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# Plant mechanosensitive ion channels: an ocean of possibilities

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- 1 Plant Mechanosensitive Ion Channels: An Ocean of Possibilities
- 2
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8

#### 9 Abstract

10

Mechanosensitive ion channels, transmembrane proteins that directly couple mechanical 11 stimuli to ion flux, serve to sense and respond to changes in membrane tension in all 12 branches of life. In plants, mechanosensitive channels have been implicated in the 13 perception of important mechanical stimuli such as osmotic pressure, touch, gravity, and 14 pathogenic invasion. Indeed, three established families of plant mechanosensitive ion 15 channels play roles in cell and organelle osmoregulation and root mechanosensing-and 16 it is likely that many other channels and functions await discovery. Inspired by recent 17 discoveries in bacterial and animal systems, we are beginning to establish the conserved 18 19 and the unique ways in which mechanosensitive channels function in plants. 20

2

#### 21 Introduction

#### 22

The ability to sense intrinsic or extrinsic mechanical cues is as basal to the tree of life as 23 the ownership of a cell membrane [1]. Several aspects of growth and development in land 24 plants involve mechanical signals, including touch, osmotic stress, vibration, and gravity 25 responses, the perception of pathogen invasion, and proprioception. One well-26 established component of the mechanosensory apparatus of cells in every kingdom of 27 life is the mechanosensitive (also called stretch-activated) (MS) ion channel [2-4]). These 28 multimeric pore-forming proteins convert mechanical force into ion flux. In some cases, 29 the flow of ions through an open MS ion channel is sufficient for the desired response to 30 mechanical stimulation. For example, the canonical bacterial MS ion channel MscS acts 31 as an osmotic safety valve to protect the cell from hypo-osmotic stress; passage of ions 32 out of the cell through channel directly accomplishes the primary function of the channel 33 [5]. In other cases, mechanosensitive ion flux generates bioelectric signals that in turn 34 trigger organismal sensory perception. For example, the MS ion channel NOMPC 35 36 mediates touch perception in *Drosophila* larvae [6]. The line between the two examples above may not be so clear, as a recent report demonstrated entry of the second 37 messenger Ca<sup>2+</sup> into the bacterial cell through MscS during hypoosmotic shock [7]. In this 38 article, we summarize recent exciting developments in the field of plant MS channels, 39 40 speculate on their evolution, describe a few areas of limited knowledge, and propose potential solutions to technical challenges. 41

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#### **The Tip of the Iceberg: Known Families of Plant Mechanosensitive Channels**

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The first MS channel activities in plant membranes were characterized by patch clamp electrophysiology [8,9] shortly after they were discovered in animal cells (see [10] for a historical perspective). Dozens of MS channel activities in the plasma and vacuolar membranes of a wide variety of cell types and species have been described over the past 30 years (summarized in [11]), suggesting that they are used broadly in plants to respond to diverse signals. Despite this apparent ubiquity, the underlying genes/proteins and physiological function of only a handful of MS ion channel activities have been elucidated.

So far, three MS channel families have so far been characterized as membrane stretch-52 activated in plant systems; as described in further detail below, these channels exhibit 53 diverse, yet overlapping localization, structure, channel properties and proposed function. 54 As a result, the activity of channels with different ionic affinities in the same or in different 55 compartments is likely to result in crosstalk and have complex effects on ion flux into and 56 out of the cytoplasm and apoplast (Figure 1). These three families are unlikely to provide 57 all observed MS channel activities in plants, and a major challenge for the field will be the 58 development of functional (rather that homology-based) screens capable of identifying 59 additional MS channels. Intriguing candidates have been identified [12-14] but have not 60 yet been shown to respond directly to membrane tension. 61

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63 MscS-Like (MSL) Channels. Escherichia coli MscS is one of the best-understood MS ion channels in any system. It is an essentially non-selective ion channel, gated directly 64 by membrane tension, with a large conductance of 1.2 nS. The classic function of EcMscS 65 is to serve as an osmotic safety valve, protecting cells from rupture during extreme hypo-66 67 osmotic downshock. MscS-Like channels, or MSLs, are found throughout bacteria, archaea, some fungi, algae, and plants [15]. MSL gene families have been described and 68 69 characterized to various degrees in Arabidopsis, papaya, rice, and common bean [16-19]. There are 10 MSL proteins in Arabidopsis, most of which are predicted to localize to 70 71 the plasma membrane. Unexpectedly, MSL1, MSL2, and MSL3 were found to localize to the inner membrane of plastids and mitochondria (Figure 1, [20-23]). 72

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Electrophysiological analyses of MSL9 and MSL10 in plant cells [22], MSL10 and MSL8 74 75 expressed heterologously in Xenopus oocytes [23,24], and MSL1 expressed heterologously in giant E. coli spheroplasts [21] all revealed channel characteristics that 76 are similar (though not identical) to *Ec*MscS. MSLs are anion-preferring (e.g. 2 to 6 anions) 77 pass for every cation) MS ion channels with conductances ranging from  $\sim 0.1$  to 1 nS, 78 depending on buffer conditions. Several lines of evidence support the model that, like 79 EcMscS, AtMSLs function to relieve osmotic stress. This was first demonstrated with 80 MSL2 and MSL3, two plastid-localized channels that directly maintain plastid 81 osmoregulation. Plastids in msl2 msl3 mutants exhibit altered size, shape and fission 82

[20,25,26]. The loss of MSL2/3 also leads to stress responses associated with drought 83 and the development of callus tissue at the apex of the plant [27,28]. While the pleiotropic 84 phenotypes associated with this mutant have illustrated the importance of plastid 85 osmoregulation during normal plant growth and development, any mechanistic insights 86 await the electrophysiological analysis of MSL2 and MSL3—a challenging prospect for 87 plastid-localized proteins. Adding to the complexity is a recent report demonstrating that 88 mitochondria-localized MSL1 is required to ameliorate the oxidative burden imposed upon 89 mitochondria during abiotic stress [21]. The potential role of membrane tension, redox 90 state, and transmembrane voltage in regulating MSL1 channel activity in vivo remains to 91 be determined. For plasma membrane-localized MSLs, recent reports both support their 92 role as osmotic safety valves and suggest more complex function, as discussed below. 93

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**Two-Pore Domain K<sup>+</sup> (TPK) Channels.** TREK1, TREK2, and TRAAK are MS channels 95 from the TPK family that are expressed in the mammalian nervous system and are 96 proposed to modulate mechanical, heat and cold-associated pain perception [29]. 97 AtTPK1 is a voltage-independent  $K^{\dagger}$  channel required for normal guard cell closure 98 kinetics [30], and, along with homologs from rice and barley, has been demonstrated to 99 be mechanosensitive [31]. Whether the mechanosensitive activity of AtTPK1 is important 100 for its function in guard cells, and how it is integrated with other regulatory signals such 101 as low pH, Ca<sup>2+</sup> and binding to 14-3-3 proteins is not yet understood [30]. 102

103

Mid1-Complementing Activity (MCA) Channels. The Mid1-Complementing Activity 104 (MCA) proteins were identified based on their ability to rescue the mating-induced lethality 105 106 of the yeast *mid1* mutant [32]. MCA proteins are plant-specific and show no homology to the yeast Mid1 channel. In fact, MCA proteins have only 1 transmembrane (TM) domain 107 [33], placing them outside the norm for ion channel subunits. Cryo-EM imaging followed 108 by single particle reconstruction of a MCA2 tetramer did not reveal a pore [34]. However, 109 heterologously expressed MCA1 and 2 produce increased current in response to osmotic 110 swelling in whole cells and to membrane stretch in excised patches [35], providing 111 evidence that they directly form a MS ion channel. MCA expression is correlated with 112 enhanced Ca<sup>2+</sup> influx in response to hypoosmotic shock and mechanical stimulus in 113

several plant species [32,36,37]. Arabidopsis *MCAs* are required for normal rates of root
penetration into hard agar and for proper response to cellulose biosynthesis inhibition,
implying a role in the maintenance/response to extracellular mechanical stress [32,38].
MCAs may be involved in the perception of developmentally imposed mechanical signals,
as a maize MCA homolog was recently identified in a screen for leaf patterning mutants
[39].

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#### 121 Getting our Sea Legs: Recent Advances in Understanding Plasma Membrane 122 Localized MSL Channels

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MSL8 Fully Meets the Criteria for a Mechanoreceptor. A recent analysis of MSL8, a 124 125 MS ion channel expressed exclusively in mature pollen grains and tubes, advanced our understanding of the function of plasma membrane-localized MSL channels and 126 underscores the essential role of osmoregulation during fertilization. The correct level of 127 MSL8 activity appears critical for pollen to survive hydration and germination and for full 128 129 male fertility. Disruption of MSL8 results in high rates of bursting during pollen hydration and germination, but the overall rate of *in vitro* germination is higher than the wild type. 130 131 On the other hand, overexpressing *MSL8* inhibits pollen germination and no bursting is observed [23]. These opposing effects can be attributed to the inability to relieve excess 132 133 turgor during hydration (in *msl8* mutants) or to maintain necessary turgor during germination, and tube growth (in lines that overexpress MSL8) (Figure 2). Lesions that 134 disrupt the ion conducting properties of MSL8 also disrupt its ability to accomplish these 135 functions in pollen [40], providing further evidence that it serves directly as an osmotic 136 137 mechanosensor in pollen membranes. MSL8 is thus the first plant protein to fill the stated criteria for a mechanoreceptor [2]. 138

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Links Between MSLs and Stress Responses. The role or roles of MSLs at the plasma membrane in cells other than pollen grains has remained stubbornly opaque. Both *MSL* and *MCA* gene expression responds to vibration [41] and nodulation [42], but the physiological relevance of these observations have yet to be demonstrated. While a mutant harboring lesions in 5 *MSL* genes (*msl4 msl5 msl6 msl9 msl10*) ablated the

primary MS channel activity in Arabidopsis root protoplasts, the quintuple mutant does 145 not produce an observable mutant phenotype in response to a wide range of mechanical, 146 touch or osmotic stimuli [22]. However, overexpression of MSL10 results in dwarfing, 147 ROS accumulation, and ectopic lesions, and all of these effects are negatively regulated 148 by phosphorylation of the N-terminus [43]. Dwarfing and ectopic lesions are also observed 149 in response to a single EMS-induced point mutation in the C-terminus of MSL10 [44]. 150 suggesting that these overexpression phenotypes reflect some aspect of the normal gene 151 function. In addition, a recent study implicated MSL4 in pathogen-triggered immunity [45], 152 and MSL6 phosphorylation was observed in response to oligo-galacturonide treatment 153 [46]. We propose that plasma membrane-localized MSLs serve as sensors of cellular 154 mechanical homeostasis, or "mechanostasis". This idea is supported by a recent meta-155 analysis of Arabidopsis microarray datasets wherein MSL10 expression levels were 156 altered in a wide range of mutant backgrounds [47]. 157

158

An intriguing aspect to the MSL10 study was the discovery that the soluble N-terminus of MSL10 is on its own able to trigger cell death in an overexpression system, indicating that the protein harbors at least one function independent of the production of a channel pore [43]. Determining if this non-conducting function is regulated by membrane tension is an important next step. If so, MSLs (and possibly other MS channels or MS channel homologs [39]) may have evolved to couple changes in membrane tension to a wide range of signaling outputs beyond ion flux.

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#### 167 Beyond the Horizon: Innovations in MS Channel Studies

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Plant MS Channel Structure and Gating Dynamics. Structural information about bacterial and animal MS channels derived from a multiplicity of approaches has led to a rapid uptick in our understanding of the structural and biophysical basis of mechanosensitivity. A number of recent reports utilizing crystallography, EPR spectroscopy, PELDOR, and/or molecular dynamics add exciting and provocative new detail to the force-from-lipid concept/principle [1], see Box 1, and suggest that lipid acyl chains filling voids or pockets in the channel surface could "drag" MS channels open under increased membrane tension [48,49] or even block the permeation pathway
[50](but see [51]). While these new ideas are sparking a great deal of discussion in the
field, MS channels from plants have yet to contribute to the conversation. The cryo-EM
structure of MCA2 provides only low resolution information (26 Å) [34], and nothing is yet
known about the structure or even oligomeric state of any MSL channel.

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Solving the structure of plant MSLs would do more than contribute to our view of MS 182 channel gating dynamics. Arabidopsis MSL family members differ substantively from 183 EcMscS (and from each other) not only in terms of the number of TM helices, but in the 184 presence of soluble domains at the N- and C-termini and in inter-TM loops [11,52]. We 185 have previously proposed that this diversity in structure within the MscS family implies 186 that MSL channels in plants may have functions and regulatory mechanisms that are 187 specific to multicellular eukaryotes [53]. A three-dimensional structure of these channels 188 189 would reveal the spatial relationship between the regions thought to serve as tension sensors, the channel pore, and soluble domains. This would help us determine how 190 191 membrane tension is transmitted from the channel-membrane interface to the channel pore—and potentially to other domains within the protein (see non-conducting functions, 192 193 above).

194

195 Closing the Gap between Channel Behavior in the Patch Pipette and in the Intact **Plant Cell**. While patch clamp electrophysiology has proven a powerful way to identify 196 197 and characterize MS ion channels, in plants takes place in the absence of a cell wall, sometimes in an isolated membrane patch, in tightly regulated and non-physiological ionic 198 199 conditions, and in the case of heterologous expression, not in the native lipid environment. Thus, the next great challenge for the field will be developing approaches that allow the 200 analysis of MS ion channel action in their native context. Controlled activation of MS 201 channels from inside a plant cell might be possible through the application of focused 202 ultrasound, as was recently demonstrated for animal TPKs expressed in oocytes [54]. 203 Integration of localized extracellular ion flux measurements with genetically encoded ion 204 or voltage biosensors may allow the study of MS channel function in some cellular 205 contexts, such as pollen tubes [55]. To date, the genetically encoded sensors for 206

transmembrane voltage used extensively in animal systems to monitor ion channel
activity *in vivo* [56] do not yet function well in plants [57].

209

#### 210 Conclusion

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Membrane tension is a force intrinsic to all cells, and every branch of life expresses ion 212 channels that serve specifically to sense and respond to it. In plants, MS ion channels are 213 widely distributed across multiple species, cell types, and intracellular compartments. In 214 Arabidopsis, MS ion channels are required for roots to penetrate hard agar and mediate 215 osmoregulation of pollen and plastids during normal growth and development. Future 216 work should reveal the physiological function of channels we know, add more channel 217 218 genes and proteins to our short list, and develop the methodologies that will allow in vivo analysis of ion channel function, regulation, and mechanism. 219

220

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222

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#### **Box 1: The force-from-lipid principle**

According to the force-from-lipid principle, anisotropic forces inherent to the lipid bilayer impinge on the conformation of membrane-embedded proteins. Ion channels classified as mechanosensitive allow the passage of ions when forces directly transmitted from the lipid bilayer are transduced into conformational rearrangements of the protein. This concept is proposed to underlie the mechanosensitivity of channels from multiple kingdoms and evolutionarily unrelated families. It follows from this principle that all channels are to some degree mechanosensitive; enhanced sensitivity, dynamic range,

- and spatio-temporal control are accomplished through structural arrangement and/or by
- tethering to cytoskeletal elements or extracellular matrix.
- 241

#### 242 Figure Legends

243

Figure 1. Subcellular Localization and Ionic Preference for Known Plant Mechanosensitive Ion Channels.

The subcellular localization of MS ion channel proteins so far identified in land plants is indicated [20-23,32,58]. The outer membrane of the chloroplast is permeable to ions [59], and Voltage-dependent Anion Channels (VDACs) are thought to mediate flux across the outer mitochondrial membrane [60]. MSL, MscS-Like; TPK, Two-pore K<sup>+</sup>; MCA, Mid1-Complementing Activity. Note that only general ion permeability preferences are indicated; these channels are likely to be permeable to additional species.

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Figure 2. Proposed Role of MSL8 in Controlling Turgor During Pollen Hydration,
 Germination, and Tube Growth.

Wild-type pollen grains successfully survive hydration in distilled water, germinate 255 effectively in germination media, produce intact pollen tubes, and are optimally fertile. 256 Pollen grains from *msl8-4* null mutants, or null mutants expressing the  $MSL8F^{720L}$  allele, 257 display reduced viability upon hydration in distilled water, and we propose that this is due 258 to an inability to relieve turgor pressure by releasing ions upon hypoosmotic shock. 259 Excess turgor after hydration leads both to germination at a rate higher than the wild type, 260 261 but also to frequent bursting, and an overall loss of fertility. When MSL8 is overexpressed from the pollen-specific, strong LAT52 promoter, pollen grains survive hydration but are 262 unable to maintain the threshold turgor pressure required for pollen germination or tube 263 elongation. Green arrows, optimal turgor; red arrows, excessive turgor; blue arrows, 264 insufficient turgor. 265

#### 266 **References**

- Anishkin A, Loukin SH, Teng J, Kung C: Feeling the hidden mechanical forces
   in lipid bilayer is an original sense. *Proc. Natl. Acad. Sci. U.S.A.* 2014,
   111:7898–7905.
- The authors of this Perspective argue that the force-from-lipid principle can and should be applied to membrane-embedded proteins in all organisms.
- Ranade SS, Syeda R, Patapoutian A: Mechanically Activated Ion Channels.
   *Neuron* 2015, 87:1162–1179.
- Peyronnet R, Tran D, Girault T, Frachisse J-M: Mechanosensitive channels:
   feeling tension in a world under pressure. *Front. Plant Sci.* 2014, 5:558.
- Martinac B: Mechanosensitive ion channels: an evolutionary and scientific tour de force in mechanobiology. *Channels (Austin)* 2012, 6:211–213.
- Levina N, Tötemeyer S, Stokes NR, Louis P, Jones MA, Booth IR: Protection of
   Escherichia coli cells against extreme turgor by activation of MscS and MscL
   mechanosensitive channels: identification of genes required for MscS
   activity. EMBO J. 1999, 18:1730–1737.
- Yan Z, Zhang W, He Y, Gorczyca D, Xiang Y, Cheng LE, Meltzer S, Jan LY, Jan
   YN: Drosophila NOMPC is a mechanotransduction channel subunit for gentletouch sensation. *Nature* 2013, **493**:221–225.
- Cox CD, Nomura T, Ziegler CS, Campbell AK, Wann KT, Martinac B: Selectivity
   mechanism of the mechanosensitive channel MscS revealed by probing
   channel subconducting states. *Nature Communications* 2013, 4:2137.
- Falke LC, Edwards KL, Pickard BG, Misler S: A stretch-activated anion channel
   in tobacco protoplasts. *FEBS Lett.* 1988, 237:141–144.
- Schroeder JI, Hedrich R: Involvement of ion channels and active transport in
   osmoregulation and signaling of higher plant cells. *Trends in Biochemical Sciences* 1989, 14:187–192.
- 10. Morris CE: Mechanosensitive ion channels. J. Membr. Biol. 1990, **113**:93–107.
- Hamilton ES, Schlegel AM, Haswell ES: United in diversity: mechanosensitive
   ion channels in plants. Annu. Rev. Plant Biol. 2015, 66:113–137.
- Yuan F, Yang H, Xue Y, Kong D, Ye R, Li C, Zhang J, Theprungsirikul L, Shrift T,
   Krichilsky B, et al.: OSCA1 mediates osmotic-stress-evoked Ca2+ increases
   vital for osmosensing in Arabidopsis. Nature 2014, 514:367–371.
- 13. Hou C, Tian W, Kleist T, He K, Garcia V, Bai F, Hao Y, Luan S, Li L: DUF221

- proteins are a family of osmosensitive calcium-permeable cation channels
   conserved across eukaryotes. *Cell Res.* 2014, **24**:632–635.
- 14. Coste B, Mathur J, Schmidt M, Earley TJ, Ranade S, Petrus MJ, Dubin AE,
   Patapoutian A: Piezo1 and Piezo2 are essential components of distinct
   mechanically activated cation channels. *Science* 2010, 330:55–60.
- Pivetti CD, Yen M-R, Miller S, Busch W, Tseng Y-H, Booth IR, Saier MH: Two
   families of mechanosensitive channel proteins. *Microbiol. Mol. Biol. Rev.* 2003,
   67:66–85– table of contents.
- Silvia GHUL de S, Adriana PDS, Tania MI, Eduardo GDS, Talita CU: Genome wide analysis of mechanosensitive channel of small conductance (MscS)-like
   gene family in common bean. *Afr. J. Biotechnol.* 2016, **15**:580–592.
- Porter BW, Zhu YJ, Webb DT, Christopher DA: Novel thigmomorphogenetic
   responses in Carica papaya: touch decreases anthocyanin levels and
   stimulates petiole cork outgrowths. *Ann. Bot.* 2009, 103:847–858.
- 18. Saddhe AA, Kumar K: In silico identification and expression analysis of MscS
   like gene family in rice. *Plant Gene* 2015, 1:8–17.
- Haswell ES: MscS-Like Proteins in Plants. In *Mechanosensitive Ion Channels, Part A.* Elsevier; 2007:329–359.
- Haswell ES, Meyerowitz EM: MscS-like proteins control plastid size and shape
   in Arabidopsis thaliana. *Curr. Biol.* 2006, 16:1–11.
- Lee CP, Maksaev G, Jensen GS, Murcha MW, Wilson ME, Fricker M, Hell R, Haswell ES, Millar AH, Sweetlove L: MSL1 is a mechanosensitive ion channel that dissipates mitochondrial membrane potential and maintains redox homeostasis in mitochondria during abiotic stress. *Plant J* 2016, doi:10.1111/tpj.13301.
- Haswell ES, Peyronnet R, Barbier-Brygoo H, Meyerowitz EM, Frachisse J-M: Two
   MscS homologs provide mechanosensitive channel activities in the
   Arabidopsis root. Curr. Biol. 2008, 18:730–734.
- Hamilton ES, Jensen GS, Maksaev G, Katims A, Sherp AM, Haswell ES:
   Mechanosensitive channel MSL8 regulates osmotic forces during pollen
   hydration and germination. *Science* 2015, 350:438–441.
- MSL8 is shown to be a pollen-specific, membrane tension-gated, ion channel required for pollen to survive the hypoosmotic shock of rehydration and for full male fertility. MSL8 negatively regulates pollen germination, but is required for cellular integrity during germination and tube growth. These data suggest that homologs of bacterial MscS have been repurposed in eukaryotes to sense and respond to mechanical stimuli in a developmental context.

Maksaev G, Haswell ES: MscS-Like10 is a stretch-activated ion channel from
 Arabidopsis thaliana with a preference for anions. *Proc. Natl. Acad. Sci. U.S.A.* 2012, 109:19015–19020.

- This study established that MSL10 is a *bona fide* mechanosensitive ion channel using
   single-channel patch clamp electrophysiology and a heterologous expression
   system, and characterized its behavior. Thus, plant genomes do indeed encode
   mechanosensitive ion channels evolutionarily related to those from bacteria.
- Wilson ME, Jensen GS, Haswell ES: Two mechanosensitive channel homologs
   influence division ring placement in Arabidopsis chloroplasts. *Plant Cell* 2011,
   23:2939–2949.
- Veley KM, Marshburn S, Clure CE, Haswell ES: Mechanosensitive channels
   protect plastids from hypoosmotic stress during normal plant growth. *Curr. Biol.* 2012, 22:408–413.
- Wilson ME, Basu MR, Bhaskara GB, Verslues PE, Haswell ES: Plastid osmotic
   stress activates cellular stress responses in Arabidopsis. *Plant Physiol.* 2014,
   165:119–128.
- Wilson ME, Mixdorf M, Berg RH, Haswell ES: Plastid osmotic stress influences
   cell differentiation at the plant shoot apex. *Development* 2016, 143:3382–3393.
- Brohawn SG: How ion channels sense mechanical force: insights from
   mechanosensitive K2P channels TRAAK, TREK1, and TREK2. Ann. N. Y. Acad.
   Sci. 2015, 1352:20–32.
- 358 30. Voelker C, Gomez-Porras JL, Becker D, Hamamoto S, Uozumi N, Gambale F,
   359 Mueller-Roeber B, Czempinski K, Dreyer I: Roles of tandem-pore K+ channels
   360 in plants a puzzle still to be solved\*. *Plant Biology* 2010, 12:56–63.
- 361 31. Maathuis FJM: Vacuolar two-pore K+ channels act as vacuolar osmosensors.
   362 New Phytol 2011, 191:84–91.
- 363 32. Nakagawa Y, Katagiri T, Shinozaki K, Qi Z, Tatsumi H, Furuichi T, Kishigami A,
   364 Sokabe M, Kojima I, Sato S, et al.: Arabidopsis plasma membrane protein
   365 crucial for Ca2+ influx and touch sensing in roots. Proc. Natl. Acad. Sci. U.S.A.
   366 2007, 104:3639–3644.
- The authors employed a novel functional screen in yeast to identify MCA1, a plant specific membrane protein that facilitates Ca<sup>2+</sup> influx in response to mechanical and
   osmotic stress. Seedlings lacking functional MCA1 are unable to efficiently penetrate hard
   agar. MCA1 and close homolog MCA2 are leading candidates for the long-sought plant
   stretch-activated Ca<sup>2+</sup> channel.
- 372

- 373 33. Kamano S, Kume S, Iida K, Lei K-J, Nakano M, Nakayama Y, Iida H:
  374 Transmembrane Topologies of Ca2+-permeable Mechanosensitive Channels
  375 MCA1 and MCA2 in Arabidopsis thaliana. J. Biol. Chem. 2015, 290:30901–
  376 30909.
- 377 34. Shigematsu H, Iida K, Nakano M, Chaudhuri P, Iida H, Nagayama K: Structural
   378 characterization of the mechanosensitive channel candidate MCA2 from
   379 Arabidopsis thaliana. *PLoS ONE* 2014, **9**:e87724.
- 380 35. Furuichi T, lida H, Sokabe M, Tatsumi H: Expression of Arabidopsis MCA1
   anhanced mechanosensitive channel activity in the Xenopus laevis oocyte
   plasma membrane. *Plant Signal Behav* 2012, **7**:1022–1026.
- 383 36. Kurusu T, Yamanaka T, Nakano M, Takiguchi A, Ogasawara Y, Hayashi T, Iida K, 384 Hanamata S, Shinozaki K, Iida H, et al.: Involvement of the putative Ca<sup>2+</sup>-385 permeable mechanosensitive channels, NtMCA1 and NtMCA2, in Ca<sup>2+</sup> uptake, 386 Ca<sup>2+</sup>-dependent cell proliferation and mechanical stress-induced gene 387 expression in tobacco (Nicotiana tabacum) BY-2 cells. *J. Plant Res.* 2012, 388 125:555–568.
- 389 37. Kurusu T, Nishikawa D, Yamazaki Y, Gotoh M, Nakano M, Hamada H, Yamanaka
   390 T, Iida K, Nakagawa Y, Saji H, et al.: Plasma membrane protein OsMCA1 is
   391 involved in regulation of hypo-osmotic shock-induced Ca2+ influx and
   392 modulates generation of reactive oxygen species in cultured rice cells. *BMC* 393 *Plant Biol.* 2012, **12**:11.
- 38. Yamanaka T, Nakagawa Y, Mori K, Nakano M, Imamura T, Kataoka H, Terashima
  A, Iida K, Kojima I, Katagiri T, et al.: MCA1 and MCA2 that mediate Ca2+ uptake
  have distinct and overlapping roles in Arabidopsis. *Plant Physiol.* 2010,
  152:1284–1296.
- 398 39. Rosa M, Abraham-Juárez MJ, Lewis MW, Fonseca JP, Tian W, Ramirez V, Luan
  399 S, Pauly M, Hake S: The Maize MID-COMPLEMENTING ACTIVITY Homolog
  400 CELL NUMBER REGULATOR13/NARROW ODD DWARF Coordinates Organ
  401 Growth and Tissue Patterning. THE PLANT CELL ONLINE 2017, 29:474–490.
- 40. Hamilton ES, Haswell ES: The Tension-sensitive Ion Transport Activity of
   403 MSL8 is Critical for its Function in Pollen Hydration and Germination. Plant
   404 and Cell Physiology 2017, doi:10.1093/pcp/pcw230.
- Point mutations were introduced into the presumptive channel pore of MSL8 and
   resulting variants assessed for channel behavior and function in pollen hydration,
   germination, and tube growth. Lesions in the pore that change channel behavior also
   change physiological function, and it is concluded that MSL8 serves as a
   mechanoreceptor in pollen.

- 411 41. Ghosh R, Gururani MA, Ponpandian LN, Mishra RC, Park S-C, Jeong M-J, Bae H:
  412 Expression Analysis of Sound Vibration-Regulated Genes by Touch
  413 Treatment in Arabidopsis. Front. Plant Sci. 2017, 8:100.
- 414 42. Damiani I, Drain A, Guichard M, Balzergue S, Boscari A, Boyer J-C, Brunaud V,
  415 Cottaz S, Rancurel C, Da Rocha M, et al.: Nod Factor Effects on Root Hair416 Specific Transcriptome of Medicago truncatula: Focus on Plasma Membrane
  417 Transport Systems and Reactive Oxygen Species Networks. Front. Plant Sci.
  418 2016, 7:3389–22.
- 43. Veley KM, Maksaev G, Frick EM, January E, Kloepper SC, Haswell ES:
  Arabidopsis MSL10 has a regulated cell death signaling activity that is
  separable from its mechanosensitive ion channel activity. *Plant Cell* 2014,
  26:3115–3131.

This study used molecular genetics, a transient expression assay, and
 electrophysiology to show that MSL10 has two genetically separable functions, each
 attributable to a different domain of the protein. These results implicate MSL10 in ROS mediated cell death, and provide evidence that some MscS homologs have non conducting functions.

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 44. Zou Y, Chintamanani S, He P, Fukushige H, Yu L, Shao M, Zhu L, Hildebrand DF,
 Tang X, Zhou J-M: A gain-of-function mutation in MsI10 triggers cell death and
 wound-induced hyperaccumulation of jasmonic acid in Arabidopsis. J Integr
 431
 Plant Biol 2016, 58:600–609.

432 45. Zhang Z, Tateda C, Jiang S-C, Shrestha J, Jelenska J, Speed DJ, Greenberg JT:
433 A Suite Of Receptor-Like Kinases and a Putative Mechano-Sensitive Channel
434 Are Involved in Autoimmunity and Plasma Membrane–Based Defenses in
435 Arabidopsis. *Mol. Plant Microbe Interact.* 2017, doi:10.1094/MPMI-09-16-0184-R.

- The authors identify MSL4 as one of several proteins that interact with ACD6, a plasma membrane-localized defense protein. *msl4* mutants show multiple defense-related defects, including reduced resistance to *Pseudomonas syringae pv. maculicola ES4326 hrcC-*.
- 440 46. Kohorn BD, Hoon D, Minkoff BB, Sussman MR, Kohorn SL: Rapid Oligo441 Galacturonide Induced Changes in Protein Phosphorylation in Arabidopsis.
  442 Mol. Cell Proteomics 2016, 15:1351–1359.
- 443 47. Di Salle P, Incerti G, Colantuono C, Chiusano ML: Gene co-expression analyses:
  444 an overview from microarray collections in Arabidopsis thaliana. Brief
  445 Bioinform 2016, doi:10.1093/bib/bbw002.
- 446 48. Bavi N, Cox CD, Perozo E, Martinac B: Towards a structural blueprint for
  447 bilayer-mediated channel mechanosensitivity. *Channels (Austin)* 2016,
  448 doi:10.1080/19336950.2016.1224624.

- 449 49. Pliotas C, Naismith JH: **Spectator no more, the role of the membrane in** 450 **regulating ion channel function.** *Curr. Opin. Struct. Biol.* 2016, **45**:59–66.
- 451 50. Brohawn SG, Campbell EB, MacKinnon R: Physical mechanism for gating and
  452 mechanosensitivity of the human TRAAK K+ channel. *Nature* 2014, 516:126–
  453 130.
- 454 51. Aryal P, Jarerattanachat V, Clausen MV, Schewe M, McClenaghan C, Argent L,
  455 Conrad LJ, Dong YY, Pike ACW, Carpenter EP, et al.: Bilayer-Mediated
  456 Structural Transitions Control Mechanosensitivity of the TREK-2 K2P
  457 Channel. Structure 2017, doi:10.1016/j.str.2017.03.006.
- 458 52. Cox CD, Nakayama Y, Nomura T, Martinac B: The evolutionary "tinkering" of
   459 MscS-like channels: generation of structural and functional diversity. *Pflugers* 460 Arch. 2015, 467:3–13.
- 461 53. Haswell ES, Phillips R, Rees DC: Mechanosensitive channels: what can they
  462 do and how do they do it? *Structure* 2011, **19**:1356–1369.
- 463 54. Kubanek J, Shi J, Marsh J, Chen D, Deng C, Cui J: **Ultrasound modulates ion** 464 **channel currents.** *Nature Publishing Group* 2016, **6**:24170.
- 55. Damineli DSC, Portes M-T, Feijó JA: Oscillatory signatures underlie growth
   regimes in Arabidopsis pollen tubes: computational methods to estimate tip
   location, periodicity, and synchronization in growing cells. Journal of
   *Experimental Botany* 2017, doi:10.1093/jxb/erx032.
- 469 56. St-Pierre F, Chavarha M, Lin MZ: Designs and sensing mechanisms of
  470 genetically encoded fluorescent voltage indicators. *Curr Opin Chem Biol* 2015,
  471 27:31–38.
- 472 57. Matzke AJM, Matzke M: Expression and testing in plants of ArcLight, a
   473 genetically-encoded voltage indicator used in neuroscience research. BMC
   474 Plant Biol. 2015, 15:245.
- 475 58. Czempinski K, Frachisse J-M, Maurel C, Barbier-Brygoo H, Mueller-Roeber B:
  476 Vacuolar membrane localization of the Arabidopsis "two-pore" K+ channel
  477 KC01. Plant J 2002, 29:809–820.
- 478 59. Heldt HW, Sauer F: **The inner membrane of the chloroplast envelope as the** 479 **site of specific metabolite transport.** *Biochim. Biophys. Acta* 1971, **234**:83–91.
- 480 60. Homblé F, Krammer E-M, Prévost M: Plant VDAC: facts and speculations.
  481 Biochim. Biophys. Acta 2012, 1818:1486–1501.

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## Fertility

### Optimal



## Compromised at germination stage