



A mechanosensitive Ca(2+) channel activity is dependent on the developmental regulator DEK1

Daniel Tran, Roberta Galletti, Enrique D. Neumann, Annick Dubois, Reza Sharif-Naeini, Anja Geitmann, Jean-Marie Frachisse, Olivier Hamant, Gwyneth Ingram

► To cite this version:

Daniel Tran, Roberta Galletti, Enrique D. Neumann, Annick Dubois, Reza Sharif-Naeini, et al.. A mechanosensitive Ca(2+) channel activity is dependent on the developmental regulator DEK1. *Nature Communications*, 2017, 8 (1), pp.1009. 10.1038/s41467-017-00878-w . hal-02392032

HAL Id: hal-02392032

<https://hal.science/hal-02392032>

Submitted on 26 May 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License

ARTICLE

DOI: 10.1038/s41467-017-00878-w

OPEN

A mechanosensitive Ca^{2+} channel activity is dependent on the developmental regulator DEK1

Daniel Tran^{1,2,3}, Roberta Galletti¹, Enrique D. Neumann¹, Annick Dubois¹, Reza Sharif-Naeini³, Anja Geitmann⁴, Jean-Marie Frachisse², Olivier Hamant¹ & Gwyneth C. Ingram¹

Responses of cells to mechanical stress are thought to be critical in coordinating growth and development. Consistent with this idea, mechanically activated channels play important roles in animal development. For example, the PIEZO1 channel controls cell division and epithelial-layer integrity and is necessary for vascular development in mammals. In plants, the actual contribution of mechanoperception to development remains questionable because very few putative mechanosensors have been identified and the phenotypes of the corresponding mutants are rather mild. Here, we show that the *Arabidopsis* Defective Kernel 1 (DEK1) protein, which is essential for development beyond early embryogenesis, is associated with a mechanically activated Ca^{2+} current *in planta*, suggesting that perception of mechanical stress plays a critical role in plant development.

¹Laboratoire Reproduction et Développement des Plantes, Université de Lyon, ENS de Lyon, UCB Lyon 1, CNRS, INRA, F-69342 Lyon, France. ²Institute for Integrative Biology of the Cell (I2BC), CEA, CNRS, Université Paris-Sud, Sciences Plant Saclay, Avenue de la Terrasse, 91198 Gif sur Yvette Cedex, France. ³Department of Physiology and Cell Information Systems, McGill University, Montreal, Québec, Canada H3G-0B1. ⁴Department of Plant Science, McGill University, Ste-Anne-de-Bellevue, Montreal, Québec, Canada H9X3V9. Daniel Tran and Roberta Galletti contributed equally to this work. Correspondence and requests for materials should be addressed to O.H. (email: Olivier.Hamant@ens-lyon.fr) or to G.C.I. (email: Gwyneth.Ingram@ENS-lyon.fr)

Multicellular development is dependent upon cell–cell communication. Recent research has highlighted the fact that, in addition to chemical signals, the perception of mechanical stress, at both the cell and tissue level, is a key factor underlying growth coordination and morphogenesis (e.g., ref. ¹). However, while in animal systems mechanosensors with important roles in development have been identified^{2–9}, the molecular basis for mechanoperception in plants remains enigmatic. As in animals, plants respond to mechanical stimuli by an elevation in cytoplasmic Ca^{2+} ^{10–12}. Based on the knowledge of plasma-membrane mechanosensing in animal systems, the most probable trigger for Ca^{2+} release from internal cellular compartments (Ca^{2+} -induced calcium release) is either the opening of plasma membrane-localized mechanosensitive Ca^{2+} permeable channels, or the opening of voltage-dependent Ca^{2+} channels in response to changes in membrane potential caused by mechanosensitive channels permeable to other ions¹². Recent research in *Arabidopsis* has led to the identification of several plasma membrane-localized mechanosensitive ion channels. These include proteins similar to the bacterial mechanosensitive channel of small conductance (the MSL family^{13–16}), the MCA1 protein that rescues the yeast Ca^{2+} channel mutant *mid1*, and its homolog MCA2^{17–20}. In addition, the membrane protein OSCA1 forms a hyperosmolarity-gated Ca^{2+} permeable channel required for osmosensing in *Arabidopsis*²¹. However, although the MSL8 protein has recently been shown to be required for pollen grains

to survive rapid rehydration during fertilization¹⁴, the very mild developmental phenotypes in single and multiple mutants of genes encoding the channels described above, suggests that they are unlikely to play a major role in mechanosensing during development. Furthermore, these proteins have not conclusively been shown to be responsible for any of the mechanosensitive Ca^{2+} currents which have been detected and extensively described by electrophysiologists over the past few decades *in planta*^{20, 22–26}. Since genes encoding voltage-sensitive Ca^{2+} channels similar to those identified in animal systems have not been found in plant genomes, it has been proposed that plants may have evolved novel systems for mediating mechanosensitive Ca^{2+} fluxes at the plasma membrane to control development (reviewed in ref. ¹²).

The Defective Kernel 1 (DEK1) protein is encoded by a highly conserved uni-gene found in all multicellular plant genomes sequenced so far, and it is absolutely required for both embryonic and post-embryonic development in angiosperms^{27–31}. Null *dek1* mutant embryos do not develop beyond the early globular stage. When plants with reduced DEK1 activity can be obtained, they show major developmental defects, notably in epidermal differentiation and adhesion^{30, 32, 33}.

The DEK1 protein contains multiple predicted transmembrane (TM) spans interrupted by a loop, and an intracellular tail including a linker domain and a C-terminal domain showing similarity to animal calpains, a class of Ca^{2+} -dependent cysteine

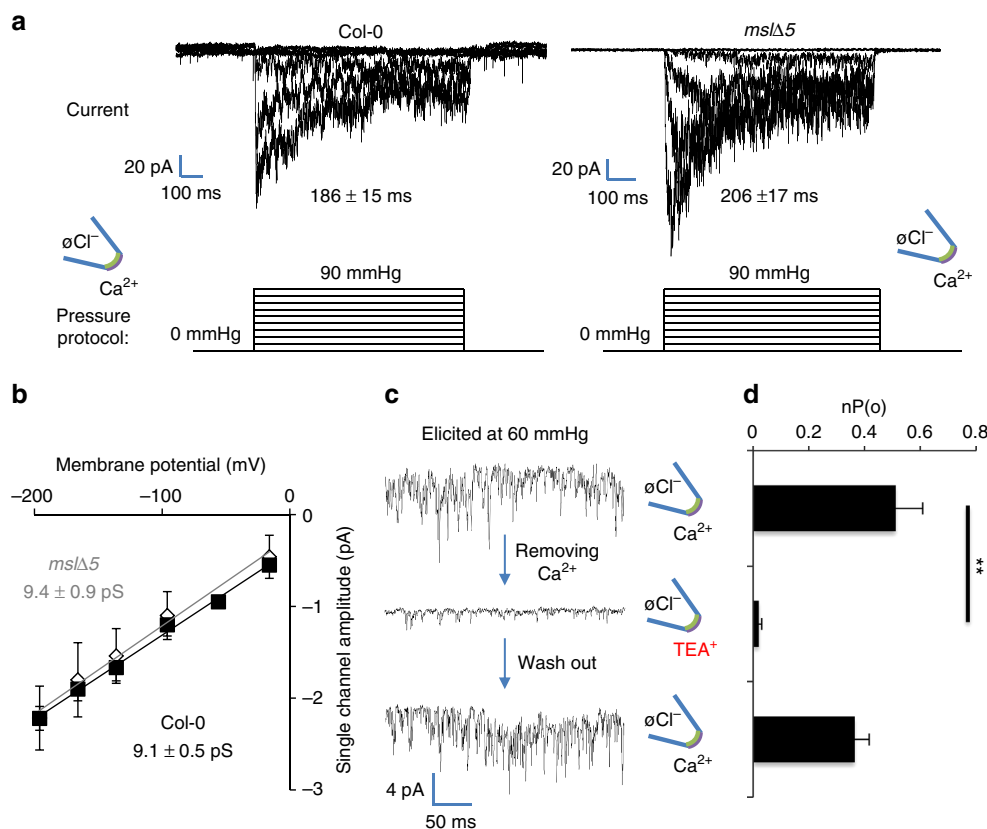


Fig. 1 A mechanically activated current permeable to Ca^{2+} is present at the plasma membrane. **a** Representative membrane patches from Col-0 (left) and an *msl* quintuple mutant *msl4;msl5;msl6;msl9;msl10* (*mslΔ5*, right), exposed to increased positive pressure steps in an outside-out patch configuration show a rapidly activated, rapidly inactivated current in ionic conditions favorable for Ca^{2+} current recording. We have named the channel responsible for this activity the RMA channel (see text). Time constants are means $\pm$ SE ($n = 6$); **b** Under the same conditions, single channel I/V curves show similar RMA channel conductance in Col-0 (solid square) and in the *mslΔ5* mutant (open square). Values are means $\pm$ SE ($n = 6$); **c** Representative single channel recordings showing that the RMA current is reversibly eliminated by exchanging Ca^{2+} ions with non-permeant TEA^+ ions. **d** Open probability (for n channels, $n\text{Po}$) is severely decreased in TEA^+ bath solution. A paired *t*-test was used to compare means (** $P < 0.01$). Values are means $\pm$ SE. For all experiments, the membrane potential was held at -196 mV. Ionic conditions are described in the Methods section

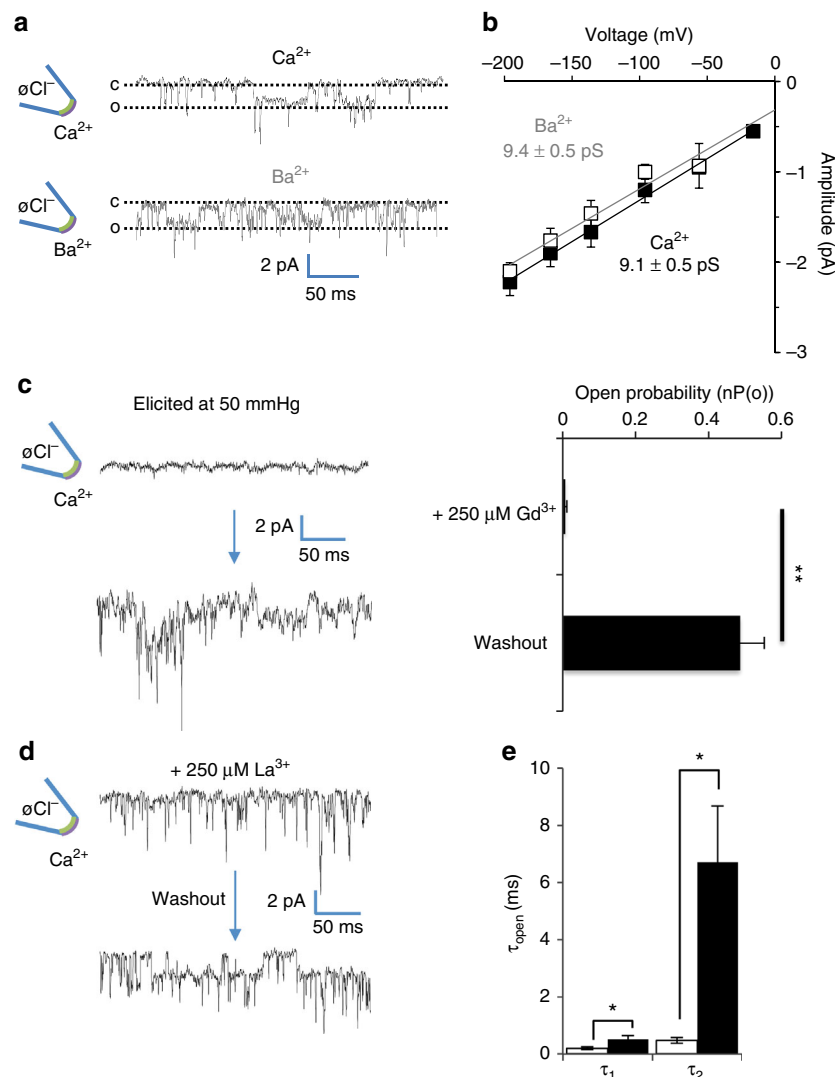


Fig. 2 The mechanically activated RMA current is preferentially permeable to Ca^{2+} and affected by both the mechanosensitive channel inhibitor Gd^{3+} and the calcium channel blocker La^{3+} . **a** Representative single channel recordings in response to positive pressure, in an outside-out patch configuration with Ca^{2+} (top) or Ba^{2+} (bottom) as the permeant bivalent cation in the bath. The dotted lines indicate the open (O) and closed (C) channel state; **b** Single channel I/V curves show similar conductance with permeant cations Ca^{2+} (solid square) and Ba^{2+} (open square). Values are means $\pm$ SE ($n = 6$); **c** The mechanically activated RMA current is inhibited by Gd^{3+} and restored after wash-out (left, $n = 6$) and the corresponding open probability (nP(o) , right). Values are means $\pm$ SE; **d** Representative recording showing the effect of La^{3+} (left, $n = 6$) on the RMA current; **e** La^{3+} ions affects the open state of the RMA channel and reducing the mean open time. Bars represent the two time constant τ_1 and τ_2 of the open state with (white bars) or without (black bars) La^{3+} . A paired t -test was used to compare means ($P < 0.01$). $E_{\text{rev}}(\text{Ca}^{2+}) = 47.5 \pm 5.4$ mV; $E_{\text{rev}}(\text{Ba}^{2+}) = 36.5 \pm 6.1$ mV. Values are means $\pm$ SE (open events, $n \geq 100$). For all experiments, the membrane potential was held at -196 mV. Ionic conditions are described in the Methods section

proteases³⁴. DEK1 localizes to the plasma membrane and to internal compartments^{32,34}. In their original model of DEK1, Lid et al.³⁴ predicted the presence of 21 TM domains with an extracellular localization for the loop and cytoplasmic localization (subsequently confirmed *in planta*) for the C-terminus. However, in a more recent analysis, Kumar et al.³⁵ proposed a consensus model with 23 TM domains and with a cytoplasmic localization for the loop. Structure-function studies have shown that the cytosolic CALPAIN domain of the protein can, alone, complement the embryo lethality of *dek1* mutants suggesting that this domain, which is removed from the rest of the protein by an autolytic cleavage event, represents an active form of the DEK1 protein³². Consistent with this scenario, and with a role for DEK1 in maintaining epidermal integrity, overexpression of the CALPAIN domain of DEK1 leads to thickening of the outer epidermal cell wall in leaves, and increased deposition of pectins³⁶, which

are important for cell adhesion (reviewed in ref. ³⁷). The epidermis is thought to be under tension during much of plant growth³⁸. One plausible function of DEK1 could, therefore, be to perceive and respond to this tension to coordinate epidermal development and maintain epidermal integrity. As the epidermis plays a critical role in organ growth in plants (e.g., refs. ^{39–42}), this, in turn, has major implications for plant development as a whole.

The activity of the CALPAIN domain has been shown to be Ca^{2+} -dependent *in vitro*⁴³. These findings are in line with results in animal systems where the activation of cytoplasmic calpains is associated with the Ca^{2+} -dependent autolytic-removal of an N-terminal extension, which in some cases can inhibit calpain activity^{44–46}. Because calpains have been shown to act downstream of mechanosensitive channels such as PIEZO in animals⁴, we tested the hypothesis, and demonstrated, that the TM domain

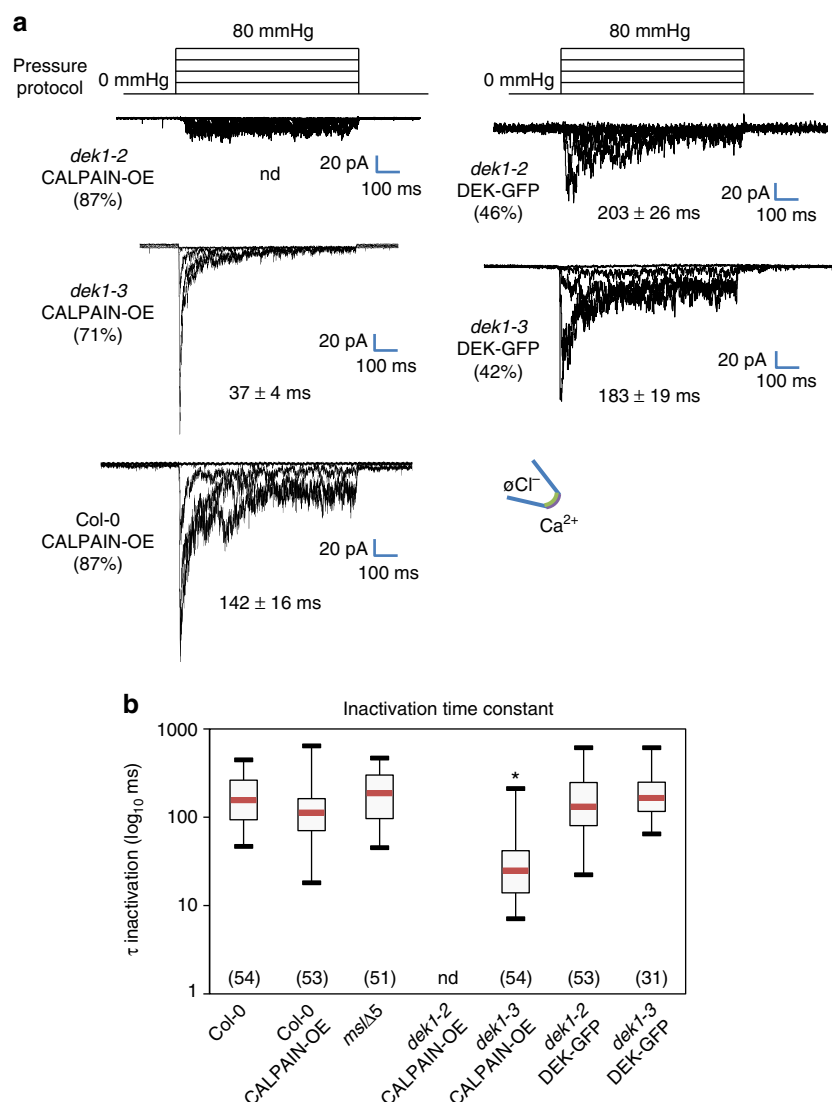


Fig. 3 RMA activation depends on the trans-membrane region of the essential DEK1 protein. **a** Mechanosensitive current kinetics in response to pulse pressure (as illustrated) in *dek1-2* CALPAIN-OE, *dek1-3* CALPAIN-OE, Col-0 CALPAIN-OE, *dek1-2* DEK-GFP and *dek1-3* DEK-GFP. Time constants are means $\pm$ SE ($n = 6$); **b** Inactivation times are fitted with an exponential function, and time constants (τ_{inact}) are represented in a box plot distribution for different lines. Number in brackets represent the number of time constants. *significantly different from Col-0 line, Anova on ranks ($P < 0.05$). For all experiments the membrane potential was held at -196 mV. Ionic conditions are described in the Methods section

of DEK1 is associated with Ca^{2+} dependent mechanoperception at the plasma membrane in plants.

Results

A rapidly mechanically activated plasma membrane Ca^{2+} channel. Combining genetic and electrophysiological approaches, the *Arabidopsis* MSL9 and MSL10 proteins have been shown to mediate a mechanically activated channel activity in protoplasts derived from wild-type plant root cells¹³. Selectivity characterization indicated that anions permeate preferentially through the wild-type MSL channels¹³. Because in this work we used excised membrane patches from callus protoplasts, rather than whole root cell-derived protoplasts, we first investigated whether mechanically activated ion currents due to MSLs and other unidentified channels could also be detected in this material.

Reverse transcription quantitative-PCR (RT-qPCR) analysis confirmed the expression of *MSL9* and *MSL10* in wild-type

(Columbia-0) callus tissue (Supplementary Fig. 1a). When Cl^- is provided at the cytosolic face of plasma membrane patches, a mechanically activated channel with a high conductance (46 ± 1.4 pS; $\pm$ indicates standard deviation, $n = 6$) is elicited upon application of positive pressure using a high-speed pressure clamp, as previously described in root protoplasts (Supplementary Fig. 1b)¹³. This activity, which is mainly due to MSL9 and MSL10, was completely abolished in protoplasts obtained from quintuple *msl4 msl5 msl6 msl9 msl10* (*msl5*) mutant callus (Supplementary Fig. 1c). In addition, to MSL activity, we observed a rapidly activated-inactivated current with a smaller conductance, elicited immediately after the pressure increase (Supplementary Fig. 1b). Unlike the situation for the MSL current, removing Cl^- from the cytosolic face of the membrane did not modify this rapid current (Fig. 1a, Supplementary Fig. 1b). The rapid current was still present in protoplasts derived from *msl5* mutant callus with or without Cl^- ions at the cytosolic face (Fig. 1a, Supplementary Fig. 1c). In addition this

mechanically activated channel showed a similar conductance to that seen in Col-0 callus (Fig. 1b), with a reversal potential at positive voltage ($E_{rev}^{WT} = 47.5 \pm 5.4$ mV and $E_{rev}^{msl/\Delta 5} = 60.6 \pm 11.5$ mV; $\pm$ indicates standard deviation, $n = 6$) suggesting that this rapid current is most likely mediated by Ca^{2+} influx (dominant cation gradient, $E_{Ca^{2+}} = 128.7$ mV). Consistent with this hypothesis, we found that it could be reversibly eliminated by replacing Ca^{2+} in the bath solution with the larger TEA^{+} cation (Fig. 1c, d). The pressure required to activate the current was higher than that required to activate MSL channels, suggesting distinct activation thresholds for these channels (Supplementary Fig. 1b, c). Thus, we can distinguish at least two independent mechanically activated channel activities in callus protoplasts: one dependent on MSLs activity and the second due to an unidentified Ca^{2+} channel which we called the rapid mechanically activated (RMA) channel.

To understand the nature of the RMA-mediated current we tested a range of pipette and bath solutions. The RMA channel conductance is unaffected when Ca^{2+} ions are replaced with Ba^{2+} ions in the bath solution (Fig. 2a, b) suggesting that the RMA channel is similarly permeant to divalent Ca^{2+} and Ba^{2+} ions, as is classically found for Ca^{2+} channels. To test whether the RMA channel shows characteristics previously described for mechanically activated channels in plants, we investigated the effect of Gd^{3+} ions, which are known to affect mechanically activated channel activity in both animal and plant systems^{23, 47–52}. RMA activity was almost totally blocked by the presence of Gd^{3+} ions, and activity could subsequently be restored by washing out Gd^{3+} (Fig. 2c). The presence of La^{3+} ions, a known blocker of Ca^{2+} channels^{53–55} strongly reduced the mean open time of the RMA channel (Fig. 2d, e) but did not modify the channel conductance. We conclude that the RMA channel activity fulfils the key criteria previously attributed to mechanically sensitive Ca^{2+} channel activities recorded at the plant plasma membrane.

DEK1 trans-membrane domains are required for RMA activation. Loss of DEK1 activity in *Arabidopsis* leads to early embryo lethality (Supplementary Fig. 2a)^{29, 30}. By expressing the cytoplasmically localized CALPAIN domain under a constitutive promoter (the *RPS5A* promoter) in *dek1* mutant alleles³², we could restore post-embryonic growth and thus investigate the contribution of the DEK1 TM domain to RMA activation. This was done in both the *dek1-2* mutant background (which has a T-DNA insertion toward the beginning of the TM encoding regions (Supplementary Fig. 2b, c)³²) and in the *dek1-3* mutant background (which contains a T-DNA downstream of the TM encoding regions of the *DEK1* gene (Supplementary Fig. 2b, c)³²). Calli were generated from both *dek1-2* CALPAIN-OE and *dek1-3* CALPAIN-OE lines and genotyped (Supplementary Fig. 3a–c). In callus from CALPAIN complemented *dek1-3* mutants (*dek1-3* CALPAIN-OE), we detected wild-type levels of the transcript corresponding to the endogenous TM-span encoding region of *DEK1* (Supplementary Figs. 2c, 4a, b). This allele therefore has the potential to encode a truncated version of the DEK1 protein, with an intact set of TM spans. However, in callus from complemented *dek1-2* mutants (*dek1-2* CALPAIN-OE), the TM domain-encoding region of the *DEK1* gene is physically disrupted (Supplementary Figs. 2c, 4a, c) and an aberrant transcript is produced downstream of the T-DNA insertion (Supplementary Fig. 4b). This transcript is unlikely to be translated to produce a functional truncated protein as the *dek1-2* allele shows a null phenotype. Overexpression of the CALPAIN transcript was detected in both backgrounds (Supplementary Fig. 4b). We also generated an antibody against the CALPAIN domain and verified overexpression at the protein level (Supplementary Figs. 5a, b, 6).

Because the *dek1-2* CALPAIN-OE background cannot produce the intact N-terminal region of DEK1, we next tested the potential effect of DEK1 TM domain disruption on the RMA channel activity. In the majority of responsive patches obtained from *dek1-2* CALPAIN-OE calli we observed a complete absence of the RMA current upon membrane stretching (Fig. 3a, Supplementary Fig. 7). Only a residual current was detectable in most patches of this background. In patches from *dek1-3* CALPAIN-OE calli the RMA channel activity was still detectable, but showed 6–8 times more rapid inactivation kinetics (Fig. 3a, b, Supplementary Fig. 7) compared to those observed in callus from Col-0 plants.

The production of viable *dek1* mutant plants, as mentioned before, requires the expression of at least the CALPAIN domain of DEK1. In our lines, the CALPAIN domain is overexpressed in the callus system at the RNA level (Supplementary Fig. 4). To confirm that the loss of the RMA channel activity in the *dek1-2* CALPAIN-OE background is not a consequence of CALPAIN overexpression, we generated calli from Col-0 plants overexpressing the CALPAIN domain (Supplementary Figs. 4d and 5)¹⁸. This material was submitted to electrophysiological analysis. The RMA current was still present in patches from this material, and behaved similarly to the current in Col-0 patches. (Fig. 3a, b, Supplementary Fig. 7). Therefore, we conclude that the CALPAIN domain overexpression *per se* is not responsible for the loss of RMA channel activity in the *dek1-2* CALPAIN-OE background.

Loss of the DEK1 TM domains does not affect MSL activity. Next, we investigated whether the removal of the DEK1 TM domains could cause a perturbation of other mechanosensitive channel activities at the plasma membrane by testing whether MSL channels can be activated in the *dek1-2* CALPAIN-OE line. We found that the MSL activity in *dek1-2* CALPAIN-OE protoplasts was still detectable upon membrane stretching, and its characteristics were comparable to that of Col-0 (Supplementary Fig. 8). Thus, loss of the DEK1 TM domains leads to a specific perturbation of the mechanosensitive RMA channel activity in our system.

DEK1 function influences root growth inhibition by Gd^{3+} ions. Based on the mild developmental defects observed in the *dek1* CALPAIN-OE lines^{32, 56}, the CALPAIN domain of DEK1 is likely the “active” domain of the protein. The DEK1 CALPAIN domain is autolytically cleaved from the full length DEK1 protein³², and its activity has been shown to be Ca^{2+} dependent *in vitro*⁴³. Because the release of the CALPAIN domain upon DEK1 cleavage could be triggered by Ca^{2+} influx via the RMA channel, blocking the RMA channel might affect development in wild-type plants, but would have a reduced impact in the *dek1-2* CALPAIN-OE line, where the expression of a “pre-cleaved” version of the DEK1 CALPAIN would partially bypass the need for the RMA channel activity. Considering the sensitivity of the RMA current to Gd^{3+} , and the absence of this current in CALPAIN complemented *dek1-2* mutants we tested whether CALPAIN complemented *dek1-2* mutants might show such reduced sensitivity to Gd^{3+} . Gd^{3+} dramatically inhibits *Arabidopsis* root growth⁵⁷. We generated a dose sensitivity curve of wild-type Col-0 seedlings to concentrations of Gd^{3+} between 0 and 200 μ M (Supplementary Fig. 9). We found strong growth inhibition (40–50%) in wild-type seedlings even at the lowest concentrations tested (30 and 60 μ M) (Supplementary Fig. 9). This inhibition was alleviated in CALPAIN complemented *dek1-2* plants (Supplementary Fig. 10), formally relating DEK1 function to Gd^{3+} sensitivity *in planta*.

Reintroducing full-length DEK1 restores RMA current. We hypothesize that the loss of RMA current observed in *dek1-2* CALPAIN-OE callus is caused by the absence of an intact DEK1 transmembrane region. We propose that in the *dek1-3* CALPAIN-OE plants a truncated DEK1 protein, which contains the DEK1 transmembrane spans but which cannot confer normal inactivation kinetics on the RMA current is produced (Supplementary Fig. 2c). To test both these hypotheses we investigated the RMA channel activity in both the previously published and characterized *dek1-3* line complemented with the full length GFP-tagged DEK1 protein expressed under the *RPS5A* promoter (*dek1-3* DEK1)³⁰, and in a *dek1-2* mutant background into which the same, complementing full length DEK1-encoding construct had been introduced by crossing (*dek1-2* DEK1). In the majority of patches obtained from these lines, we recorded an RMA channel activity with inactivation kinetics and current amplitude resembling those of wild-type plants (Fig. 3a, b; Supplementary Fig. 7). The properties of this restored current were tested in the complemented *dek1-2* DEK1 mutant callus and found to be strikingly similar to those of the RMA current (Supplementary Fig. 11). Our results show that reintroducing the full length DEK1 into *dek1* mutant backgrounds leads to the restoration of an RMA current similar to that observed in wild-type callus.

Discussion

In summary, we show that the mechanosensitive activity of a Ca^{2+} -permeable channel present in the plasma membrane of *Arabidopsis* callus-derived protoplast requires the TM domains of the DEK1 protein. It has previously been demonstrated that the CALPAIN domain of DEK1 is released from the plasma membrane by an autolytic cleavage event³², and that the CALPAIN activity is enhanced by Ca^{2+} ions⁴³. We therefore propose a model in which DEK1 activity leads to transient elevation of cytoplasmic Ca^{2+} concentration during mechanical stimulation, which is locally transduced by autolytic cleavage (and thus activation) of the CALPAIN domain³². Based on the phenotype of lines with reduced DEK1 activity, this mechanotransduction pathway is likely required for the maintenance of cell–cell contacts and epidermis integrity³³, consistent with embryo lethality in loss of function alleles. In turn, DEK1-dependent epidermis integrity is required for the propagation of mechanical signals between neighboring cells, in a feedback loop. Given the absolute requirement for DEK1 in both coherent embryogenesis and post-embryonic growth^{29–31, 33} (summarized in Supplementary Fig. 12), our work suggests a key contribution of mechanoperception to plant development. The absence of cell migration in plant tissues may explain why the perception of tension between adjacent cells plays such an essential role in development in this kingdom.

This study, together with previously published results from both our work and the work of others^{32, 56} supports the idea that the CALPAIN domain of DEK1 is the effector component of the protein in terms of mechanotransduction. Whether the transmembrane portion of DEK1 forms a mechanically activated Ca^{2+} channel per se, or it is the mechano-sensor associated with an independent Ca^{2+} channel remains to be determined. However, the published Ca^{2+} dependence of CALPAIN catalytic activity, combined with our electrophysiological results and the fact that CALPAIN complemented *dek1* null mutants develop relatively normally, suggests that once activated by cleavage, the CALPAIN domain of DEK1 can respond to changes in cytoplasmic Ca^{2+} levels mediated by other Ca^{2+} channels. These could include proteins such as MCA1 and MCA2, or potentially the plant PIEZO protein, which has yet to be functionally characterized. The presence of such channels is consistent with our finding that

CALPAIN complemented *dek1-2* plants retain some Gd^{3+} sensitivity in *planta*.

Interestingly, intracellularly localized calpains have been proposed to act downstream of mechanosensitive ion channels in animals, to regulate a variety of cellular processes, including cell-to-cell adhesion^{58–61}. In this context, the association of a calpain protease with a domain influencing a mechanosensitive Ca^{2+} channel activity in the membranes of multicellular plants may reflect evolutive convergence between these two kingdoms.

In animals, the characterization of mechanosensitive channels Piezo1 and Piezo2 has revealed novel roles of mechanoperception in development and physiology. For instance, respiration in lungs relies on Piezo2-expressing sensory neurons, which use mechanical signals to sense airway stretching⁶². Axon growth and trajectory was also shown to depend on Piezo1-dependent perception of the mechanical environment of neurons⁶³. The association of DEK1, a key developmental regulator, with a mechanosensitive channel activity paves the way to the identification of new roles of mechanical signals in plant development.

Methods

Plant material. *Arabidopsis* (*Arabidopsis thaliana*) Columbia-0 (Col-0) wild-type seeds were obtained from Nottingham *Arabidopsis* Stock Centre (NASC, School of Biosciences, University of Nottingham, United Kingdom). Plants expressing the *pRPS5A:CALPAIN-HIS* in a wild-type background (CALPAIN-OE), *pRPS5A:CALPAIN-GFP*, or *pRPS5A:DEK1* in the *dek1-3* mutant background and *pRPS5A:CALPAIN-GFP* in the *dek1-2* background have been previously described^{18, 30}. The *pRPS5A:DEK1* construct was transferred from the *dek1-3* to the *dek1-2* background by crossing, and plants were genotyped as described in Supplementary Fig. 3.

Plant and callus growth conditions. For *in vitro* cultures, seeds were surface-sterilized with chlorine gas, sown on square plates, and stratified for 2 or 3 days in the dark at 4 °C. After stratification, seeds were germinated in a growth chamber under a 16-h light/8-h dark cycle at 21 °C.

Callus generation. Surface-sterilized seeds were sown on “initiation medium” containing 4.3 g/L Murashige and Skoog salts (MS, Sigma-Aldrich), 2% sucrose, 10 mg/L myo-inositol, 100 µg/L nicotinic acid, 1 mg/L thiamine-HCl, 100 µg/L pyridoxine-HCl, 400 µg/L glycine, 0.23 µM kinetin, 4.5 µM 2,4-D, 1% Phytigel, (pH 5.7). For callus generation, seeds were cultured in a growth chamber for 15 days. Calli were then transferred onto “maintenance medium” containing 4.3 g/L MS salts (Sigma-Aldrich), 2% sucrose, 10 mg/L myo-inositol, 100 µg/L nicotinic acid, 1 mg/L thiamine-HCl, 100 µg/L pyridoxine-HCl, 400 µg/L glycine, 0.46 µM kinetin, 2.25 µM 2,4-D, 1% phytigel, (pH 5.7), and sub-cultured every 15 days onto fresh “maintenance medium”.

Protoplasting protocol. Calli were digested for 15 min at 22 °C under hyper-osmotic conditions (2 mM CaCl_2 , 2 mM MgCl_2 , 1 mM KCl, 10 mM MESs (pH 5.5), 0.2% cellulysin (Calbochem), 0.2% cellulase RS (Onozuka RS, Yakult Honsha Co.), 0.004% pectolyase Y23 (Kikkoman Corporation), 0.35% bovine serum albumine and mannitol to 600 mOsmol. For enzyme removal, the preparation was washed twice with 2 mM CaCl_2 , 2 mM MgCl_2 , 10 mM MES (pH 5.5), and mannitol to 600 mOsmol. For protoplast release, the preparation was incubated with 2 mM CaCl_2 , 2 mM MgCl_2 , 10 mM MES (pH 5.5), and mannitol to 280 mOsmol. The suspension was filtered through a 50 µm nylon mesh.

Electrophysiology. Patch-clamp experiments were performed at room temperature with a patch-clamp amplifier (model 200 A, Axon Instruments, Foster City, CA) and a Digidata 1322 A interface (Axon Instruments). Currents were filtered at 5 kHz, digitized at 20 kHz, and analyzed with pCLAMP8.1 and Clampfit 10 software. During patch-clamp recordings, cells were held at a holding potential (corrected from liquid junction potential) of –16 or –6 mV depending on the composition of the pipette solution and pressure was applied with a high speed pressure-clamp system (ALA Scientific Instrument, NY), allowing the application of precise and controlled pressure pulses in the pipette^{15, 64}. For Ca^{2+} and Ba^{2+} current recordings, bath solutions contained: 50 mM CaCl_2 or 50 mM BaCl_2 , respectively, and 5 mM MgCl_2 , 10 mM MES-Tris (pH 5.6); while pipettes were filled with: 150 mM CsMES, 2 mM MgCl_2 , 5 mM EGTA, 4.2 mM CaCl_2 , and 10 mM Tris-HEPES (pH 7.2), supplemented with 5 mM MgATP. To remove Ca^{2+} , a solution containing: 100 mM TeacCl, 5 mM MgCl_2 and 10 mM MES-Tris (pH 5.6) was used.

For Cl^- current recordings, bath solution contained (mM): 50 CaCl_2 , 5 MgCl_2 , 10 MES-Tris (pH 5.6) and pipettes were filled with (mM): 150 CsCl, 2 MgCl_2 , 5 EGTA, 4.2 CaCl_2 , and 10 Tris-HEPES (pH 7.2), supplemented with 5 MgATP. For

inhibitor treatments, 0.25 mM LaCl_3 or 0.25 mM GdCl_3 were added to the bath solution, osmolarity was adjusted with mannitol to 450 mOsmol for the bath solution and to 460 mOsmol for the pipette solution using an osmometer (Type 15, Löser Meßtechnik). Gigaohm resistance seals between pipettes (pipette resistance, 0.8–1.5 M Ω) (coated with Sylgard (General Electric) pulled from capillaries (Kimax-51, Kimble Glass)) and protoplast membranes were obtained with gentle suction leading to the whole-cell configuration, then excised to an outside-out configuration. The current inactivation kinetics were fitted with a mono-exponential function: $F(t) = A \times e^{-t/\tau} + C$, where A is the coefficient, τ is the time constant, and C represents the maximum current intensity.

Genomic DNA extraction and genotyping. Plant callus DNA was extracted using a rapid CTAB isolation technique⁶⁵. *dek1-2* and *dek1-3* mutants were genotyped using the primers shown in Supplementary Table 1 and Supplementary Fig. 3. The following thermal profile was used: 95 °C for 5 min, 35 cycles of 95 °C for 45 s, 60 °C for 45 s, 72 °C for 1 min, and a final extension step of 5 min at 72 °C.

RNA extraction and quantitative gene expression analysis. Callus material was collected and snap-frozen in liquid nitrogen for gene expression analysis. For each experiment, at least two independent biological replicates were used. Total RNA was extracted using the Spectrum Plant Total RNA Kit (Sigma-Aldrich). Total RNAs were digested with Turbo DNA-free DNase I (Ambion) according to the manufacturer's protocol. RNA concentrations were measured with a NanoDrop ND-1000 UV–Vis spectrophotometer (NanoDrop Technologies). One microgram of total RNA was reverse transcribed (RT) using the SuperScript VILO cDNA Synthesis Kit (Invitrogen) according to the manufacturer's instructions. PCR reactions were performed in optical 96-well plates in the StepOne Plus Real Time PCR System (Applied Biosystems). Five microliters of a 1:10 dilution of cDNA was amplified in 20 μL of reaction mix. The following thermal profile was used: 95 °C for 10 min, 40 cycles of 95 °C for 15 s, 60 °C for 30 s. Amplicon melting curves, were recorded after cycle 40 by heating from 60 to 95 °C with a ramp speed of 1 °C min^{-1} . Expression levels of each gene, relative to *EIF4A*, were determined using a modification of the Pfaffl method³³. Technical triplicates were performed for each biological sample. Primers used for RT-quantitative (q)PCR analysis are shown in Supplementary Table 1.

Antibody production. Anti-Ila1b rabbit polyclonal antibodies were generated against a peptide from the predicted catalytic domain (RGDKQFTDQEPFNC) of the *Arabidopsis* DEK1 protein using the PolyExpress Custom Polyclonal Antibody Production Package Service (GenScript).

Western blot analysis. Callus material for each genotype was snap-frozen in liquid nitrogen, thoroughly ground in pre-cooled porcelain mortars, and proteins were extracted using an extraction buffer containing 50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% Nonidet P-40, 10% glycerol, 1% deoxycholate, 1 mM EDTA and 1 $\times$ protease inhibitor cocktail P9599 (Sigma-Aldrich). Samples were incubated on ice for 1 h, centrifuged for 15 min at 14,000 $\times$ g at 4 °C to remove cell debris, and protein concentrations in supernatants were determined using a Bio-Rad protein assay (Bio-Rad). Equal amounts of proteins were loaded and resolved on 7.5% polyacrylamide/0.1% SDS gels. Proteins were transferred onto nylon membranes using an iBlot II dry blotting system (Invitrogen). Membranes were blocked over night at 4 °C using 1 $\times$ PBS/0.2% Tween-20 (Sigma-Aldrich) with 5% non-fat milk (Regila), incubated for 2 h at room temperature with rabbit polyclonal anti-Ila1b (Genescript) or mouse monoclonal anti-alpha tubulin clone B-5-1-2 (Sigma-Aldrich, catalogue number T5168) primary antibodies at dilutions of 1:1000 and 1:2000, respectively. Secondary horseradish peroxidase-conjugated anti-rabbit and anti-mouse antibodies (Promega, catalogue numbers S3731 and S3721, respectively) were used at a dilution of 1:45,000. Blots were incubated with Super Signal West Femto reagents (Thermo Scientific) according to the manufacturer's instructions, and then exposed to Super RX film (Fujifilm). Membranes were first incubated with anti-Ila1b antibodies, and then re-probed with anti-alpha tubulin antibodies.

Gadolinium sensitivity experiments. Seeds were surface-sterilized with chlorine gas, sown on square plates (in a single row in the upper part of the plates), and stratified for 2 days in the dark at 4 °C. After stratification, seeds were germinated in a growth chamber under a 16-h light/8-h dark cycle at 21 °C, with plates kept in a vertical position. Plates containing 1/5 strength (0.86 gr/L) MS medium (Duchefa) pH = 5.7, 1% sucrose and 0.8% phytoagar, were supplemented with gadolinium (III) chloride hexahydrate (Sigma-Aldrich). Primary root length was scored after 5 or 6 days of growth with ImageJ software. Different genotypes were sown side by side on the same plate and in multiple combinations to buffer possible positional effects.

Data availability. The authors declare that all data supporting the findings of this study are available within the manuscript and its Supplementary Files or are available from the corresponding authors upon request.

Received: 3 July 2015 Accepted: 2 August 2017

Published online: 18 October 2017

References

- Dahmann, C., Oates, A. C. & Brand, M. Boundary formation and maintenance in tissue development. *Nat. Rev. Genet.* **12**, 43–55 (2011).
- Gudipaty, S. A. et al. Mechanical stretch triggers rapid epithelial cell division through Piezo1. *Nature* **543**, 118–121 (2017).
- Ranade, S. S. et al. Piezo1, a mechanically activated ion channel, is required for vascular development in mice. *Proc. Natl Acad. Sci. USA* **111**, 10347–10352 (2014).
- Li, J. et al. Piezo1 integration of vascular architecture with physiological force. *Nature* **515**, 279–282 (2014).
- Porazinski, S. et al. YAP is essential for tissue tension to ensure vertebrate 3D body shape. *Nature* **521**, 217–221 (2015).
- Pouille, P. A., Ahmadi, P., Brunet, A. C. & Farge, E. Mechanical signals trigger Myosin II redistribution and mesoderm invagination in *Drosophila* embryos. *Sci. Signal.* **2**, ra16 (2009).
- Fernandez-Gonzalez, R., Simoes Sde, M., Roper, J. C., Eaton, S. & Zallen, J. A. Myosin II dynamics are regulated by tension in intercalating cells. *Dev. Cell* **17**, 736–743 (2009).
- Koser, D. E. et al. Mechanosensing is critical for axon growth in the developing brain. *Nat. Neurosci.* **19**, 1592–1598 (2016).
- Bagriantsev, S. N., Gracheva, E. O. & Gallagher, P. G. Piezo proteins: regulators of mechanosensation and other cellular processes. *J. Biol. Chem.* **289**, 31673–31681 (2014).
- Monshausen, G. B. & Haswell, E. S. A force of nature: molecular mechanisms of mechanoperception in plants. *J. Exp. Bot.* **64**, 4663–4680 (2013).
- Monshausen, G. B. & Gilroy, S. Feeling green: mechanosensing in plants. *Trends Cell Biol.* **19**, 228–235 (2009).
- Hedrich, R. Ion channels in plants. *Physiol. Rev.* **92**, 1777–1811 (2012).
- Haswell, E. S., Peyronnet, R., Barbier-Brygoo, H., Meyerowitz, E. M. & Frachisse, J. M. Two MscS homologs provide mechanosensitive channel activities in the *Arabidopsis* root. *Curr. Biol.* **18**, 730–734 (2008).
- Hamilton, E. S. et al. Mechanosensitive channel MSL8 regulates osmotic forces during pollen hydration and germination. *Science* **350**, 438–441 (2015).
- Peyronnet, R., Tran, D., Girault, T. & Frachisse, J. M. Mechanosensitive channels: feeling tension in a world under pressure. *Front. Plant Sci.* **5**, 558 (2014).
- Maksaev, G. & Haswell, E. S. MscS-Like10 is a stretch-activated ion channel from *Arabidopsis thaliana* with a preference for anions. *Proc. Natl Acad. Sci. USA* **109**, 19015–19020 (2012).
- Furuichi, T., Iida, H., Sokabe, M. & Tatsumi, H. Expression of *Arabidopsis* MCA1 enhanced mechanosensitive channel activity in the *Xenopus laevis* oocyte plasma membrane. *Plant Signal. Behav.* **7**, 1022–1026 (2012).
- Yamanaka, T. et al. MCA1 and MCA2 that mediate Ca^{2+} uptake have distinct and overlapping roles in *Arabidopsis*. *Plant Physiol.* **152**, 1284–1296 (2010).
- Nakano, M., Iida, K., Nyunoya, H. & Iida, H. Determination of structural regions important for Ca^{2+} uptake activity in *Arabidopsis* MCA1 and MCA2 expressed in yeast. *Plant Cell Physiol.* **52**, 1915–1930 (2011).
- Nakagawa, Y. et al. *Arabidopsis* plasma membrane protein crucial for Ca^{2+} influx and touch sensing in roots. *Proc. Natl Acad. Sci. USA* **104**, 3639–3644 (2007).
- Yuan, F. et al. OSCA1 mediates osmotic-stress-evoked Ca^{2+} increases vital for osmosensing in *Arabidopsis*. *Nature* **514**, 367–371 (2014).
- Cosgrove, D. J. & Hedrich, R. Stretch-activated chloride, potassium, and calcium channels coexisting in plasma membranes of guard cells of *Vicia faba* L. *Planta* **186**, 143–153 (1991).
- Ding, J. P. & Pickard, B. G. Mechanosensory calcium-selective cation channels in epidermal cells. *Plant J.* **3**, 83–110 (1993).
- Dutta, R. & Robinson, K. R. Identification and characterization of stretch-activated ion channels in pollen protoplasts. *Plant Physiol.* **135**, 1398–1406 (2004).
- Furuichi, T., Tatsumi, H. & Sokabe, M. Mechano-sensitive channels regulate the stomatal aperture in *Vicia faba*. *Biochem. Biophys. Res. Commun.* **366**, 758–762 (2008).
- Zhang, W., Fan, L. M. & Wu, W. H. Osmo-sensitive and stretch-activated calcium-permeable channels in *Vicia faba* guard cells are regulated by actin dynamics. *Plant Physiol.* **143**, 1140–1151 (2007).
- Ahn, J. W., Kim, M., Lim, J. H., Kim, G. T. & Pai, H. S. Phytocain controls the proliferation and differentiation fates of cells in plant organ development. *Plant J.* **38**, 969–981 (2004).
- Perroud, P. F. et al. Defective Kernel 1 (DEK1) is required for three-dimensional growth in *Physcomitrella patens*. *New Phytol.* **203**, 794–804 (2014).
- Lid, S. E. et al. Mutation in the *Arabidopsis thaliana* DEK1 calpain gene perturbs endosperm and embryo development while over-expression affects organ development globally. *Planta* **221**, 339–351 (2005).

30. Johnson, K. L., Degnan, K. A., Ross Walker, J. & Ingram, G. C. AtDEK1 is essential for specification of embryonic epidermal cell fate. *Plant J.* **44**, 114–127 (2005).
31. Becraft, P. W., Li, K., Dey, N. & Asuncion-Crabb, Y. The maize dek1 gene functions in embryonic pattern formation and cell fate specification. *Development* **129**, 5217–5225 (2002).
32. Johnson, K. L., Faulkner, C., Jeffree, C. E. & Ingram, G. C. The phyto-calpain defective kernel 1 is a novel *Arabidopsis* growth regulator whose activity is regulated by proteolytic processing. *Plant Cell* **20**, 2619–2630 (2008).
33. Galletti, R. et al. DEFECTIVE KERNEL 1 promotes and maintains plant epidermal differentiation. *Development* **142**, 1978–1983 (2015).
34. Lid, S. E. et al. The defective kernel 1 (*dek1*) gene required for aleurone cell development in the endosperm of maize grains encodes a membrane protein of the calpain gene superfamily. *Proc. Natl Acad. Sci. USA* **99**, 5460–5465 (2002).
35. Kumar, B. K., Venkateswaran, K. & Kundu, S. Alternative conformational model of a seed protein DeK1 for better understanding of structure-function relationship. *J. Proteins Proteom.* **1**, 77–90 (2010).
36. Amanda, D. et al. DEFECTIVE KERNEL1 (DEK1) regulates cell walls in the leaf epidermis. *Plant Physiol.* **172**, 2204–2218 (2016).
37. Galletti R., Verger S., Hamant O. & Ingram G. Developing a thick skin: a paradoxical role for mechanical tension in maintaining epidermis integrity? *Development* **143**, 3249–3258 (2016).
38. Kutschera, U. & Niklas, K. J. The epidermal-growth-control theory of stem elongation: an old and a new perspective. *J. Plant Physiol.* **164**, 1395–1409 (2007).
39. Hamant, O. et al. Developmental patterning by mechanical signals in *Arabidopsis*. *Science* **322**, 1650–1655 (2008).
40. Savaldi-Goldstein, S., Peto, C. & Chory, J. The epidermis both drives and restricts plant shoot growth. *Nature* **446**, 199–202 (2007).
41. Reinhardt, D., Frenz, M., Mandel, T. & Kuhlmeier, C. Microsurgical and laser ablation analysis of interactions between the zones and layers of the tomato shoot apical meristem. *Development* **130**, 4073–4083 (2003).
42. Fridman, Y. et al. Root growth is modulated by differential hormonal sensitivity in neighboring cells. *Genes Dev.* **28**, 912–920 (2014).
43. Wang, C. et al. The calpain domain of the maize DEK1 protein contains the conserved catalytic triad and functions as a cysteine proteinase. *J. Biol. Chem.* **278**, 34467–34474 (2003).
44. Friedrich, P., Tompa, P. & Farkas, A. The calpain-system of *Drosophila melanogaster*: coming of age. *Bioessays* **26**, 1088–1096 (2004).
45. Farkas, A. et al. Autolytic activation and localization in schneider cells (S2) of calpain B from *drosophila*. *Biochem. J.* **378**, 299–305 (2004).
46. Ermolova, N., Kramerova, I. & Spencer, M. J. Autolytic activation of calpain 3 proteinase is facilitated by calmodulin protein. *J. Biol. Chem.* **290**, 996–1004 (2015).
47. Ermakov, Y. A., Kamaraju, K., Sengupta, K. & Sukharev, S. Gadolinium ions block mechanosensitive channels by altering the packing and lateral pressure of anionic lipids. *Biophys. J.* **98**, 1018–1027 (2010).
48. Millet, B. & Pickard, B. G. Early wrong-way response occurs in orthogravitropism of maize roots treated with lithium. *Physiol. Plant* **72**, 555–559 (1988).
49. Yang, X. C. & Sachs, F. Block of stretch-activated ion channels in *Xenopus* oocytes by gadolinium and calcium ions. *Science* **243**, 1068–1071 (1989).
50. Ruknudin, A., Sachs, F. & Bustamante, J. O. Stretch-activated ion channels in tissue-cultured chick heart. *Am. J. Physiol.* **264**, H960–H972 (1993).
51. Hamill, O. P. & McBride, D. W. Jr. The pharmacology of mechanogated membrane ion channels. *Pharmacol. Rev.* **48**, 231–252 (1996).
52. Haley, A. et al. Effects of mechanical signaling on plant cell cytosolic calcium. *Proc. Natl Acad. Sci. USA* **92**, 4124–4128 (1995).
53. Allen, G. J. & Sanders, D. Two voltage-gated, calcium release channels coreside in the vacuolar membrane of broad bean guard cells. *Plant Cell* **6**, 685–694 (1994).
54. Grabov, A. & Blatt, M. R. A steep dependence of inward-rectifying potassium channels on cytosolic free calcium concentration increase evoked by hyperpolarization in guard cells. *Plant Physiol.* **119**, 277–288 (1999).
55. Lemtiri-Chlieh, F. & Berkowitz, G. A. Cyclic adenosine monophosphate regulates calcium channels in the plasma membrane of *Arabidopsis* leaf guard and mesophyll cells. *J. Biol. Chem.* **279**, 35306–35312 (2004).
56. Liang, Z. et al. The catalytic domain CysPc of the DEK1 calpain is functionally conserved in land plants. *Plant J.* **75**, 742–754 (2013).
57. Ruiz-Herrera, L., Sanchez-Calderon, L., Herrera-Estrella, L. & Lopez-Bucio, J. Rare earth elements lanthanum and gadolinium induce phosphate-deficiency responses in *Arabidopsis thaliana* seedlings. *Plant Soil* **353**, 231–247 (2012).
58. Sorimachi, H., Mamitsuka, H. & Ono, Y. Understanding the substrate specificity of conventional calpains. *Biol. Chem.* **393**, 853–871 (2012).
59. Wells, A., Huttenlocher, A. & Lauffenburger, D. A. Calpain proteases in cell adhesion and motility. *Int. Rev. Cytol.* **245**, 1–16 (2005).
60. Franco, S. J. et al. Calpain-mediated proteolysis of talin regulates adhesion dynamics. *Nat. Cell Biol.* **6**, 977–983 (2004).
61. Zhao, Y. et al. Regulation of cell adhesion and migration by Kindlin-3 cleavage by calpain. *J. Biol. Chem.* **287**, 40012–40020 (2012).
62. Nonomura, K. et al. Piezo2 senses airway stretch and mediates lung inflation-induced apnoea. *Nature* **541**, 176–181 (2017).
63. Koser, D. E. et al. Mechanosensing is critical for axon growth in the developing brain. *Nat. Neurosci.* **19**, 1592–1598 (2016).
64. Peyronnet, R. et al. Piezo1-dependent stretch-activated channels are inhibited by Polycystin-2 in renal tubular epithelial cells. *EMBO Rep.* **14**, 1143–1148 (2013).
65. Stewart, C. N. & Via, L. E. A rapid CTAB DNA isolation technique useful for RAPD fingerprinting and other PCR applications. *Biotechniques* **14**, 748–750 (1993).

Acknowledgements

This work was funded by the European Research Council consolidator grant #615739 “MechanoDevo” to O.H. and the ANR Chaire d’excellence #ANR-10-CHEX-0011 “Mechanograin” to G.C.I. as well as a team grant from the Fonds de Recherche du Québec-Nature et Technologies to R.S.-N. and A.G. E.D.N. was supported by a PhD studentship from the BBSRC. R.G. was also supported by the European Research Council consolidator grants #307387 “PhyMorph”. We would like to thank Professor Mike Blatt (University of Glasgow, UK), Dr. Anne-Aliénor Very (BPMP Montpellier, France), Prof. Arezki Boudaoud (ENS de Lyon, France) and Dr Kim Johnson (University of Melbourne, Australia) for their help and for insightful discussions. We would like to thank Claire Lionnet of the Platim microscopy facility and the RDP growth facility team (coordinated by Alexis Lacroix, logistics team, and secretarial team for their invaluable technical support.

Author contributions

D.T., R.G., and A.D. performed the experiments and prepared the figures. E.N. carried out important groundwork. G.C.I., O.H., R.S.-N., A.G., and J.-M.F. supervised the work. The paper was written by G.C.I. with active input from R.G., O.H., J.-M.F., and D.T.

Additional information

Supplementary Information accompanies this paper at doi:10.1038/s41467-017-00878-w.

Competing interests: The authors declare no competing financial interests.

Reprints and permission information is available online at <http://npg.nature.com/reprintsandpermissions/>

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2017