UNIVERSITY of York

This is a repository copy of *KIN7 kinase regulates the vacuolar TPK1 K+ channel during stomatal closure*.

White Rose Research Online URL for this paper: https://eprints.whiterose.ac.uk/126750/

Version: Accepted Version

Article:

Isner, Jean-Charles Eric Francois, Begum, Afroza, Nuehse, Thomas et al. (2 more authors) (2018) KIN7 kinase regulates the vacuolar TPK1 K+ channel during stomatal closure. Current Biology. 466-472.e4. ISSN 0960-9822

https://doi.org/10.1016/j.cub.2017.12.046

Reuse

This article is distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs (CC BY-NC-ND) licence. This licence only allows you to download this work and share it with others as long as you credit the authors, but you can't change the article in any way or use it commercially. More information and the full terms of the licence here: https://creativecommons.org/licenses/

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



eprints@whiterose.ac.uk https://eprints.whiterose.ac.uk/

	1	
2	KIN7 kinase regulates the vacuolar TPK1 K $^+$ channel during stomatal	
3	closure.	
4		
5	AUTHORS: Jean Charles Isner ¹ , Afroza Begum ² , Thomas Nuehse ³ , Alistair M	
6	Hetherington ¹ and Frans J.M. Maathuis ²	
7		
8	Affiliations: ¹⁾ School of Biological Sciences, University of Bristol, Life Sciences Building,	
9	24 Tyndall Avenue, Bristol BS8 1TQ, UK,	
10	²⁾ Department of Biology, Wentworth Way, University of York, York, YO10 5DD, UK	
11	³⁾ Faculty of Life Sciences, Michael Smith Building, Oxford Road, Manchester, M13 9PT,	
12	UK	
13		
14	Corresponding author and lead contact: Frans J.M. Maathuis, Department of Biology,	
15	University of York, York, YO10 5DD, United Kingdom. Tel: +44-1904-328652, Fax:	
16	+44-1904-328505, email: fjm3@york.ac.uk	
17		
18		
19		
20		
21		
22		

23 Summary

24 Stomata are leaf pores that regulate CO₂ uptake and evapotranspirational water loss. By 25 controlling CO₂ uptake stomata impact on photosynthesis and dry matter accumulation. The regulation of evapotranspiration is equally important because it impacts on nutrient 26 27 accumulation, leaf cooling and enables the plant to limit water loss during drought [1]. Our work centres on stomatal closure [2-6]. This involves loss of potassium from the guard cell 28 29 by a two-step process. Salt is released across the plasma membrane via anion channels 30 such as SLAC1 [7-9] and depolarisation-activated channels such as GORK [10,11] with 31 the net result that cations and anions exit guard cells. However, this critically depends on 32 K⁺ release from the vacuole; With ~160 and 100 mM K⁺ in cytoplasm and vacuole of open 33 guard cells [12], vacuolar K⁺ efflux is driven by the negative tonoplast potential and this 34 expels K⁺ from the vacuole via tonoplast K⁺ channels like TPK1. In all, guard cell salt release leads to a loss of turgor that brings about stomatal closure. First we show that the 35 36 TPK1 vacuolar K⁺ channel is important for ABA and CO₂-mediated stomatal closure. Next we reveal that during ABA and CO2-mediated closure, TPK1 is phosphorylated and 37 38 activated by the KIN7 receptor like protein kinase (RLK) which co-expresses in the 39 tonoplast and plasma membrane. The net result is K⁺ release from the vacuole. Taken 40 together our work reveals new components involved in guard cell signalling and describes 41 a new mechanism potentially involved in fine-tuning ABA and CO₂-induced stomatal closure. 42

43

44 Results and Discussion

The tonoplast located channel AtTPK1 [13,14] was previously shown to affect ABAinduced stomatal closure in *Arabidopsis* [15]. In this investigation we used patch-clamp

47 electrophysiology to show that GC vacuolar preparations exhibited TPK1 activity (Fig. 1A). Tonoplast channels may show tissue specific properties as was shown for TPC1 [16]. 48 49 However, single channel conductance and weak voltage dependence of GC TPK1 activity were the same as previously reported for TPK1 currents from mesophyll cell vacuoles [15]. 50 51 Addition of 0.5 mM Mg-ATP to the cytoplasmic side of the membrane led to a rapid but moderate increase in channel activity. Channel activity was further increased when both 52 53 Mg-ATP and 14-3-3 protein were present as was shown previously for TPK1 in mesophyll 54 cells [17,18]. On average, ATP alone caused an increase in channel activity that was 55 equivalent to an increase in open time (and therefore current) of ~50% whereas ATP+14-56 3-3 more than tripled open probability and current (Figure 1B).

57

The S42 residue forms part of the 14-3-3 binding domain in the TPK1 N-terminus and by 58 using a reconstituted TPK1 N-terminus, Latz, et al. [18] showed that S42 phosphorylation 59 60 is required for TPK1-14-3-3 interaction. To test if the same residue is responsible for the phosphorylation-dependent changes in GC TPK1 activity, we transiently expressed a 61 62 mutated (S42A) version of TPK1 in the tpk1 T-DNA knock out background [15]. The S42A 63 version mimics a constitutively non-phosphorylated form. Normal currents were recorded showing that the channel is fully functional (Figure 1A) but the stimulating effects of both 64 65 ATP and ATP+14-3-3 were abolished (Figure 1B). Taken together these data show that in GCs TPK1 shows low basal activity, which increases after phosphorylation at S42 and 66 is further elevated after binding of 14-3-3. Figure S1 shows that these characteristics were 67 68 retained when measuring macroscopic currents.

69

70 ABA causes phosphorylation of TPK1 in planta

3

71 Knowing that ABA induces stomatal closure we decided to test whether TPK1 phosphorylation is ABA-dependent. GC protoplasts were isolated from the tpk1 mutant 72 that had been transiently transformed with TPK1::YFP (MW 67.7 kD) and were then 73 probed with a phosphoserine specific antibody that recognised the phosphorylated 14-3-74 75 3 binding domain [19]. Figure 1C shows a band around 70 kD that increased in intensity after ABA treatment and was absent in the (non-transformed) tpk1 null mutant. We also 76 77 tested if ABA increased TPK1 phosphorylation in intact tissue: treating leaves with 40 µM 78 ABA for one hour led to increased TPK1 phosphorylation detected in isolated GC 79 protoplasts (Figure 1D). On the basis of densitometry, 2.5-3-fold higher phosphorylation 80 signals were recorded after ABA treatment (Figure 1E). These data allow us to conclude 81 that phosphorylation of TPK1 is regulated by ABA and opens up the possibility that this is 82 part of the GC ABA signalling network and may be necessary for stomatal closure. It would be interesting to further test this idea using a phosphomimic TPK1 version, for example 83 84 through substitution of S42 with glutamate. Such genotypes could be evaluated for their transpirational flux, steady state stomatal conductance and responses to various stimuli. 85

86 A receptor like kinase is involved in TPK1 phosphorylation

To help identify the protein kinase that was responsible for TPK1 phosphorylation we 87 88 interrogated the SUBA database (http://suba3.plantenergy.uwa.edu.au/) for protein 89 kinases that are annotated as tonoplast-localised. A total of 22 kinase isoforms was found 90 and loss of function mutants were obtained for 16 of these (Table S1). Mutants were tested for an altered channel 'activation response' to ATP and 14-3-3 (Figures S2A and S2B and 91 Data S1). The amount of activation generated by ATP varied but was not significantly 92 93 different between any of the genotypes (Figure S2B). However, activation by ATP+14-3-3 was significantly (p<0.001) lower in the kin7 mutant compared to that observed in WT. 94 4

Furthermore, a role for KIN7 in TPK1 phosphorylation was confirmed by the large reduction in ABA-induced TPK1 phosphorylation in the *kin7* loss of function mutants (Figure 1E).

KIN7 is ubiquitously expressed in all leaf tissues but to investigate whether KIN7 and TPK1 98 99 physically interact at the GC tonoplast we employed a BiFC strategy [20,21]. Figure 2A 100 shows an intact and an osmotically ruptured GC protoplast co-transformed with TPK1-101 YFP_{Nt} and TPK1-YFP_{Ct}. As expected, fluorescence is clearly localised in the tonoplast. 102 Similar to the results with TPK1 only, when TPK1-YFPct was co-transformed with KIN7-103 YFP_{Nt} there was a prominent fluorescence signal (Figure 2B, bottom left) in many 104 protoplasts. After osmotic lysis, clear but low intensity fluorescence signal could be 105 observed in the tonoplast of some cells (Figure 2B, bottom right). In contrast, when TPK1-106 YFP_{Ct} was co-transformed with KIN8-YFP_{Nt} (a kinase very similar to KIN7, see Table S1) 107 no signal was observed in intact or lysed protoplasts (Figure 2C, n>200). BiFC 108 experimentation with KIN12, another comparable kinase (Table S1), produced 109 fluorescence signal but never in the tonoplast (Figure S2, n>200). These results suggest 110 that TPK1 and KIN7 can directly interact at the tonoplast but that either the incidence is 111 low (see Figure S2), or alternatively, that relatively few proteins are involved.

Apart from electrophysiology and BiFC, we employed a third strategy to probe TPK1-KIN7 interaction. Pull down assays were carried out where the TPK1 N- terminus containing the 14-3-3 domain was used as bait [19] and plant extract derived from shoot tissue expressing KIN7::YFP as prey. Figure S3 shows that, in addition to the BiFC and the electrophysiological data, pull down assays too confirm the interaction between TPK1 and KIN7.

118

5

119 kin7 and tpk1 are compromised in ABA and CO₂-induced stomatal closure

120 Previously, we showed that in tpk1 loss of function mutants ABA-induced stomatal closure 121 is slower than wild type [15]. In the light of our data that TPK1-mediated K⁺ release is likely 122 to depend on KIN7-mediated TPK1 phosphorylation we compared the kinetics of stomatal 123 closure in wild type, *tpk1* and *kin* loss of function mutants. When ABA-induced stomatal 124 closure was measured in tpk1 a reduced closing response was observed (Figure 2D) as 125 was previously observed [15]. In two independent mutant alleles of kin7 delayed ABAinduced closure was also observed. When the kin7-1 line was rescued with a 126 127 35S:KIN7:YFP construct, the transformed line reverted to the wildtype phenotype. These 128 results suggest that TPK1 and KIN7 are part of the GC ABA signalling network. This idea 129 is supported by the observation that ABA dependent TPK1 phosphorylation was greatly reduced in both kin7 mutants (Figure 1E). To investigate whether TPK1 and KIN7 are 130 involved in other stimuli that cause stomatal closure we exposed the WT, tpk1 and kin7 131 132 mutants to elevated levels of CO₂. Figure 2E shows that tpk1 and kin7 mutants are 133 markedly unresponsive to 1000 ppm CO₂ whether assayed after 3 hours (Figure 2E) or during continuous conductance measurements (Figure S3). These data suggest that 134 135 TPK1 and KIN7 are also involved in the GC CO₂ signalling network.

The latter begs the question whether the KIN7-14-3-3-TPK1 pathway also pertains to other closing stimuli such as the transition from light to dark or a reduction in relative humidity. Preliminary experiments did not show any drought related phenotype in *tpk1* mutants but, since the *tpk1* stomatal phenotype primarily affects closing dynamics rather than the absolute conductance levels [15], more subtle treatments such as alternating relative humidity may be required. Transition to darkness is another stimulus which so far has not been investigated in either the *tpk1* or *kin7* genetic background. Future testing of these

and other closing stimuli should help determine whether the signalling mechanism wedescribe has more general validity.

145

146 The KIN7 kinase shows dual membrane localisation

147 As there are reports that KIN7 is localized to the plasma membrane (SUBA: suba2.plantenergy.uwa.edu.au/) we decided to test whether this protein had a dual 148 149 membrane localization in GCs. To test this we used KIN7::YFP fusion proteins which were 150 stably expressed under control of the 35S promoter or the endogenous KIN7 promoter. In 151 GC protoplasts derived from transgenic lines, KIN7-YFP was occasionally observed in 152 both plasma membrane and tonoplast (Figure 3A-F). However, in most cases plasma 153 membrane signal greatly predominated (Figure 3C and D) to the extent that the tonoplast signal was only detected after osmotic lysis (Figure 3E and F) when weak but distinct 154 155 fluorescence can be distinguished in a proportion of cells as was seen in the BiFC 156 experiments. This pattern was consistent for KIN7-YFP expression, irrespective of the 157 promoter driving expression (Figure S4) and clearly point to dual localisation of KIN7. The latter prompted us to investigate whether KIN7 localisation is sensitive to ABA. One half 158 159 of a KIN7-YFP transformed GC population was treated with 40 µM ABA and subsequently 160 the proportion of protoplasts with tonoplast located signal was determined. Figure 3G 161 shows that the fraction of cells with tonoplast signal more than doubles after exposure to 162 ABA in a time dependent manner. To independently confirm these findings, we carried out Western analyses on tonoplast enriched membrane fractions using the tonoplast 163 164 aguaporin TIP1;1 as a tonoplast specific marker [22]. Figure 3H shows that initially there 165 is a minimal KIN7 signal in the lower phase of a two phase partitioned membrane prep. 166 However, relative to the tonoplast marker TIP1;1, and within 30 min ABA exposure, the

167 KIN7 signal is greatly enhanced, to more than 4-fold the initial value (Figure 3I) while 168 similar experiments using high CO₂ treatment (1000 ppm, 3h) showed comparable results 169 with an approximately 3-fold increase towards tonoplast expression (Figure S4). These results show that ABA may affect TPK1 activity in less than 30 minutes. Cation flux 170 171 measurements from Commelina epidermal strips [23] show remarkably similar kinetics of 10-20 minutes between ABA addition and cation release. However, these values are 172 173 considerably slower than what has been seen for some plasma membrane anion 174 channels. For example, Levshenko et al [24] recorded anion channel activation 1-2 175 minutes after ABA exposure. There may be several explanations for this difference; Anion 176 channel activation is one of the first responses to ABA and may therefore precede slower 177 cation channel activation. TPK1 activation in intact tissue may be accelerated by unknown cell wall components or, alternatively, different ABA receptors may be involved in the 178 179 coupling to various membranes.

180 The above results show that ABA and CO₂ treatment led to an increase in tonoplast KIN7 181 signal. The data do not allow to distinguish whether elevated tonoplast expression was 182 due to de novo expression or a consequence of intracellular trafficking. However, 183 preliminary experiments where tissue was treated with cycloheximide (a protein synthesis 184 inhibitor) or chlorpromazine (an endocytosis inhibitor) suggest that tonoplast KIN7 185 expression was not affected by cycloheximide (Figure S4) but sensitive to chlorpromazine. 186 Such results suggest that endocytosis, rather than de novo protein synthesis, is an essential feature of the shift in KIN7 expression toward the tonoplast. 187

188 Conclusions

There is a number of conclusions that can be drawn from our work. Our data showing that TPK1 and KIN7 are involved in CO₂ and ABA is further evidence that both these closure 8 191 signals are able to access a common set of signalling components whose role it is to bring 192 about stomatal closure [25]. In addition to identifying that TPK1 activity is regulated by 193 protein phosphorylation we also report the identity of the regulatory protein kinase. KIN7 194 has all the hallmarks of an LRR-receptor kinase. It is ubiquitously expressed in many 195 tissues, including mesophyll cells. In addition to GCs, mesophyll cells have been shown 196 to respond to ABA, for example by reducing cell volume (e.g. [26]). This opens the 197 possibility that in mesophyll cells too, KIN7-mediated TPK1 activation plays a role in ABA 198 signalling. Preliminary patch clamp recordings suggest that KIN7 does impact on TPK1 199 activity in mesophyll cells which supports the above idea but whether this is linked to 200 phosphorylation and 14-3-3 binding of mesophyll cell TPK1 remains to be tested.

201 Another major questions to emerge from our work is, what is the link between perception 202 of ABA and CO₂ on the one hand, and activation of KIN7, binding of 14-3-3 and activation 203 of TPK1 on the other? Upstream signalling components could include well known players 204 such as ABI and OST gene products. Further experimentation with ABA signalling 205 mutants, or ones in CO₂ signalling components such as HT1 kinase, RHC1 and RBOH-D/F, will help reveal such interactors. Downstream, 14-3-3 must bring about a 206 207 conformational change that greatly stimulates channel opening. One mechanism suggests that the TPK1 gate is directly controlled via Ca²⁺ binding to C-terminal EF domains [27] 208 and a model where 14-3-3 sensitises TPK1 Ca2+ dependence not only provides a 209 210 mechanistic explanation but could be tested using electrophysiology.

It is noteworthy that ABA treatment has been reported to result in KIN7 phosphorylation at
its C-terminus [28]. If the phosphorylation results in alterations to KIN7 activity this would
suggest that at least one additional protein kinase is involved in this signalling network.
Our data showing that KIN7 is localized at both the plasma and tonoplast membranes is

215 supportive of a dynamic, stimulus-induced, mechanism of TPK1 regulation. Inhibition by 216 the endocytotic inhibitor chlorpromazine of the relative shift toward tonoplast expression 217 of KIN7 suggests that the observed increase in KIN7 tonoplast localisation does not originate from de novo KIN7 biosynthesis, but occurs via a hitherto uncharacterised 218 219 trafficking pathway. Although such findings can only be preliminary and will need further 220 retrograde endocytotic trafficking of plasma membrane proteins to support, 221 endocompartments has been reported: In animals compartmentalisation of receptor 222 kinases generates endosome specific signal transduction complexes [29]. In plants too, 223 trafficking of the steroid receptor kinase BRI1 to endosomal vesicles is believed to be 224 important in intracellular signalling [30].

225 Control of TPK1 activity through the stimulus-induced localization of KIN7, be it via modulation of expression or via trafficking, represents an attractive mechanism for exerting 226 227 control over K⁺ efflux from vacuoles. We have summarised what we know about this 228 pathway in the schema described in Figure 4. What might be the function of this form of regulation? Our phenotypic data showing that mutants in this pathway are distinguished 229 230 by exhibiting slower rates of closure suggest that this pathway might be in the fine-tuning 231 of stomatal responses rather than switching closure on (or off). However, a better 232 understanding awaits the discovery of additional components in the network.

ACKNOWLEDGEMENTS: We thank Bert de Boer (Vrije Universiteit Amsterdam) for his kind gift of 14-3-3 protein and Ingo Dreyer (Universidad de Talca, Chile) for his kind gift of BiFC vectors. We thank Wioletta Pijacka (University of Bristol, UK) for her technical help with Western blotting. AMH and J-C I wish to acknowledge grant support from the UK BBSRC (BB/J002364/1) and the Leverhulme Trust.

AUTHOR CONTRIBUTIONS:

10

- JCI and FJM designed research; JCI, FJM, AB and TSN performed research; JCI and
- FJM analysed data; FJM, AMH and JCI wrote the paper.

241 **DECLARATION OF INTERESTS**

242 The authors declare no competing interests.

243 **REFERENCES**

- Hetherington, A.M., and Woodward, F.I. (2003). The role of stomata in sensing and driving
 environmental change. Nature. 424, 901-908.
- 246 2. Roelfsema, M.R., Levchenko, V., and Hedrich, R. (2004). ABA depolarizes guard cells in 247 intact plants, through a transient activation of R- and S-type anion channels. Plant J. *37*, 578-588.
- 3. Schroeder, J., Gethyn J Allen, Veronique Hugouvieux, June M Kwak, a., and Waner, D.
 (2001). Guard cell signal transduction. Annu Rev Plant Physiol Plant Mol Biol. *52*, 627-658.
- Assmann, S.M., and Jegla, T. (2016) Guard cell sensory systems: recent insights on stomatal
 responses to light, ABA and CO2. Curr Opinion Plant Biology 33, 157-167.
- 5. Raghavendra, A.S., Gonugunta, V.K., Christmann, A., and Grill, E. (2010) ABA perception and signalling. Trends in Plant Sciences 15, 395-401.
- Munemasa, S., Hauser, F., Park, J., Waadt, R., Brandt, B., and Schroeder, J.I. (2015)
 Mechanisms of ABA-mediated control of stomatal aperture. Current Opinion in Plant Biology 28,
 154-162.
- 7. Geiger, D., Maierhofer, T., AL-Rasheid, K.A.S., Scherzer, S., Mumm, P., Liese, A., Ache,
 P., Wellmann, C., Marten, I., Grill, E., et al. (2011). Stomatal closure by fast abscisic acid signaling
 is mediated by the guard cell anion channel SLAH3 and the receptor RCAR1. Sci Signal. *4*, ra32.
- S. Geiger, D., Scherzer, S., Mumm, P., Stange, A., Marten, I., Bauer, H., Ache, P., Matschi,
 S., Liese, A., Al-Rasheid, K.A.S., et al. (2009). Activity of guard cell anion channel SLAC1 is
 controlled by drought-stress signaling kinase-phosphatase pair. Proc Natl Acad Sci USA. *106*,
 21425-21430.
- 9. Imes, D., Mumm, P., Bohm, J., Al-Rasheid, K.A., Marten, I., Geiger, D., and Hedrich, R.
 (2013). Open stomata 1 (OST1) kinase controls R-type anion channel QUAC1 in *Arabidopsis* guard
 cells. Plant J. 74, 372-382.
- 267 10. Ache, P., Becker, D., Ivashikina, N., Dietrich, P., Roelfsema, M.R., Hedrich, R. (2000).
 268 GORK, a delayed outward rectifier expressed in guard cells of Arabidopsis thaliana, is a K(+)269 selective, K(+)-sensing ion channel. FEBS Lett. 486, 93-98.
- Hosy, E., Vavasseur, A., Mouline, K., Dreyer, I., Gaymard, F., Porée, F., Boucherez, J.,
 Lebaudy, A., Bouchez, D., Véry, A.-A., et al. (2003). The *Arabidopsis* outward K⁺ channel GORK
 is involved in regulation of stomatal movements and plant transpiration. Proc Natl Acad Sci USA.
- 273 100, 5549-5554.
- 12. Hills, A., Chen, Z.H., Amtmann, A., Blatt, M.R., Lew, V.L. (2012). OnGuard, a
 Computational Platform for Quantitative Kinetic Modeling of Guard Cell Physiology. Plant
 Physiol. 159(3):1026-42.
- 277 13. Dunkel, M., Latz, A., Schumacher, K., Muller, T., Becker, D., and Hedrich, R. (2008).
- Targeting of vacuolar membrane localized members of the TPK channel family. Mol Plant. 1, 938 949.

- 14. Voelker, C., Gomez-Porras, J.L., Becker, D., Hamamoto, S., Uozumi, N., Gambale, F.,
 Mueller-Roeber, B., Czempinski, K., and Dreyer, I. (2010). Roles of tandem-pore K⁺ channels in
- 282 plants a puzzle still to be solved. Plant Biol *12 Suppl 1*, 56-63.
- 15. Gobert, A., Isayenkov, S., Voelker, C., Czempinski, K., and Maathuis, F.J. (2007). The twopore channel TPK1 gene encodes the vacuolar K⁺ conductance and plays a role in K⁺ homeostasis.
 Proc Natl Acad Sci USA. *104*, 10726-10731.
- 286 16. Rienmüller, F., Beyhl, D., Lautner, S., Fromm, J., Al-Rasheid, K.A., Ache, P., Farmer, E.E.,
- Marten, I., Hedrich, R. (2010) Guard cell-specific calcium sensitivity of high density and activity
 SV/TPC1 channels. Plant Cell Physiol. *51*, 1548-54.
- 17. Isayenkov, S., Isner, J.C., and Maathuis, F.J. (2011). Rice two-pore K⁺ channels are
 expressed in different types of vacuoles. Plant Cell. 23, 756-768.
- 18. Latz, A., Becker, D., Hekman, M., Muller, T., Beyhl, D., Marten, I., Eing, C., Fischer, A.,
 Dunkel, M., Bertl, A., et al. (2007). TPK1, a Ca²⁺-regulated *Arabidopsis* vacuole two-pore K⁺
- channel is activated by 14-3-3 proteins. Plant J. 52, 449-459.
- Latz, A., Mehlmer, N., Zapf, S., Mueller, T.D., Wurzinger, B., Pfister, B., Csaszar, E.,
 Hedrich, R., Teige, M., and Becker, D. (2013). Salt stress triggers phosphorylation of the *Arabidopsis* vacuolar K⁺ channel TPK1 by calcium-dependent protein kinases (CDPKs). Mol Plant.
 6, 1274-1289.
- 298 20. Waadt, R., Schmidt, L.K., Lohse, M., Hashimoto, K., Bock, R., and Kudla, J. (2008).
 299 Multicolor bimolecular fluorescence complementation reveals simultaneous formation of 300 alternative CBL/CIPK complexes in planta. Plant J. *56*, 505-516.
- 301 21. Kudla, J., and Bock, R. (2016). Lighting the way to protein-protein interactions:
 302 Recommendations on best practices for bimolecular fluorescence complementation analyses. Plant
 303 Cell. 28, 1002-1008.
- Ludevid, D., Hofte, H., Himelblau, E., and Chrispeels, M.J. (1992). The expression pattern
 of the tonoplast intrinsic protein gamma-tip in *Arabidopsis thaliana* is correlated with cell
 enlargement. Plant Physiol. *100*, 1633-1639.
- 307 23. MacRobbie EA (2006) Control of volume and turgor in stomatal guard cells. J Membr Biol
 308 210: 131-42
- 309 24. Levchenko, V., Konrad, K.R., Dietrich, P., Roelfsema, M.R., Hedrich, R.(2005). Cytosolic
- abscisic acid activates guard cell anion channels without preceding Ca2+ signals. Proc Natl Acad
 Sci U S A. 15;102(11):4203-8.
- 25. Chater, C., Peng, K., Movahedi, M., Dunn, Jessica A., Walker, Heather J., Liang, Y.-K.,
 McLachlan, Deirdre H., Casson, S., Isner, Jean C., Wilson, I., et al. Elevated CO₂-induced
 responses in stomata require ABA and ABA signaling. Curr Biol. 25, 2709-2716.
- Kolla, V.A., Suhita, D., Raghavendra, A.S. (2004) Marked changes in volume of mesophyll
 protoplasts of pea (Pisum sativum) on exposure to growth hormones. J Plant Physiol. *161*, 557-562.
- 317 27. Hartley, T.N., Maathuis, F.J.M. (2015) Allelic variation in the vacuolar TPK1 channel
 318 affects its calcium dependence. FEBS Lett. *590*, 110-117.
- 28. Chen, Y., Hoehenwarter, W., and Weckwerth, W. (2010). Comparative analysis of phytohormone-responsive phosphoproteins in *Arabidopsis* thaliana using TiO₂-phosphopeptide enrichment and mass accuracy precursor alignment. Plant J. *63*, 1-17.
- Villaseñor, R., Nonaka, H., Del Conte-Zerial, P., Kalaidzidis, Y., and Zerial, M. (2015).
 Regulation of EGFR signal transduction by analogue-to-digital conversion in endosomes. eLife. *4*, e06156.
- 325 30. Geldner, N., and Robatzek, S. (2008). Plant receptors go endosomal: a moving view on 326 signal transduction. Plant Physiol. *147*, 1565-1574.

- 327 31. Schindelin, J., Rueden, C.T., Hiner, M.C., and Eliceiri, K.W. (2015). The ImageJ
 328 ecosystem: An open platform for biomedical image analysis. Mol Reprod Dev. 82, 518-529.
- 329 32. Pijacka, W., Clifford, B., Tilburgs, C., Joles, J.A., Langley-Evans, S., and McMullen, S.
 330 (2015). Protective role of female gender in programmed accelerated renal aging in the rat. Physiol
 331 Rep. 3.
- 332 33. Hellens, R., Edwards, E.A., Leyland, N., Bean, S., and Mullineaux, P. (2000). pGreen: a
- versatile and flexible binary Ti vector for *Agrobacterium*-mediated plant transformation. Plant Mol
 Biol. 42, 819-832.
- 335 34. Pandey, S., Wang, X.Q., Coursol, S.A., and Assmann, S.M. (2002). Preparation and applications of Arabidopsis thaliana guard cell protoplasts. New Phytol. *153*, 517-526.
- 337 35. Maathuis, F.J.M., May, S.T., Graham, N.S., Bowen, H.C., Jelitto, T.C., Trimmer, P.,
 338 Bennett, M.J., Sanders, D., and White, P.J. (1998). Cell marking in *Arabidopsis thaliana* and its
 339 application to patch–clamp studies. Plant J. *15*, 843-851.
- 340 36. Sinnige, M.P., Roobeek, I., Bunney, T.D., Visser, A.J., Mol, J.N., and de Boer, A.H. (2005).
- Single amino acid variation in barley 14-3-3 proteins leads to functional isoform specificity in the
 regulation of nitrate reductase. Plant J. 44, 1001-1009.
- 343 37. Lund, A., and Fuglsang, A.T. (2012). Purification of plant plasma membranes by two-phase
 344 partitioning and measurement of H⁺ pumping. Methods Mol Biol. *913*, 217-223.
- 38. Rea, P.A., Britten, C.J., and Sarafian, V. (1992). Common identity of substrate binding
 subunit of vacuolar H⁺-translocating inorganic pyrophosphatase of higher plant cells. Plant Physiol. *100*, 723-732.
- 348

349 Figure legends:

350 Figure 1: GC TPK1 currents and ABA induced TPK1 phosphorylation.

351 (A) Representative TPK1 currents from one cytoplasmic side out excised patch shows

352 TPK1 currents in control buffer, after addition of MgATP (0.5 mM) or MgATP plus 14-3-

- 353 3 (100 nM). Mutation of the N-terminal serine 42 to alanine (S42A) leaves channel
- activity intact but abrogates the effect of ATP and 14-3-3. Currents were recorded at -

355 40 mV and left side arrows indicate closed levels. Amplitude histograms on right show

356 increase in channel openings for wildtype but not S42A. 'Po' open probability

- 357 quantification based on 60 sec records. (B) Normalised relative channel activity, based
- 358 on open probability data obtained at -80 mV, showing ATP stimulates activity by around
- 45% and ATP plus 14-3-3 stimulates activity by over 300%, while no significant effect of
- 360 ATP+14-3-3 is observed for the S42A mutant protein (n=3 independent membrane

361 patches for each condition, error bars are standard errors.) Data were analysed using 362 one way ANOVA with Tukey post-hoc analysis comparison with control conditions, 363 asterisks denoting p<0.05. (C) GCs from tpk1 plants were isolated and transiently 364 transformed with TPK1-YFP. Approx 12-16h after transformation half the protoplasts 365 were treated with ABA (1h, 40 µM). Right hand panel: non transformed (NT) protoplasts 366 show complete absence of signal ('T' are transformed protoplasts from the same prep). 367 (D) Guard cells isolated from ABA-treated tpk1 loss of function plants, control leaves 368 (WT plants) or ABA treated leaves (WT plants). In (C) and (D), top panels show typical 369 example of blots with antibody against the phosphorylated TPK1 N-terminus. Bottom 370 panels show total protein staining. (E) Densitometry based (n=3, bars denote standard 371 errors) fold-changes in TPK1 phosphorylation in control (WT-con) and ABA treated (WT ABA L) leaves or protoplasts (WT ABA P) and in null mutants of the LRR kinase 372 KIN7 (kin7 ABA). Data were analysed using one way ANOVA with Tukey post-hoc 373 analysis comparison with control conditions, asterisks denoting p<0.05. (See also 374 375 Figures S1 and Table S1).

Figure 2: Bimolecular fluorescence complementation and loss of function phenotypes in t*pk1* and *kin7*.

(A) Representative image for Arabidopsis GCs cotransformed with TPK1_{YFP-Nt} plus
TPK1_{YFP-Ct}. Note clear vacuolar fluorescence. (B) GC protoplasts expressing KIN7_{YFP-}
Nt plus TPK1_{YFP-Ct}. Note clear signal in the tonoplast. (C) GC protoplast expressing
KIN8_{YFP-Nt} plus TPK1_{YFP-Ct}. Note the absence of any fluorescence signal. In all cases,
top two panels show DIG images of intact and osmotically ruptured protoplast (releasing
the large central vacuole). Bottom panels show corresponding YFP fluorescence signal.
Scale bar is 5 µm. (D) Leaves of wild type (WT), TPK1 and KIN7 loss of function mutants

KIN7 complemented plants were exposed to 100, 10 or 1 μ M ABA and start ('Control') and end (30 minute ABA exposure) conductance values were recorded. **(E)** Stomata were opened by exposure of leaves to 400 ppm CO₂ for 2 hours. Subsequently, peels were either aerated with 1000 ppm [CO₂] or continued to be aerated with 400 ppm [CO₂]. Three hours later, stomatal apertures were measured. Data in D and E were analysed for significance using a one way ANOVA with Tukey post-hoc analysis. Asterisk denotes p<0.05. (See also Figures S2 and S3).

Figure 3: KIN7:YFP expression patterns.

15

393 (A) DIC image of intact Arabidopsis GC transformed with KIN7:YFP. (B) Corresponding 394 fluorescence image showing expression in both plasma membrane and tonoplast 395 (arrows). (C and D) DIC and fluorescence image of GC protoplast showing prominent expression in the plasma membrane only. (E) DIC image of osmotically ruptured 396 397 protoplast releasing the large vacuole. (F) Corresponding fluorescence image showing 398 weak but distinct tonoplast expression. (G) The proportion of lysed guard cell 399 protoplasts that shows tonoplast KIN7 expression increases after ABA treatment. (H) Western immunoblot showing increasing level of KIN7-GFP expression in response to 400 401 ABA. The vacuole specific aquaporin TIP1;1 was used as tonoplast marker whereas the 402 lack of cross reactivity with the H⁺-ATPase AHA1 (a plasma membrane specific marker) 403 in the 'lower phase' (LP) shows absence of plasma membrane. 'MF': microsomal 404 fraction showing positive reactivity for all three probes. (I) Quantification based on 405 densitometry measurements (using ImageJ) of relative increase in KIN7 expression in 406 the tonoplast. Scale bar in A-F is 7 µm. Data depicted in G and I are based on 3 or more 407 independent experiments, bars denote standard errors and data were analysed using

15

- 408 one way ANOVA with Tukey post-hoc analysis comparison with control conditions.
 409 Asterisk denotes p<0.05. (See also Figure S4).
- 410 Figure 4: A model for coupling ABA and TPK1.
- 411 ABA perception leads to activation of the LRR kinase KIN7. Increased tonoplast
- 412 expression of KIN7 and/or KIN7 traffic from plasma membrane (PM) to tonoplast (TO)
- 413 brings KIN7 in the vicinity of TPK1. At the tonoplast, phosphorylation of S42 in the N-
- 414 terminal 14-3-3 binding domain allows 14-3-3 binding to TPK1, which leads to drastically
- 415 increased TPK1 activity and stomatal closure.

416 **METHODS**

417 CONTACT FOR REAGENT AND RESOURCE SHARING

- 418 Further information and requests for resources and reagents should be directed to and
- 419 will be fulfilled by the Lead Contact, Frans Maathuis (frans.maathuis@york.ac.uk)

420 EXPERIMENTAL MODEL AND SUBJECT DETAILS

- 421 Plant material and growth. Arabidopsis thaliana (L) ecotype Columbia (0) wild type,
- 422 *tpk1* (SALK line 146903; [15] and kinase mutants and kinase mutants obtained from
- 423 NASC (see Table S1).

424 METHOD DETAILS

425 <u>Plants</u>

- 426 Arabidopsis thaliana (L) ecotype Columbia (0) wild type, tpk1 and kinase mutants were grown for
- 427 3 to 4 weeks in soil (F2, Levington, UK) at 18/22°C night/day temperature in a glasshouse with day
- 428 lengths of 14 h, supplemented with artificial light of around 200 μ mol m⁻² sec⁻¹ as described [17].
- 429 T-DNA insertion lines for kinase mutants and the forward and reverse primers that were used to test
- 430 for homozygosity and transcript can be found in Table S1 and S2.
- 431 Cloning of kinases in the BiFC vector
- 432 KIN8 and KIN12 were cloned from cDNA produced from total RNA: Total RNA was extracted
- 433 from Arabidopsis leaves with RNeasy Plant Mini Kit (Qiagen GmbH, Germany). First strand cDNA

- 434 was synthesised using SuperScript II Reverse Transcriptase kit (Life Technologies Ltd, UK) with
- the following primers: kin8bifcfwd and kin8bifcrev, kin12bifcfwd and kin12bifcrev (Table S2).
- 436 KIN8 and KIN12 fragments were amplified using kin8bifcfwd+kin8bifcrev and kin12bifcfwd+
- 437 kin12bifcrev primers respectively (Table S2). KIN7 was PCR-amplified from the full length cDNA
- 438 clone U12357 (<u>http://abrc.osu.edu/</u>) using the kin7bifcfwd and kin7bifcrev primers (Table S2).
- 439 Amplified fragments were digested using SpeI and XhoI for KIN7 and KIN8 or BamHI and XhoI
- 440 for KIN12 and inserted in pSPYNE [20].

441 **Quantification of BiFC signals**

442 To be able to compare BiFC signals from various combinations, ImageJ software [31]) was used to 443 measure signal intensity from the vacuolar and plasma membranes. Values were corrected by 444 subtracting signal intensity from nearby background. Signal from TPK1_YFP_Nt+TPK1_YFP_Ct 445 (which forms a dimer) was used as positive control and the signal from 446 kin12_YFP_Nt+TPK1_YFP_Ct (which shows signal in ER and PM but not, or extremely little, in 447 the tonoplast) used as negative control. Quantitative data for BiFC fluorescence signal are shown in 448 Figure S2.

449 TPK1 pull-down assay

450 Pull-down assays were used to confirm the interaction between the N-terminus part of TPK1 and 451 KIN7. The sequence corresponding to the first 81 amino acids of TPK1 (NTPK1) was amplified by PCR using tpk1bamhIf and tpk1xhoir primers (Table S2) and cloned into the pGEX6P-1 vector 452 453 (GE Healthcare, Amersham, UK). GST and GST::NTPK1 expression was induced in 11 culture of 454 E. coli BL21 at 37°C with 1mM IPTG for 4 hours. Cells were collected and lysed in PBS-T buffer 455 (PBS pH 7.3, 0.1% Trion X-100) by sonication. After centrifugation, the lysate was cleared using 456 0.45 µm filters. GST or GST::NTPK1 was bound to 500 µl of Glutathione Sepharose 4B (GE 457 Healthcare, Amersham, UK) for 2h. 50 µl bead aliquots washed with buffer P (HEPES 50mM 458 pH7.3, 2mM CaCl2, 2mM MgCl2, 2mM KCl, 100mM NaCl, 1% CHAPS) and protease inhibitor 459 cocktail IV (Calbiochem, Merck, Feltham, UK) were incubated overnight with either proteins 460 extracted with buffer P from plants expressing wild type KIN7 or KIN7::YFP. The next day, beads 461 were washed 4 times with the same buffer. Bound proteins were eluted with SDS-PAGE Protein 462 Sample Buffer (2x) and loaded on a 10% acrylamide gel. Western blotting was performed as 463 described previously [32]. Rabbit anti-GFP (Thermofisher, Paisley, UK) and swine anti-rabbit 464 Immunoglobulins HRP (Agilent, Stockport, UK) were used and the resulting signal in the presence

of Luminata Forte substrate (Merck, Feltham, UK) was imaged with Fusion Pulse imaging system
(Vilber Lourmat, Marne-la-Vallée, France).

467 Cloning of KIN7:YFP and Arabidopsis plant transformation

468 KIN7 was PCR amplified from the clone U12357 (http://abrc.osu.edu/) with the kin7yfpfwd and 469 kin7yfprev primers (Table S2) and inserted in the pART7 vector [33]. The NotI fragment containing 470 the 35S:KIN7:YFP fragment was subsequently inserted in pGREEN0229. Arabidopsis plants were 471 transformed by floral spraying as described in [33]. Briefly, Agrobacterium was grown in 50 mL 472 liquid YEB medium for two days or until the OD reached 3. The cells were spun down (5 min, 473 4000g) and resuspended in 20 mL of 0.1x MS medium, 5% sugar, 0.1% Silwet L-77, pH 5.7. Every 474 two weeks, flowering plants were sprayed with Agrobacterium using an airbrush. For 475 pKin7::KIN7:YFP, 589bp upstream of the ATG was amplified from genomic DNA with 476 promkin7fwd and promkin7rev primer (Table S2). This Fragment was inserted in the 477 35S::KIN7::YFP vector instead of the 35S promoter, which was removed using StuI and XhoI.

478 **Protoplast isolation:**

479 Guard cell protoplasts were isolated as described by Pandey, et al. [34]. 40 fully expanded leaves 480 were blended in water for 1 minute using a waring SS515 Blender (Cole-Parmer UK) and poured 481 onto a 200 μ M mesh to collect epidermes. Epidermes were incubated in 45% H₂O, 55% basic 482 solution (0.5 mM CaCl₂, 0.5mM MgCl₂, 0.01 mM KH₂PO₄, 0.5 mM ascorbic acid, 550 mM 483 sorbitol, 0.2% BSA, 0.7% cellulysin, 5 mM MES/Tris pH 5.5) for 1.5 h at 30°C. The epidermes 484 were then incubated in basic solution supplemented with cellulase Onozuka RS 0.01% and 485 pectolyase Y23 for 1 h at 30°C. Guard cell protoplasts were collected after passing the solution 486 through a 20 µm mesh.

- 487 Mesophyll protoplast were extracted from Arabidopsis leaves according to {Isayenkov, 2011 #4}.
- 488 Leaves were cut in 1-mm sections and digested for 4 h in an enzyme solution containing 1.5%
- 489 cellulose RS, 0.75% macerozyme, 0.6 m mannitol, 10 mm 2-(N-morpholine)-ethanesulphonic acid
- 490 (MES), pH 5.6 and 1 mM CaCl₂, in which proteases were heat inactivated for 10 min at 55°C.
- 491 Protoplasts were filtered and centrifuged for 3 min at 500 g and resuspended in protoplast incubation
- 492 solution (0.6 M mannitol, 4 mM MES, pH 5.7, 4 mM KCl and 3 mm CaCl₂)

493 Electrophysiology

494 Vacuole release, equipment and analyses were as described in Maathuis, et al. [35]. After transfer 495 of protoplasts to the recording chamber, vacuoles were released by washing protoplasts with a

496 solution containing 10 mM EDTA, 10 mM EGTA, pH 8 with an osmolarity of 350 mOsM. Standard 497 experimental solutions for bath and pipette contained 100 mM KCl, 0.1 mM CaCl₂, 5 mM MES/Tris 498 pH 7 and sorbitol adjusted to 430 mOsm total osmolarity. Open probability (Po) was calculated as 499 described in Gobert, et al. [15] and defined as: Po = (t open/t total)/n where 'n' is an estimate of 500 the number of channels in the membrane patch derived from the maximum number of open levels 501 observed in the recording. Recordings of 60 s duration (t_total) at a membrane potential of -80 mV 502 were analysed by using a 50% threshold technique to define current transitions and calculate t_open 503 to determine Po. Open probability data were obtained from 3 to 10 individual protoplasts (see Figure 504 S2 and Data S1). To compare different genotypes, the increase in Po in response to ATP and 505 ATP+14-3-3 was calculated for each experiment and 'fold changes' were subsequently averaged 506 across experiments. ATP was added as Mg-ATP at a final concentration of 0.5 mM. 14-3-3 protein 507 (as Hv14-3-3B or Hv14-3-3C; GenBank accessions X93170 and Y14200 respectively [36] was a 508 kind gift from Bert de Boer (Vrije Universiteit, Amsterdam) and added to a final concentration of 509 $0.2\mu g/ml$.

510 Imaging

511 Intact and osmotically disrupted guard cells were photographed on a Zeiss epifluorescence 512 microscope using bright light, DIC or epifluorescence (465-495 nm excitation and 515 nm emission 513 wavelength) with a 20x, 40x or 63x objective. Confocal imaging was performed using a Zeiss 514 LSM510 Meta microscope (Carl Zeiss, http://www.zeiss.com).

515 Leaf Stomatal conductance and apertures

516 To record responses to ABA, leaves of mature plants were removed at approx 10:00h and incubated 517 in 'opening' buffer (10 mM KCl, 10 mM MES-KOH pH 6.15) for 2 h in the light to induce 518 maximum opening. Subsequently, H₂O gas exchange was determined (in the same buffer) using an 519 infrared gas analyser, Li-Cor 6400 (LI-COR, Cambridge, UK) to obtain a starting conductance 520 value. Subsequent values were obtained after 30 min incubation in control (no ABA) or ABA (1, 521 10 or 100 µM final concentration) buffer. Each experiment was carried out using three leaves and 522 in total 12-24 leaves from 3-4 individual plants were used for each treatment. Changes recorded in 523 control treatments (no ABA) were subtracted from those obtained from ABA treated leaves.

- 524 To obtain time courses for the response to elevated $[CO_2]$, stomatal conductance was measured
- 525 using infrared gas analysis. Measurements were performed using a portable photosynthesis system
- 526 attached to a leaf chamber with a 2.5 cm^2 leaf area (Walz GFS-3000). CO₂ was scrubbed from

527 external air using soda lime and resupplied from a liquid CO_2 cartridge to maintain CO_2 528 concentrations of either 400 or 1000 ppm. Temperature was maintained at 22°C and the absolute 529 humidity to 16000 ppm to obtain a relative humidity of 64.5%. Air flow was 400 µmol.s⁻¹, light intensity was 120 µmol.m⁻² s⁻¹. For each measurement, an individual mature leaf was placed in the 530 531 leaf chamber, while still attached to the plant. Leaves were left in the chamber for 30 min before 532 measurements were taken to allow them to acclimatise to chamber conditions and for gas exchange 533 to stabilise. Measurements were then logged every 10 secs and averaged every 2.5 min. Data 534 represent the mean +-SEM from 3 different plants.

535 To obtain stomatal apertures, leaf epidermes were removed from fully expanded leaves of 5 to 6 536 week old plants. They were collected cuticle-side up on CO₂-free 10 mM MES/KOH (pH 6.15) in 5cm Petri dishes (Sterilin, UK) at 22°C for 30min. Epidermal peels were transferred to fresh Petri 537 dishes and incubated in the light under a fluence rate of 120 µmol.m⁻². s⁻¹ in 50mM KCl, 10 mM 538 539 MES/KOH (pH 6.15) at 22°C 2 hours whilst being aerated with an air stream containing 400 ppm 540 [CO₂] from a pressurised cylinder (BOC, Special Gasses, UK) by bubbling directly into the buffer. 541 Subsequently, peels were either aerated with 1000ppm [CO₂] or continued to be aerated with 542 400ppm [CO₂]. After 3h, peels were removed, mounted on slides and measurements of stomatal 543 aperture were recorded using an inverted microscope (Leica DM-IRB, Leica UK). Forty stomatal 544 pores were measured per treatment in three separate replicated experiments (total stomatal number 545 = 120; n = 3). To avoid bias, experiments were performed without knowing the identity of the plants 546 and the treatments until data were collected. Data were analysed using one-way ANOVA.

547 Immunoblotting

548 Guard cell protoplast preparations were visually inspected and only used when containing fewer 549 than 1% mesophyll protoplasts. ABA treatment (40 µM, 1 hour) was either done on intact leaves in buffer (containing 20 mM KCl, 10 mM MES/KOH pH 6.1) at ~150 µmol.s⁻¹.m⁻² light, or on isolated 550 551 protoplasts in buffer (containing 500 mM Sorbitol, 10 mM KCl, 1 mM CaCl₂, 10 mM MES/Tris 552 pH 5.5). In the latter case, protoplasts were isolated from *tpk1* mutants, and transiently transformed 553 with pART7:TPK1:YFP [15]. Around 16h after transformation, protoplasts were divided in two 554 population with one exposed to ABA (conditions as above). After treatment, protoplasts were resuspended in 'RIPA' buffer (1% Triton X 100, 0.1% SDS, 100 mM NaCl, 10 mM Na₂HPO₄ and 555 556 10 mM NaH₂PO₄, pH 7.2 plus protease inhibitor cocktail 'cOmplete EDTA free' from Roche UK) 557 and stored at -20°C.

558 For analysis of phosphorylation levels, protein samples $(40 \mu g)$ were precipitated by methanol-559 chloroform extraction, separated by SDS-10% acrylamide gel electrophoresis and transferred onto 560 nitrocellulose (Hybond-ECL, GE Healthcare). Prior to blocking (5% BSA, TBS), protein levels 561 were visualised with Ponceau S (0.1% w/v in 1% Acetic acid). TPK1 phosphorylation was detected 562 by immunoblotting overnight with anti-pBAD-Ser136 (1:200, i.e. 75 µL in 15 ml, rabbit polyclonal 563 serum, Santa Cruz sc-12969; secondary antibody IRDye 800CW Donkey anti-rabbit, LI-COR 926-564 32213) and visualised by infrared fluorescence (ODYSSEY CLx, LI-COR). Average 565 phosphorylation signal was calculated using densitometry (ImageJ v 1.48).

566 Isolation of tonoplasts and Western blotting

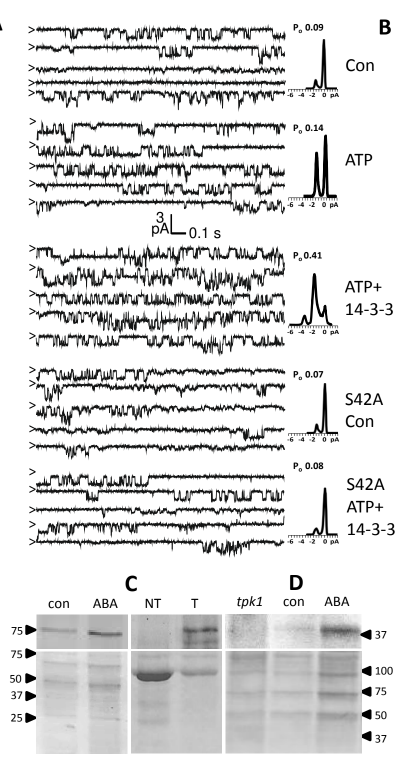
567 Five plants at the rosette stage were coated using a paint brush with tween 0.05%, cycloheximide 568 (CHX) 50 µM or chlorpromazine (CPZ) 50 µM in tween 0.05%, one hour before being sprayed 569 with ABA (100 µM) or with a mock solution (tween 0.05%). One hour after ABA treatment, the 570 samples were frozen in liquid nitrogen and ground using a mortar and a pestle. Microsomes were 571 extracted according to Lund and Fuglsang [37] with an altered homogenisation buffer according to 572 Rea, et al. [38]. Briefly, ground material was extracted in 1.1 M glycerol, 5 mM Tris-EDTA, 5 mM 573 DTT, 1% [w/v] PVP-40, 1 mM PMSF, 70 mM Tris-Mes [pH 8.0]) and centrifuged at 6,000g for 10 574 min. The pellet was resuspended in 3.5 mL buffer 330/5 (0.33 M sucrose, 5 mM potassium 575 phosphate pH 7.8) using a glass homogenizer. Three grams of microsomal fraction was added on top of 9 g polymer solution (3.72 g of 20% Dextran T500 solution, 1.86 g of 40% PEG4000 solution, 576 577 1.08 g sucrose, 225 µL potassium phosphate buffer 0.2 M pH 7.8, 18 µL KCl 2 M, H₂O up to 9 g). 578 In parallel, a blank tube was made with 3 g of buffer 330/5 that was added on top of the polymer 579 solution. The tubes were gently inverted 12 times and centrifuged at 1,000g for 5 min. The lower 580 phase enriched in tonoplast was collected and re-purified using the top phase of the blank tube. The 581 bottom phase was collected and diluted 10 times with buffer 300/5. The tonoplasts were centrifuged 582 at 100,000g for 60 min. The lower phase enriched in tonoplast was resuspended in 0.1 ml RIPA 583 buffer (Fisher scientific, Nottingham, UK) and loaded onto a 4-12% SDS-PAGE NuPAGE™ 584 (Thermofisher, UK). The western blotting was performed according to Pijacka, et al. [32]. Briefly, 585 an Amersham ECL Plex Western blotting system was used on which the gel was run for 2 hours at 586 120V and protein was subsequently transferred onto a low-fluorescent PVDF membrane (GE 587 Healthcare, Buckinghamshire, UK) using NuPAGE Transfer Buffer (Thermofisher, UK) for 1 hour 588 at 30V in a mini gel tank (Thermofisher, UK). The membrane was cut horizontally in two parts,

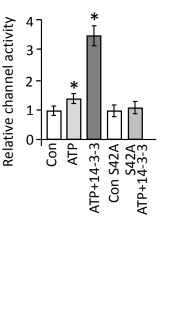
with the upper part incubated overnight at 4°C with anti-GFP antibody (at 1:2000 dilution) and the bottom part with anti TIP1-1 antibody (at 1:5000 dilution), ensuring equal protein ratio between lanes. Fluorescent secondary antibodies were incubated for one hour and the blots were imaged using fluorescent laser scanner (Typhoon, GE Healthcare, Buckinghamshire, UK). Quantification of bands was performed using CLIQS software (Totallab, Newcastle upon Tyne, UK).

- 594 To determine change in tonoplast expression in response to ABA or CO₂, plants were treated for up
- 595 to 2h with 50 µM ABA or 1000 ppm CO₂. Immunoblotting was carried out as described above and
- relative expression levels were determined using TIP1;1 as marker and ImageJ [31] software to
- 597 quantify band intensities.

598 QUANTIFICATION AND STATISTICAL ANALYSIS

- 599 Where statistical testing of data was applied, it is indicated in the legend of the respective figure, as
- 600 is the number ('n') of experimental replicates. For ANOVA analyses, Prism 6 (Graphpad software)
- 601 was used. If ANOVA revealed significant (p < 0.05) effect of group, post hoc test used to determine
- 602 p values for all relevant comparisons is mentioned in the figure legends.
- Data S1 'Patch clamp open probabilities' (related to Figures 1E and S2A). The spreadsheet
 shows open probabilities for each kinase genotype in control, plus ATP or plus ATP+14-3-3
 conditions.

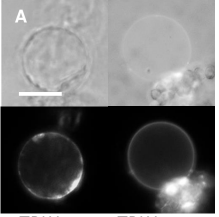




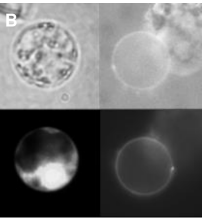
Phosphorylation WT_Con WT_ABA_L WT_ABA_L WT_ABA_L WT_ABA_L *w*T_ABA_L *kin7-1_*ABA_L *kin7-2_*ABA_L

Figure1

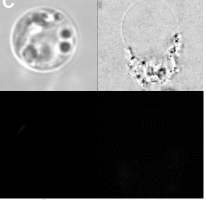
Figure2



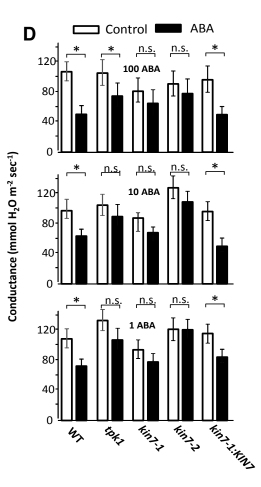
TPK1_{YFP-Nt}+ TPK1_{YFP-Ct}

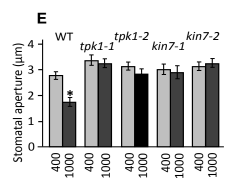


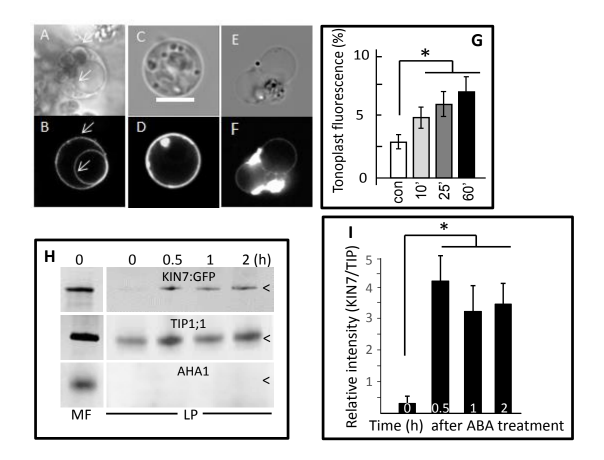
KIN7_{YFP-Nt}+ TPK1_{YFP-Ct}

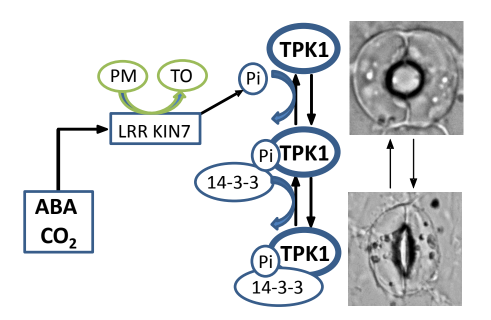


KIN8_{YFP-Nt}+ TPK1_{YFP-Ct}









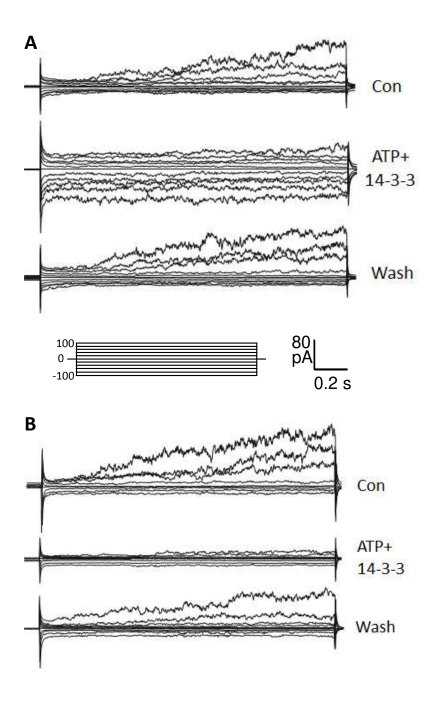


Figure S1: Whole vacuole currents (relating to Figure 1).

(A), WT guard cell whole vacuole currents in standard buffer, after exposure to buffer containing ATP+14-3-3 and after washout of ATP and 14-3-3. (B) Same protocol as shown in (A) for a vacuole that expresses TPK1 with the S42A mutation. Note voltage dependent activation of the SV-type current at positive voltages in control conditions and its inhibition by 14-3-3. Also note absence of TPK1 current activation in the S42A mutant. Inset shows applied voltage clamp protocol (-100 to 100 mV).

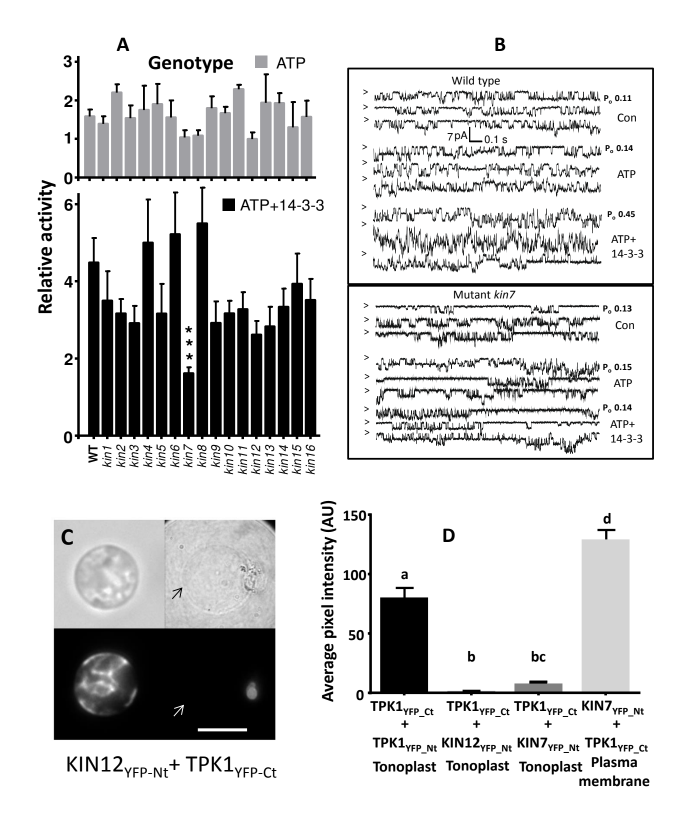


Figure S2: Kin mutant currents and BiFC (relating to Figure 2).

(A) Vacuolar cytoplasmic side out patches (n=5 to 10), isolated from wild type plants and from 16 different kinase loss of function mutants were recorded in control buffer, in the presence of MgATP, and MgATP plus 14-3-3 protein. Channel activity in the presence of ATP (or ATP+14-3-3) was compared to that measured in the control solution. The ratios of activities between ATP (or ATP+14-3-3) and control solution were then used for analysis. Data show the average for each genotype +/- SEM. The *kin7* genotype stimulatory effect of 14-3-3 (bottom panel) was significantly lowered compared to the WT. Data were tested using

a one-way ANOVA with Dunnett post hoc test analysis ***p<0.001. (B) Example traces that reflect the different levels of current stimulation by ATP and by ATP+14-3-3 for wild type, and *kin7* KO mutants. (C) BiFC fluorescence in guard cell cotransformed with KIN12_{YFP-Nt} and TPK1_{YFP-CT}. Top two panels show DIC images of intact and osmotically ruptured protoplast. Bottom panels show corresponding fluorescence signal. Scale bar is 10 μ m. (D) Quantification of BiFC fluorescence signals. Data were analysed using one way ANOVA with Tukey post-hoc analysis, different letters denoting p<0.05.

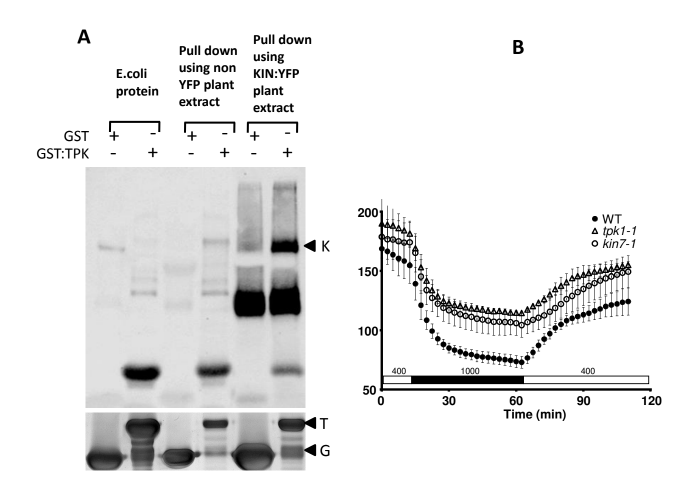


Figure S3: Pull-down assays and stomatal conductance in response to elevated CO_2 (relating to Figure 2).

(A) Pull down assays were carried out using the N-terminal part of TPK1 fused to GST as bait (or GST alone) and leaf extract from wild type plants or plants expressing KIN7::YFP as bait. A primary anti-GFP antibody and HRP-coupled secondary antibody were used to visualise bands on the blot. Arrow head on the top panel (K) depicts 100 kDa (expected KIN7::YFP size ~95 kDa), bottom panel G for GST (~26 kDa) and T for GST:TPK1 (~37 kDa). The top panel represents immunoblot using anti-GFP antibody and the bottom panel represents the corresponding gel stained using Coomassie. (B) Leaf conductance time courses (n=3 leaves per genotype) in response to changing ambient CO₂ which is 400 ppm for the first 10 minutes, raised to 1000 ppm for one hour and brought back to 400 ppm for another hour. Note the unresponsiveness for both the *tpk1* and *kin7* loss of function mutants.

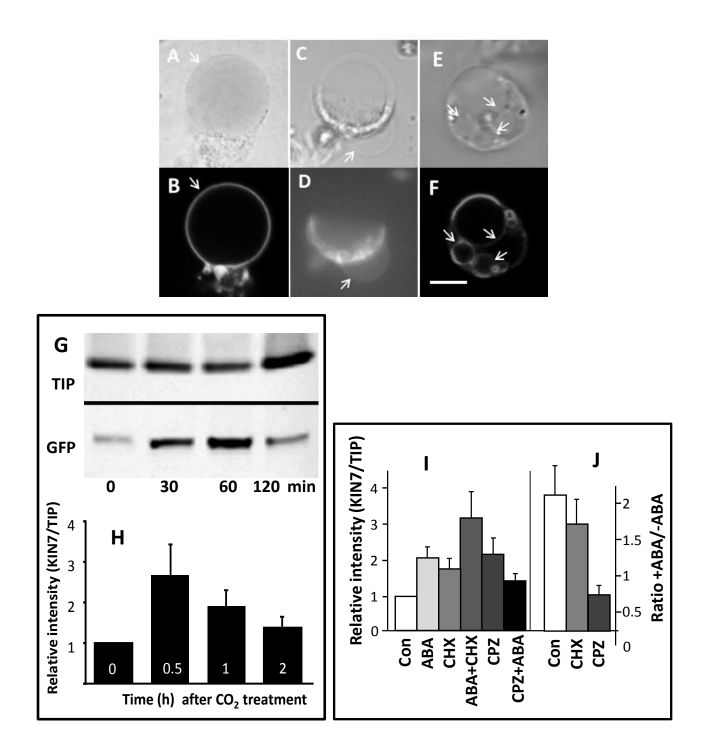


Figure S4: KIN7-YFP membrane expression pattern, CO_2 induced shift in expression and the effect of protein synthesis and endocytosis inhibitors (relating to Figure 3). (A, B) Arabidopsis guard cells transiently transformed with 35S:KIN7-YFP show KIN7 expression in tonoplast (white arrows). (C, D) Partially disrupted guard cell protoplast transiently transformed with Kin7p:KIN7-YFP shows fluorescence in large vacuolar bulge (arrows). (E, F), Guard cell protoplasts isolated from plants stably transformed with Kin7p:KIN7-YFP shows fluorescence large and multiple small vacuoles (arrows). Top row shows DIC images, bottom row shows corresponding epifluorescence images. Scale bar is 3 µm. (G) Example Western blot using antibodies against KIN7-GFP and TIP1;1, showing that CO_2 treatment (1000 ppm, 3h) causes an increase in KIN7 expression relative to TIP1;1, a marker for the tonoplast. (H) Data were quantified on the basis of 3 independent biological replicates, error bars denote standard errors. (I) Treatment of plants with ABA, cycloheximide (CHX, a protein synthesis inhibitor) or chlorpromazine (CPZ, an inhibitor of endocytosis) on their own, causes a shift in KIN7 expression towards tonoplast location. Cycloheximide and chlorpromazine stimulate an increased KIN7:TIP ratio when used on their own, pointing to non-specific effects of these inhibitors. The effect of ABA is abrogated in plants pre-treated with chlorpromazine. This becomes obvious when plus/minus ABA ratios are compared for the various treatments (Figure 3B). (J) Same data as in (A) but expressed as a ratio of plus ABA to minus ABA treatment. This shows that cycloheximide does not affect the ABA-dependent shift toward tonoplast expression whereas chlorpromazine strongly inhibits it. Data are from 3 independent experiments and error bars denote standard errors.

KIN# Gene ID Annotation Mutant line TM				
1	AT1G21250	WAK1, wall associated kinase	SALK_209668	1
2	AT1G48480	RKL1 (Receptor-like kinase 1)	SALK_099094	2
3	AT1G51805	Leucine-rich repeat protein kinase,	SALK_003231	1
4	AT1G53730	SRF6 (Strubbelig-receptor family 6)	SALK_054337	2
5	AT1G66150	TMK1 (Transmembrane kinase 1)	SALK_008771	1
6	AT2G26730	Leucine-rich repeat transmembrane protein kinase	SALK_034004	1
7	AT3G02880	Leucine-rich repeat transmembrane protein kinase	SALK_019840/FLAG_321B	08 1
8	AT3G14840	Leucine-rich repeat family protein	SALK_094512	2
9	AT3G22060	Receptor protein kinase-related	SALK_151902	1
10	AT4G01330	Protein kinase family protein	SALK_081284	1
11	AT4G21380	ARK3 (Arabidopsis Receptor Kinase 3)	SALK_001986	2
12	AT4G21410	Protein kinase family protein (CRK29)	SAIL_447_F06	1
13	AT4G29130	AtHXK1 (GLUCOSE INSENSITIVE 2) hexokinase	SALK_018086	1
14	AT4G34220	Leucine-rich repeat transmembrane protein kinase	SALK_112336	2
15	AT5G38990	Protein kinase family protein	SALK_139579	1
16	AT5G49760	Leucine-rich repeat family protein kinase	SALK_118908	2

Table S1: Vacuolar kinases (related to Figure 1, Table S2 and Figure S2).

Membrane kinases identified in the SUBA database (suba2.plantenergy.uwa.edu.au/) with a 'vacuolar' localisation based on either 'GFP' or MS/MS analyses. 'TMD': putative number of transmembrane domains. For each, loss of function mutants were obtained which were genotyped using T-DNA and gene specific primers (see Table S2). RT-PCR on RNA from homozygous lines was carried out, using gene specific primers, to ensure no transcript was present.

Mutant Line	Forward and Reverse genotyping primers
SALK 209668	5'AATCTGCGAAATGTCATGAGG3'
0/1211_200000	5'TTGGTATCAGCCTTGAAGCAC3'
SALK 099094	5'TCTCTGTTTTCCTCTCCCCCC3'
	5'GTCAAGCACTGCCTTATACGC3'
SALK 003231	5'TTCTTACATGACAAAAATTCAGGG3'
	5'AGCTCAAGATCAACCCGGTAC3'
SALK 054337	5'AACGACTTTCACGGTATGCAC3'
—	5'TGTCAAATGGTTTTCTCCCAG3'
SALK 008771	5'CGAGAAAACCGGGTAAAGAAC3'
—	5'TTTTTGTGCGAATAAACAACTTG3'
SALK_034004	5'GCCCACCAGATAAATATTTGC3'
	5'AGTGAGTCGACCTAAGGAGCC3'
SALK_019840	5'CATGAAAACAAAGGGATGAGG3'
	5'CACGCAGAGGATATCGAGAAG3'
GABI_047F11	5'TAAATGGTTTAAGCGGTGTGC3'
	5'ATCAGCAACGGACATTTCAAC3'
FLAG_321B08	5'TATTCCGAGTTCGTTGTCGTC3'
	5'GGAAACGGGAGAACTCCGTT3'
SALK_094512	5'CATCGATCATGCCATAAATCC3'
	5'GAGCAGTTACAAGTAACGGCG3'
SALK_151902	5'TTTACCGTCGCAACAGTTAGG3'
	5'TTAAAACGCATCGTTTGGTTC3'
SALK_081284	5'TTCGTCTCGATATGGACAAGG3'
	5'GACGACCCAAAGCTTTAGACC3'
SALK_001986	5'TCAAAGACATCAGTTCAGGGG3'
0.411 4.47500	5'TTCACTGTCCATGATTCATCG3'
SAIL_447F06	
	5'AAATACAGCAGGAGGGATGTG3'
SALK_018086	5'CTCGAAACTGAGACAAGTGGC3' 5'GTCTTCGCATTCTGTAGCGAC3'
SALK_112336	5'CTCGAAACTGAGACAAGTGGC3'
SALK_112550	5'GTCTTCGCATTCTGTAGCGAC3'
SALK 139579	5'CTCTCGAATTCCTTGGGTTTC3'
OALIN_100010	5'CAATCTCTCTGGTGATTTGCC3'
SALK 118908	5'TCAATGGACGTAACTTTGAGG3'
OALN_110000	5'AGTGAAACGGTTGACGTTGAC3'
	Cloning primers
kin8bifcfwd	CCACTAGTGGAACAATGTCGTTAAATCGACAACTTCT
kin8bifcrev	CGACTCGAGAGTTCTAGTGTTCCAATAGG
kin12bifcfwd	CCGGATCCGGAACAATGGAACATGTCAGAGTTATCT
kin12bifcrev	CGACTCGAGACGAGGAGAAAACTCAGAAA
kin7bifcfwd	CCACTAGTGGAACAATGAAGTATAAGCGTAAGCT
kin7bifcrev	CGACTCGAGGTCGGATACAGGATTTGGGG
tpk1bamhlf	GGATCCATGTCGAGTGATGCAGCT
tpk1xhoir	CTCGAGCTACATGATCACTCGCCTGAG
kin7yfpfwd	CGACTCGAGAACAATGAAGTATAAGCGTAAGCT
promkin7fwd	ATGTCTTTGGTCACACTCAACCTGC
promkin7rev	CACTCGAGCTTCTTCTTCTTCTTCAAAAAC
kin7yfpfwd	CGACTCGAGAACAATGAAGTATAAGCGTAAGCT
kin7yfprev	ACCCGGGGTCGGATACAGGATTTGGGG

Table S2: Genotyping and cloning primers (related to Star Method section 'Plants' and 'Cloning of kinases in the BiFC vector').

Gene specific primers used to identify homozygous lines for kinase mutant lines listed in Table S1 and cloning primers used to make kinase-YFP fusion proteins.