel behavior provides insight into the evolutionary conservation of structure and function between MscS homologs. Fig. S1 shows the known or predicted topology of MscS, MSL10, MSC1, and YbdG, a MscS homolog from *E. coli* to which we refer here as MscM (37, 38). MSL10 has an extended N terminus and a total of six TM helices, whereas MSC1 and MscM have five and MscS has three TM helices. The conserved "MscS domain" as defined here comprises the most C- terminal TM helix (TM3 in MscS) and the upper portion of the hollow cytoplasmic domain, called the  $\beta$ -domain (Fig. S1 *A* and *B*; ref. 39.

Numerous studies on MscS have indicated the functional importance of the residues within the conserved region, primarily in the pore-lining TM3 helix (summarized in ref. 40). For example, the characteristic alteration between small and large hydrophobic residues appear to be responsible for proper TM3 packing in the MscS heptamer (41), and the hydrophobic seal residues L105 and L109 are essential for complete channel closure (39, 41–44). Although these structural motifs are for the most part preserved in MscM and MSC1, MSL10 shares very little homology in the predicted pore-lining region (Fig. S1*C*), instead exhibiting many bulky hydrophobic residues—including six phenylalanines—in the TM3 region. It is therefore surprising that MSL10 has MS channel activity at all, and perhaps even more surprising how closely MSL10 behavior resembles MscS, MscM, and/or MSC1.

**Unitary Conductance.** MSL10 had a unitary conductance of  $103 \pm 3$  pS in symmetric 100 mM NaCl, whereas MscS has a conductance of 330 pS under the same conditions (21) (approximately 1 nS when measured in 200 mM KCl, 90 mM MgCl<sub>2</sub>, and 10 mM CaCl<sub>2</sub>; refs. 32, 41, and 45). Although MscM and MSC1 are highly similar to MscS with respect to the sequence of the pore-lining TM3 helix, they have conductances similar to that of MSL10: 100–150 pS in 100 mM KCl for MscM (37, 38) and 400 pS for MSC1 measured in 200 mM KCl with 40 mM MgCl<sub>2</sub> and 10 mM CaCl<sub>2</sub> (17) (120–130 pS if measured in 100 mM salt). Although it is tempting to speculate that the many bulky hydrophobic residues in the pore-lining helix of MSL10 may be responsible for a smaller pore size—and therefore a smaller conductance than MscS—the smaller conductances of MscM and MSC1 require a different explanation.

**Ion Selectivity.** Although formally nonselective, the MSL10 channel showed a preference for anions, with a  $P_{CI}$ : $P_{Na}$  ratio of 5.9 based on reversal potential and on its conductance when Na<sup>+</sup> was replaced with TEA<sup>+</sup> (Figs. 2*A* and 3*A*). Other MscS family members display diverse ion selectivity: MscS demonstrates a weak preference for anions with  $P_{CI}$ : $P_{K} = 1.2-3.0$  (45–48), MscM a weak preference for cations, with  $P_{CI}$ : $P_{K} = 0.4$  (37), and MSC1 is as anion-selective as MSL10, with  $P_{CI}$ : $P_{K} = 7$  (17). MSL10 also

showed saturation with increased solution conductivity at both positive and negative membrane potentials (Fig. 3*B*). A conductance ratio of approximately 1.3 (negative membrane potential to positive membrane potential) was observed in these experiments, similar to that measured in symmetric ND96 (Fig. 2*B*). Although MscS does not show saturation up to 1.5 M KCl (46), a MscS homolog from the soil bacterium *Corynebacterium glutamicum* saturated at negative, but not at positive, membrane voltages (49).

Gadolinium is a potent inhibitor of mechanosensitive channels of various types (27, 30, 50, 51), but different mechanisms are likely to be involved in each case. Gd<sup>3+</sup> ions inhibit Ca<sup>2+</sup>-selective stretch-activated channels at concentrations as low as 1 µM (30), but concentrations above 100 µM are required to inhibit the essentially nonselective bacterial channels MscS and MscL (31, 51). In the later case, gadolinium ions have been shown to inhibit MS channels through interactions with negatively charged lipids in the membrane (51). Efficient inhibition of MSL10 in inside-out patches was produced only at the highest concentration of gadolinium tested, 100  $\mu$ M (Fig. 3 *D*–*F*). Even in this case, increased tension evoked partial restoration of channel activity. Because the inside but not the outside of the oocyte membranes contain the negatively charged lipids proposed to interact with Gd<sup>3+</sup> (PS, PG, and PI) (51, 52), only Gd<sup>3+</sup> treatment of inside-out patches would be expected to show lipid-mediated inhibition. Indeed, we did not observe consistent  $Gd^{3+}$  inhibition in outside-out patches (Fig. 3C), and conclude that, like MscL, MSL10 is likely to be inhibited by gadolinium ions indirectly through changes in lipid packing or increased membrane stiffness.

Gating Dynamics. The hallmark of MSL10 activation is a dramatic asymmetry of current (hysteresis) with respect to the pressure ramp, suggesting that the membrane tension at which MSL10 opens is higher than the tension at which it subsequently closes. We reproducibly observed this behavior in experiments with pipettes of various diameters (BN 4-7), at opposite membrane potentials, under both positive and negative pipette pressures, with different ramp speeds, buffer compositions, and number of channels activated, making it unlikely to be an artifact (Figs. 1 and 4 and Figs. S2 and S3). The threshold tension for MSL10 opening depended on ramp speed (in the 1-25s range), decreasing on longer ramps; almost no effect on midpoint tension was seen with MscS opening (33), and no dependence on ramp speed was reported for MSC1 opening (17). Unlike MscS, the unusually slow closing kinetics of both MSL10 and MSC1 were not substantially affected by pressure ramp speed. These observations could be explained by different tension dependencies of opening and closing. The relaxation of the outer leaflet of the membrane in an excised patch described in ref. 53 has little effect on the tension at which MSL10 closes, because it stayed very close to zero in the majority of our experiments regardless of the ramp speed.

A related feature of MSL10 activity is its behavior in response to application of sharp pressure steps (Fig. 4*C* and Fig. S2*B*). Under these conditions, residual MSL10 channel activity in the absence of applied pressure (corresponding to no or very weak membrane tension; ref. 54) was frequently observed after application of the threshold tension, although never before it. MscS demonstrates similar behavior upon closing when the G113A or G121A mutations are introduced (35), indicating that only a small change in identity at a key position (G113 forms a sharp kink in the TM3 of MscS; ref. 39) can produce this phenomenon. Only G121 is conserved in MSC1; neither G113 nor G121 is conserved in MSL10 (Fig. S1*C*). Bulky residues at these positions may make the porelining helix of MSL10 stiffer and allow the channel to maintain the open state for a longer period, even when little or no membrane tension is applied.

In summary, MSL10 resembles MSC1 and MscM with respect to unitary channel conductance, MSC1 with respect to ion selectivity, and MSC1 and MscS G113A/MscS 121A with respect to gating kinetics. The similarities and differences between these four channels cannot be easily attributed to sequences previously identified as conserved among MscS family members or important for MscS channel function (summarized in ref. 40). Instead, these comparisons of sequence and electrophysiological characteristics show that there are still discoveries to be made regarding the relationship between structure and function in the MscS family of MS channels.

Our characterization of MSL10 channel behavior also provides insight into its possible *in planta* function. Increased tension in the plasma membrane of a plant cell could result from hypoosmotic swelling, invasion of the cell by a pathogen, or bending of a plant organ. As demonstrated for MscS and MscM (12, 38), the immediate consequences of MSL10 opening could include the release of osmolytes, thereby preventing cell lysis under hypoosmotic shock or mechanical strain. However, its preference for anions leads us to speculate that MSL10 opening would also result in the depolarization of the cellular membrane via Cl<sup>-</sup> efflux. Once open, MSL10 would allow chloride ions to exit the plant cell until membrane tension was completely relieved. A negative feedback mechanism not present in oocyte membranes, such as interaction with signaling molecules, could promote MSL10 channel closing (as proposed for MSC1; ref. 17). In planta, MSL10 gating could lead to the activation of depolarization-activated Shaker-type potassium channels and depolarization-activated Ca<sup>2+</sup> channels, leading to K<sup>+</sup> efflux from the cell, Ca<sup>2+</sup> influx, and possibly the propagation of a systemic signal (55, 56). Thus, the electrophysiological characterization of a MscS homolog from a multicellular system opens up the exciting possibility that some members of this family of MS channels may not only release osmolytes from swelling cells and organelles, but also alter cell physiology and potentially participate in intercellular signaling.

## Methods

**Molecular Biology.** To obtain pOO2-MSL10-GFP, the ORF of MSL10 was introduced into the pOO2-GFP vector (21) between the Xmal and BamHI cites. Site-directed mutagenesis was used to introduce two stop codons between MSL10 and GFP sequences in pOO2-MSL10-GFP, creating pOO2-MSL10. Capped cRNA was transcribed in vitro by SP6 polymerase using the mMessenger mMachine kit (Ambion) and stored at -80 °C until use.

**Oocyte Preparation.** X. *laevis* oocytes (Dumont stage V or VI) were collected and handled as described (21). GFP signal was visible by confocal microscopy within 48 h of injection, but we observed increased channel activity after longer incubation times, oocytes were patched 1–3 wk after injection.

**Confocal Microscopy.** Two to 10 days after injection with pOO2-MSL10-GFP cRNA, devitellinized oocytes (57) were placed on cavity slides and covered with thin coverslips. An Olympus Fluoview-1000 confocal with BX-61 microscope and FV10-ASW Olympus application software suite were used for image acquisition.

**Electrophysiology.** The buffers used were as follows: complete ND96 (96 mM NaCl, 2 mM KCl, 2 mM CaCl<sub>2</sub>, 1 mM MgCl<sub>2</sub>, and 5 mM Hepes at pH 7.38; specific conductivity 13 mS/cm), TEA-Cl (98 mM TEA-Cl, 5 mM Hepes, 2 mM MgCl<sub>2</sub> at pH 7.38 adjusted with TEA-OH), and 60 mM MgCl<sub>2</sub> (with 2 mM Hepes). All traces were obtained from inside-out (excised) patches except for that shown in Fig. 3C, which came from an outside-out excised patch. Experiments in asymmetric buffers, symmetric high salt buffers, and gado-linium-containing buffer used Rainin Minipulse3 peristaltic pumps. In all measurements with asymmetric buffers, liquid junction potentials were corrected after the patch was broken. Electrode potential drift was tested before the experiments and was less than 0.1 mV per 10 min. The rest of materials and methods are as in ref. 21.

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