1	SN	IF1-related protein kinase 2 directly regulate group C Raf-like protein
2	kir	nases in abscisic acid signaling
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# 30 **Distribution statement**

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34

#### 35 ABSTRUCT

A phytohormone abscisic acid (ABA) has a major role in abiotic stress responses in plants, and subclass III SNF1-related protein kinase 2 (SnRK2) mediates ABA signaling. In this study, we identified Raf36, a group C Raf-like protein kinase in Arabidopsis, as an interacting protein with SnRK2. A series of reverse genetic and biochemical analyses revealed that Raf36 negatively regulates ABA responses and is directly phosphorylated by SnRK2s. In addition, we found that Raf22, another C-type Raf-like kinase, functions

partially redundantly with Raf36 to regulate ABA responses. Comparative
 phosphoproteomic analysis using Arabidopsis wild-type and *raf22raf36-1* plants identified proteins that are phosphorylated downstream of Raf36 and
 Raf22 *in planta*. Together, these results reveal a novel subsection of
 ABA-responsive phosphosignaling pathways branching from SnRK2.

48

### 49 **INTRODUCTION**

50 Environmental stresses, such as drought, high salinity and low temperature, have adverse effects on plant growth and development. Abscisic acid (ABA) is a 51 52 phytohormone that plays important roles in responses and adaptations to these 53 stresses, as well as in embryo maturation and seed dormancy (Finkelstein, 2013; Shinozaki et al., 2003). The major ABA signaling pathway consists of three core 54 55 components: ABA receptors, type 2C protein phosphatases (PP2Cs) and SNF1-related protein kinase 2s (SnRK2s) (Cutler et al., 2010; Umezawa et al., 56 57 2010). In this pathway, SnRK2s transmit ABA- or osmostress induced-signals thorough phosphorylation of downstream substrates, thereby promoting ABA- or 58 59 stress-inducible gene expression and stomatal closure (Furihata et al., 2006; Geiger et al., 2009; Umezawa et al., 2013; Wang et al., 2013). The Arabidopsis 60 61 genome contains 10 members of SnRK2, and they are classified into three subclasses (Hrabak et al., 2003; Yoshida et al., 2002). Among them, subclass III 62 members, SRK2D/SnRK2.2, SRK2E/OST1/SnRK2.6 and SRK2I/SnRK2.3, are 63

essential for ABA responses (Fujii and Zhu, 2009; Fujita et al., 2009; Nakashima et
al., 2009; Umezawa et al., 2009).

Raf-like protein kinases were recently identified as regulators of ABA 66 signaling. Among 80 putative MAPKKKs in Arabidopsis, 48 members are 67 68 categorized as Raf-like subfamilies and can further be divided into 11 subgroups 69 (B1- B4 and C1- C7) (Ichimura et al., 2002). In *Physcomitrella patens*, the ARK 70 (also named ANR or CTR1L) gene is required for ABA-responsive 71 SnRK2-activation, gene expression and drought, osmotic and freezing tolerance 72 (Saruhashi et al., 2015; Stevenson et al., 2016; Yasumura et al., 2015). ARK 73 encodes a B3 subgroup Raf-like protein kinase that phosphorylates SnRK2s in vitro, 74 suggesting that ARK functions as a upstream kinase of SnRK2s (Saruhashi et al., 2015). In Arabidopsis, the B2 subgroup kinases Raf10 and Raf11 positively regulate 75 76 seed dormancy and ABA responses by directly phosphorylating and activating 77 subclass III SnRK2s (Lee et al., 2015; Nguyen et al., 2019). In addition to group B 78 kinases, several group C Raf-like kinases have been associated with ABA responses. For example, Arabidopsis Raf43, a C5 kinase, promotes ABA sensitivity 79 80 during seed germination and seedling root growth (Virk et al., 2015), whereas Raf22, a member of C6 subgroup, negatively regulates stress- or ABA-induced growth 81 82 arrest (Hwang et al., 2018). However, despite these ABA-related genetic phenotypes, it is still unclear whether group C kinases directly regulate 83 84 SnRK2-dependent signaling pathways.

In this study, we identified Raf36, a C5 group Raf-like kinase, as a protein 85 86 that directly interacts with, and is phosphorylated by, SnRK2. Our evidence 87 indicates that Raf36 functions as a negative regulator of ABA signaling pathway 88 during post-germinative growth stage under the control of SnRK2. In addition, we revealed that Raf22, a C6 Raf-like kinase, functions partially redundantly with Raf36. 89 Comparative phosphoproteomic analysis revealed that Raf36 and Raf22 are 90 91 required for a subset of ABA-responsive phosphosignaling pathways. Collectively, 92 unlike group B Rafs, which have been recently reported as an "accelerator" of ABA 93 response upstream of SnRK2s, our results demonstrate that Arabidopsis group C 94 Rafs, Raf22 and Raf36, functions as a "brake" of ABA response downstream of 95 SnRK2s.

96

#### 98 **RESULTS**

# 99 Raf36 interacts with subclass III SnRK2

To identify additional kinases that regulate ABA signaling pathways, we 100 101 used the AlphaScreen<sup>®</sup> assay to screen a collection of Arabidopsis MAPKKK 102 proteins for their ability to physically interact with SRK2I (SnRK2.3), a subclass III 103 SnRK2. From a pilot experiment, several Raf-like protein kinases were identified as 104 candidate interactors with SRK2I (Supplemental Figure 1). Raf36, which belongs to 105 a C5 subgroup kinase (Supplemental Figure 2), was one of the SRK2I-interacting 106 proteins. Interaction between Raf36 and SRK2I, as well as between Raf36 and additional subclass III SnRK2s, SRK2D (SnRK2.2) and SRK2E (OST1/SnRK2.6), 107 was confirmed by AlphaScreen<sup>®</sup> assay (Figure 1A) and yeast two-hybrid assay 108 (Figure 1B). SnRK2s were previously found within the cytosol and nuclei of 109 110 Arabidopsis cells (Umezawa et al., 2009). We observed that Raf36-GFP is localized 111 mainly in the cytosol (Figure 1C), and bimolecular fluorescence complementation 112 (BiFC) assay confirmed that the interactions between SnRK2s and Raf36 take 113 place in cytosol (Figure 1D) Together, these results demonstrate that Raf36 114 physically interacts with ABA-responsive SnRK2s both in vitro and in vivo.

115 Next, we investigated which domain(s) of Raf36 may be responsible for the 116 interaction with SnRK2s. According to the PROSITE database 117 (https://prosite.expasy.org/), Raf36 contains an unknown N-terminal stretch (N, 1-118 206 aa), a predicted kinase catalytic domain (KD, 207- 467 aa) and a short

119 C-terminal domain (C, 468- 525 aa) (Figure 1E). In yeast two-hybrid assay, SRK2E 120 strongly interacted with Raf36 full-length protein (FL), but just slightly or not 121 interacted with Raf36 N and KD+C alone, respectively (Figure 1E). These results 122 indicated that the complete structure of Raf36 protein is required for the interaction 123 with SnRK2.

124

# 125 **Raf36 negatively regulates ABA response at post-germination growth stage**

126 To characterize the role of Raf36 in ABA signaling, we performed a series of 127 functional analyses in Arabidopsis. Using gRT-PCR, we measured the abundance 128 of *Raf36* mRNA in seedlings and detected a slight yet significant increase in *Raf36* 129 transcripts after ABA treatment (Figure 2A). Next, we obtained two Raf36 T-DNA insertion lines, GK-459C10 and SALK 044426C, designated as raf36-1 and raf36-2, 130 131 respectively (Supplemental Figure 3A). Using RT-PCR we confirmed the loss of 132 *Raf36* transcripts in both mutants (Supplemental Figure 3B). We then measured 133 rates of seed germination and cotyledon greening in the presence or absence of 134 exogenous ABA. No difference in greening rate was observed between wild-type 135 and mutant seedlings in the absence of ABA. However, with plants treated with 0.5 136 µM ABA, the greening rate of *raf36* mutants was significantly slower than wild-type 137 (Figures 2B and 2C). This ABA-hypersensitive phenotype was complemented by 138 CaMV35S:Raf36-GFP (Figure 2D and Supplemental Figure 3C). To assess if the 139 delayed greening of raf36 requires SnRK2 signaling, we generated a triple mutant,

*raf36-1srk2dsrk2e*, and observed that it was less sensitive to ABA than *raf36-1*(Figures 2E and 2F). This result indicates that SRK2D and SRK2E are genetic
modifiers of *raf36*-dependent ABA hyper-sensitivity in the greening response.
Notably, seed germination rates were not significantly changed in either *raf36*mutant in the presence or absence of ABA (Supplemental Figure 3D). Taken
together, our results suggested that Raf36 functions as a negative regulator of
SnRK2-dependent ABA signaling during post-germinative growth.

147

# 148 **Raf36 is phosphorylated by SnRK2**

149 To further examine the biochemical relationship between SnRK2 and Raf36, we prepared Raf36 and SRK2E recombinant proteins as maltose-binding protein 150 151 (MBP)or glutathione S-transferase (GST)-fusions. Raf36 protein 152 auto-phosphorylated, demonstrating that Raf36 is an active kinase (Figure 3A). We found that Raf36 prefers Mn<sup>2+</sup> for its kinase activity (Supplemental Figure 4A), as 153 154 shown for other C-group Raf (Lamberti et al., 2011; Reddy and Rajasekharan, 2006: Rudrabhatla et al., 2006), and SRK2E prefers Mg<sup>2+</sup> for its kinase activity 155 156 (Supplemental Figure 4B). We then performed *in vitro* phosphorylation assays using kinase-dead forms of Raf36 and SRK2E as substrates. SRK2E 157 158 phosphorylated Raf36 (K234N), while Raf36 did not phosphorylate SRK2E (K50N) (Figure 3A). This result suggested that Raf36 is a potential substrate of SnRK2, but 159 160 not vice versa.

161 Additional *in vitro* kinase assays were performed using a series of truncated 162 versions of Raf36 to identify the phosphorylation site(s) of Raf36. First, we observed that Raf36 proteins lacking the N-terminal region (Raf 36 KD+C and Raf36 KD) 163 164 were not phosphorylated, suggesting this region is important for both auto-phosphorylation by Raf36 and trans-phosphorylation by SRK2E (Figure 3B). 165 166 Second, Raf36 N (1-206) and Raf36 N (1-156) recombinant proteins were strongly 167 phosphorylated, but Raf36 N (1-140) was slightly phosphorylated by SRK2E 168 (Supplemental Figure 4C). These data suggested that the major phosphorylation 169 site is located within 141-156 aa of the N-terminal region. Four serine (Ser) residues (i.e., Ser<sup>141</sup>, Ser<sup>145</sup>, Ser<sup>150</sup> and Ser<sup>155</sup>) are present within this region. To identify 170 171 which Ser residue(s) may be phosphorylated, we generated six peptides spanning 172 this region of Raf36 (134-163). The peptides either contained all four serine 173 residues (peptide #1), or had only a single Ser residue, with alanine substituted for the remaining Ser residues (peptides #2- #6) (Figure 3C). Ser<sup>157</sup> was replaced with 174 175 alanine in each peptide because it was outside of phosphorylated 141-156 aa 176 region. Of these synthetic peptides, only peptides #1 and #3 were strongly phosphorylated by SRK2E, indicating Ser<sup>145</sup> is the phosphorylation site (Figure 3D). 177 SRK2D and SRK2I also phosphorylated Ser<sup>145</sup> of Raf36 *in vitro* (Figure 3E). Taken 178 together, these data show that subclass III SnRK2s phosphorylate Ser<sup>145</sup> of Raf36. 179 180

#### 181 **Raf22 functions partially redundantly with Raf36**

182 We next tested if Raf kinases closely-related to Raf36 are also SnRK2 183 substrates. There are five kinases within Raf subgroups C5 and C6 (Figure 4A). 184 Among them, HIGH LEAF TEMPERATURE 1 (HT1/ Raf19) functions independently 185 of ABA (Hashimoto et al., 2006; Hashimoto-Sugimoto et al., 2016). Therefore, we 186 focused our analyses on Raf43, Raf22 and Raf28. As shown in Figure 4B, SRK2E strongly phosphorylated Raf22, despite having no equivalent of Ser<sup>145</sup> of Raf36 and 187 sharing only 29 % identity with Raf36. In addition, SRK2E only weakly or not 188 189 phosphorylated Raf43 and Raf28, respectively. We next tried to identify the 190 phosphorylation site in Raf22. As described above, Raf28 was not phosphorylated by SnRK2, albeit with having 88 % identity with Raf22. Because SnRK2 kinases 191 prefer [-(R/K)-x-x-(S/T)-] or [-(S/T)-x-x-x-(E/D)-] (Furihata et al., 2006; Umezawa 192 et al., 2013), we searched the amino acid sequences of Raf22 and Raf28 for 193 potential SnRK2 phosphorylation sites, and identified Ser<sup>81</sup> within an [-R-H-Y-S-] 194 195 motif in Raf22 that is converted to [-R-H-P-Y-S-] in Raf28. We introduced an alanine substitution at Ser<sup>81</sup> in recombinant Raf22, and observed that this substitution 196 197 nearly abolished phosphorylation by SRK2E (Figure 4C), indicating this serine 198 residue is a SnRK2-phosphorylation site.

199 Next, we functionally characterized the role of Raf22 in ABA-related 200 phenotypes. Similar to *Raf36*, transcriptional level of *Raf22* was slightly 201 up-regulated after exogenous ABA treatment (Supplemental Figure 5A). Using BiFC 202 assay, we observed interaction between Raf22 and SnRK2s *in vivo* (Figure 4D and

Supplemental Figure 5B). A T-DNA insertional mutant (SALK\_105195C), *raf22*, showed a similar phenotype to *raf36*, i.e. ABA hypersensitivity in the post-germination growth (Figure 4E and 4F) but not in seed germination (Supplemental Figure 5C).

To test potential functional redundancy between Raf36 and Raf22, a 207 208 raf22raf36-1 double knockout mutant was generated. In the presence of ABA, 209 raf22raf36-1 showed a stronger ABA-hypersensitive phenotype relative to individual 210 *raf22* and *raf36-1* mutants (Figures 4E and 4F). In addition, expression of ABA- and 211 stress-responsive genes RD29B and RAB18 were hyper-induced in raf22raf36-1 212 seedlings (Figures 4G and 4H). Moreover, leaf water loss was examined because 213 ABA also controls stomatal closure. However, leaf water loss of raf22raf36-1 and 214 individual raf22 and raf36-1 plants was similar to that of wild-type plants 215 (Supplemental Figure 5D), suggesting that Raf36 and/or Raf22 has a minor role in 216 stomatal movements. Taken together, these results demonstrated that Raf36 and 217 Raf22 function redundantly in ABA signaling during post-germinative growth stage.

To examine whether the protein kinase activity of Raf36 and Raf22 are required for its negative regulation of ABA response, several complemented lines with kinase-dead form were generated. As shown in Figure 5A, the wild-type Raf36 complemented the ABA-hypersensitive phenotype of *raf36-1*, while the kinase-dead form of Raf36 (Raf36 K234N) could not. Similar results were also observed for *Raf22* (Supplemental Figure 6). These results suggested that the protein kinase

activities of Raf36 and Raf22 are required for its function in ABA signaling.

225

# 226 Raf36 and Raf22 regulate a subset of protein phosphorylation network in ABA

227 response

228 To gain insight about ABA-responsive signaling pathway(s) that may be regulated by Raf36 and/or Raf22, we performed a comparative phosphoproteomic 229 230 analysis of wild-type and raf22raf36-1 seedlings treated with 50 µM ABA for 0, 15, 231 30 and 90 min. LC-MS/MS analysis identified a total of 1,500 phosphopeptides in 232 both wild-type and the *raf22raf36-1* double mutant (Supplemental Dataset 1). Within this dataset 99% of identified phosphopeptides were singly phosphorylated 233 234 7A). (Supplemental Figure Phosphoserine, phosphothreonine, and phosphotyrosine accounted for 91.2%, 8.4% and 0.3% of phosphorylated residues, 235 236 respectively (Supplemental Figure 7B). Consistent with the ABA-hypersensitive 237 phenotype observed in the raf22raf36-1 double mutant (Figures 4E and 4F), 238 principal component analysis demonstrated that the phosphoproteome profiles of 239 the ABA-treated raf22raf36-1 mutant were different from that of ABA-treated 240 wild-type (Figure 5B). In addition, the profiles of raf22raf36-1 were significantly 241 altered even before ABA treatment (Figure 5B).

242 Next, we identified phosphopeptides differentially regulated between 243 wild-type and *raf22raf36-1*. ABA-responsive and Raf36 and/or Raf22 dependent 244 phosphopeptides were screened by *P* value. After exogenous ABA treatment, 130

245 phosphopeptides were upregulated and 36 phosphopeptides were downregulated 246 in wild-type plants (Figure 5C and Supplemental Dataset 2). In comparison with 247 wild-type at each time point, a total of 416 and 175 phosphopeptides were 248 upregulated and downregulated in raf22raf36-1, respectively (Figure 5C and Supplemental Dataset 2). Intriguingly, 53.8 % of ABA-upregulated phosphopeptides 249 250 in wild-type were up- or down-regulated in *raf22raf36-1* double mutant, while 61.1 % 251 of ABA-downregulated phosphopeptides in wild-type were up- or down-regulated in 252 raf22raf36-1, indicating that large part of ABA-regulated phosphopeptides may be 253 under the control of Raf36 and/or Raf22.

254 Subsets of phosphopeptides were further analyzed to evaluate potential differences in cellular responses to ABA between wild-type and raf22raf36-1 plants. 255 256 First, gene ontology (GO) analysis reported the term "response to abscisic acid" as 257 significantly up- and down-regulated in the raf22raf36-1 double mutant 258 (Supplemental Figure 8C and 8D). In raf22raf36-1, RNA- and metabolism-related 259 GO terms were significantly up- and down-regulated groups before ABA treatment, 260 respectively (Supplemental Figures 9A and 9B). Second, we analyzed 261 differentially-accumulating phosphopeptides for enrichment of motifs that 262 correspond to kinase recognition sequences. Analysis using the algorithm motif-x 263 (Wagih et al., 2016) identified two motifs, [-p(S/T)-P-] and [-(R/K)-x-x-p(S/T)-], that are MAPK- and SnRK2-/Calcium Dependent Protein Kinase (CDPK)-targeted 264 265 sequences, respectively, as enriched in phosphopeptides from ABA-treated

wild-type (Supplemental Figure 10). In addition, [-p(S/T)-P-] and [-(R/K)-x-x-p(S/T)-] containing phosphopeptides were also enriched in *raf22raf36-1* double mutant as compared to wild-type at each time point. Together, these two motifs account for over 70% of phosphopeptides that differentially accumulate in response to ABA in both wild-type and *raf22raf36-1*. This indicates Raf36 and Raf22 are directly or indirectly related to regulation of [-p(S/T)-P-] and/or [-(R/K)-x-x-p(S/T)-] *in vivo*.

272 To confirm our phosphoproteomic data, a subset of peptides were selected 273 that were upregulated by ABA in wild-type but downregulated in raf22raf36-1. These 274 candidates were synthesized as GST-fused 31 amino acid peptides and subjected 275 to *in vitro* phosphorylation assay. The amino acids 143-173 of OLEOSIN1 (OLE1) 276 was used as a positive control, because a previous study reported OLE1 as a 277 substrate of Raf22 at seed stage (Ramachandiran et al., 2018). Consistent with our 278 phosphorylation motif analysis (Supplemental Figure 10), both [-(R/K)-x-x-p(S/T)-] and [-p(S/T)-P-] containing phosphopeptides (AT1G21630.1: Calcium-binding EF 279 280 hand family protein, AT1G20760.1: Calcium-binding EF hand family protein and AT4G33050.3: calmodulin-binding family protein EDA39 for [-(R/K)-x-x-p(S/T)-] 281 282 motif; AT1G60200.1: splicing factor PWI domain-containing protein RBM25, 283 AT3G01540.2: DEAD-box RNA helicase 1 (DRH1), AT1G20440.1: cold-regulated 284 47 (COR47) for [-p(S/T)-P-] motif) were directly phosphorylated by Raf22 (Figure 285 5D).

286

#### 288 **DISCUSSION**

SnRK2s are core components of abiotic stress signaling in plants, yet the 289 290 full complement of SnRK2 substrates leading to responses necessary for stress 291 tolerance remain unknown. To gain a better understanding of SnRK2-mediated 292 signaling pathway(s), we aimed to identify signaling factors associated with SnRK2. 293 In this study, Raf36 was identified as a candidate of SnRK2-interacting protein. 294 Using several reverse genetic and biochemical analyses, we revealed that Raf36 295 and Raf22 are novel SnRK2 substrates which negatively regulate ABA responses in 296 a partially redundant manner at post-germination growth stage through their protein 297 kinase activities.

298 Our loss-of-function analyses revealed that Raf36 negatively regulates ABA 299 responses (Figures 2B, 2C, 2D, 2E and 2F). Notably, loss of *Raf36* altered the rate 300 of cotyledon greening, but did not affect seed germination or leaf water loss 301 (Supplemental Figures 3D, 5C and 5D). These results suggest that Raf36 may 302 function specifically in regulating ABA-induced post-germination growth arrest, a 303 physiological response that may allow germinated seeds to survive under 304 unfavorable conditions (Hwang et al., 2018). Raf36 is a member of subgroup C5 305 Raf-like MAPKKK family in Arabidopsis. Previous studies also reported that other 306 group C kinases, Raf22 and Raf43, are associated with ABA responses, e.g. raf22 307 mutant showed ABA-hypersensitive phenotype at post-germinative growth stage 308 (Hwang et al., 2018) and raf43 mutant showed ABA-hypersensitive phenotype at

309 seed germination stage (Virk et al., 2015). Our results demonstrated that Raf36 and 310 Raf22 functions partially redundantly in ABA-mediated post-germination growth 311 arrest, because raf22raf36-1 double knockout mutant showed a stronger ABA 312 hypersensitivity than in the individual single mutants (Figures 4E and 4F). In 313 addition to regulating post-germination growth arrest, our results show that Raf36 314 and Raf22 also regulate ABA-induced changes in gene expression and protein 315 phosphorylation in older 1- or 2-week-old seedlings (Figures 4G, 4H, 5B, 5C, 316 Supplemental Figures 8C and 8D). It has been known that some group C Raf-like 317 kinases function in various tissues, e.g. HT1 or BLUE LIGHT-DEPENDENT 318 H<sup>+</sup>-ATPASE PHOSPHORYLATION (BHP) in guard cells and Raf28 in 319 embryogenesis (Hashimoto et al., 2006; Hayashi et al., 2017; Wang et al., 2018). 320 Thus, Raf36 and Raf22 may also have different functions in different tissues and 321 developmental stages.

322 In our experiments, kinase-dead forms of Raf36 or Raf22 did not 323 complement phenotypes of raf36-1 or raf22 mutants, indicating that in vivo kinase 324 activity of both proteins is necessary for regulating ABA responses (Figure 5A and 325 Supplemental Figure 6). To identify possible substrates of Raf36 or Raf22, we 326 performed a comparative phosphoproteomic analysis of ABA responses in wild-type 327 and *raf22raf36-1* double mutant plants. Among phosphopeptides downregulated in 328 raf22raf36-1, GO terms "Response to abscisic acid" or "Response to osmotic stress" 329 were enriched (Supplemental Figure 8D), suggesting that Raf36 or Raf22 are

involved in ABA-responsive phosphosignaling pathways. Consistent with this, 330 331 raf22raf36-1 double mutant showed enhanced salt tolerance compared to wild-type 332 or the single mutant plants (Supplemental Figure 11), indicating that Raf36 and 333 Raf22 are involved in ABA signaling at least under certain stress conditions. Functional analyses of detected phosphoproteins will be required for further 334 335 understanding of phosphosignaling pathways under the control of Raf36 or Raf22. Unexpectedly, phosphoproteomic profiling revealed that Raf36 or Raf22 regulate 336 337 some phosphosignaling pathways even in the absence of exogenous ABA 338 treatment (Figure 5B). This suggests that Raf36 or Raf22 may have some roles 339 under non-stressed conditions. A recent investigation proposed that basal ABA levels under well-watered conditions modulate plant metabolism and growth 340 (Yoshida et al., 2019). Given that not only GO terms "Response to abscisic acid" but 341 342 also metabolism-related GO terms were significantly enriched in ABA-nontreated 343 raf22raf36-1 mutant (Supplemental Figure 9B) and that raf36 or raf22raf36-1 344 showed a slight growth retardation in a normal condition (Supplemental Figure 12), 345 Raf36 or Raf22 may regulate responses to basal ABA levels under normal 346 conditions. Further analysis will be required to address this possibility in the future.

347 Several previous studies proposed the relationship between SnRK2 and 348 Raf-like kinases, e.g. HT1 or PpARK (Hõrak et al., 2016; Matrosova et al., 2015; 349 Saruhashi et al., 2015; Tian et al., 2015). Actually, PpARK, a subgroup B3 Raf, has 350 been known as a direct upstream regulator of SnRK2, i.e. PpARK can activate

351 SnRK2s by phosphorylating the activation loop of SnRK2 (Saruhashi et al., 2015). 352 In this case, ARK acts as a positive regulator in ABA signaling in *Physcomitrella* 353 patens. In addition, recent studies reported that Raf10, a subgroup B2 Raf, also 354 acts as a positive regulator upstream of SnRK2s in Arabidopsis (Lee et al., 2015; 355 Nguyen et al., 2019). In contrast to these results observed in group B Rafs, our 356 results suggest that group C Rafs negatively regulate ABA signaling and that they 357 are directly phosphorylated by SnRK2s. To clarify how they negatively regulate ABA 358 responses, functional analysis of substrate candidates of Raf36 or Raf22, such as 359 those from our phosphoproteomic data, will provide useful information for further 360 understanding of group C Rafs in ABA signaling.

361

#### 363 METHODS

#### 364 **Plant Materials**

Arabidopsis thaliana ecotype Columbia (Col-0) was used as the wild-type. T-DNA 365 366 insertion mutant lines, raf36-1 (GK-459C10), raf36-2 (SALK 044426C) and raf22 367 (SALK 105195C) were obtained from ABRC or GABI-Kat. The raf36-1raf22 double knockout mutant was generated by crossing raf36-1 and raf22. The SRK2E/OST1 368 knockout mutant, srk2e (SALK 008068), was used as described previously 369 370 (Yoshida et al., 2002). The *srk2dsrk2e* double mutant was established by crossing 371 *srk2d* (GABI-Kat 807G04) and *srk2e*, and the *raf36-1srk2dsrk2e* triple mutant was 372 generated by crossing *srk2dsrk2e* double mutant and *raf36-1*. Seeds of wild-type, 373 mutants or transgenic plants were sterilized, and sown on GM agar plates as 374 described (Umezawa et al., 2009). After vernalization at 4 °C in the dark for 4 days, 375 they were incubated in a growth chamber under a continuous light condition at 22 °C for indicated periods. To test ABA sensitivity, seeds were sown on GM agar 376 377 medium with or without 0.5 µM ABA (Sigma, MO). Germination and greening rate 378 were scored daily for 14 days according to previous study (Fujii et al., 2007).

379

# 380 Plasmids

In this study, vector constructions were performed with Gateway cloning technology
 (Invitrogen, CA) unless otherwise noted. SRK2D, SRK2E and SRK2I cDNAs were
 previously cloned into pENTR/D-TOPO vector (Invitrogen, CA) (Umezawa et al.,

2009). In addition, Raf36, Raf36 N (amino acid residues 1-206), Raf36 KD+C (207-525) and Raf22 cDNAs were cloned into pENTR/D-TOPO or pENTR1A vector and sequenced. Site-directed mutagenesis was carried out as previously described (Umezawa et al., 2013). Those cDNAs were transferred to destination vectors, such as pGBKT7 and pGADT7 (Takara Bio, Japan), pSITE-nEYFP-C1 and pSITE-cEYFP-N1 (ABRC), pGEX6p-2 (GE healthcare, IL) and pBE2113-GFP (Yoshida et al., 2002).

391

# 392 Transgenic Plants

The pBE2113 vector, which drives C-terminal GFP-tagged Raf36 or Raf22 protein under the control of the Cauliflower Mosaic Virus (CaMV) 35S promoter, was constructed as described above. The transformation vector was introduced into *Agrobacterium tumefaciens* strain GV3101 by electroporation and transformed into Arabidopsis plants as described (Umezawa et al., 2004). The transformation lines were selected on GM agar medium containing 50 µg/ mL kanamycin and 200 µg/ mL claforan. Expression levels of transgene were checked by RT-PCR.

400

# 401 AlphaScreen<sup>®</sup>

The AlphaScreen<sup>®</sup> (Amplified Luminescent Proximity Homogeneous Assay) was carried out using an AlphaScreen<sup>®</sup> FLAG<sup>®</sup> (M2) Detection Kit (Perkin Elmer, MA) to detect protein-protein interactions. For screening of SnRK2-interacting MAPKKKs,

15 MAPKKKs were selected at random from the entire family members. Then, the 405 C-terminal FLAG (DYKDDDDK)-tagged MAPKKK proteins were expressed in 406 wheat germ extract (WGE) from in vitro synthesized mRNA obtained from 407 408 PCR-amplified cDNAs (Nomoto and Tada, 2018). The N-terminal biotinylated SnRK2 proteins, such as SRK2D, E and I, were also synthesized in WGE. The 409 410 protein quality (i.e., efficient synthesis with the expected molecular weight) of FLAG-tagged MAPKKKs and biotinylated SnRK2s was confirmed by western 411 412 blotting with an anti-FLAG antibody and streptavidin, respectively. The 413 FLAG-tagged MAPKKKs and biotinylated SnRK2s were mixed with acceptor beads coated with anti-FLAG antibody, donor beads coated with streptavidin, 0.01 % 414 415 Tween-20 and 0.1 % bovine serum albumin (BSA) in sterilized water-diluted control buffer provided in the kit and then incubated at 21 °C for 12 h. The AlphaScreen<sup>®</sup> 416 luminescence was detected with the infinite<sup>®</sup> M1000 Pro (TECAN, Switzerland). 417 WGE with no expressed proteins was employed as negative control (NC) to 418 419 estimate the luminescence caused by endogenous wheat germ proteins.

420

#### 421 Yeast two-hybrid analysis

Yeast two-hybrid analysis was employed using the MatchMaker GAL4 Two-Hybrid
System 3 (Takara Bio, Japan) as previously described (Umezawa et al., 2009). *Saccharomyces cerevisiae* strain AH109 was co-transformed with various pairs of
pGBKT7 vectors harboring SnRK2s (i.e., SRK2D, SRK2E and SRK2I) and pGADT7

vectors harboring Raf36. A single colony for each transformant grown on SD/-leucine (L)/-tryptophan (W) media was incubated in liquid media, and then evaluated on SD media supplemented with or without 3-amino-1,2,4-triazole (3-AT) and lacking combinations of amino acids leucine (L), tryptophan (W) and histidine (H), as follows: -LW, -LWH, -LWH +10 mM 3-AT, -LWH +50 mM 3-AT or on SD media lacking L, W, H, and adenine (A), as follows: -LW, -LWH, -LWHA. The plates were incubated at 30 °C for the optimal period.

433

# 434 Microscopy analyses of fluorescent proteins

435 To perform Agrobacterium-mediated bimolecular fluorescence complementation (BiFC) assay, pSITE-nEYFP-C1 vectors harboring SnRK2s (i.e., SRK2D, SRK2E 436 437 and SRK2I) or pSITE-cEYFP-N1 vectors harboring Raf-like kinases (i.e., Raf36 and Raf22) were introduced to A. tumefaciens strain GV3101(p19) by electroporation. A 438 439 single colony for each transformant was cultured in LB media, and the media was 440 substituted by 1/2 GM liquid media supplemented with 0.1 mM acetosyringone. 441 SnRK2 and Raf transformants were mixed with various pairs, and then infiltrated 442 into Nicotiana benthamiana leaves. Complemented YFP fluorescence of each 443 samples was observed in epidermall cells of N. benthamiana at 3 days after 444 infiltration with a fluorescence microscope BX53 (Olympus, Japan). For analysis of subcellular localization of Raf36-GFP, mesophyll cells of 2-week-old transgenic 445 Arabidopsis plants expressing Raf36-GFP were observed with a confocal 446

447 microscope SP8X (Leica Microsystems) with the time-gating method (Kodama, 448 2016), which completely eliminates chlorophyll autofluorescence when GFP 449 imaging. GFP fluorescence was observed with 484 nm excitation and 494-545 nm 450 emission with a gating time of 0.3–12.0 nsec. Chlorophyll autofluorescence was 451 separately observed with 554 nm excitation and 640–729 nm emission.

452

# 453 **Preparation of Recombinant Proteins**

DNA fragments of Raf36, Raf36 N (1-206), Raf36 KD+C (207-525), Raf36 KD 454 455 (207-467), Raf36 N (1-156), Raf36 N (1-140), Raf22, Raf43, Raf28, SRK2D, 456 SRK2E and SRK2I were amplified from cDNAs, and they were fused in-frame to pMAL-c5X vector (New England Biolabs, MA). Amino acid substitutions, such as 457 Raf36 K234N, Raf22 K157N, Raf22 S81A K157N, Raf43 K228N, Raf28 K158N and 458 SRK2E K50N, were introduced by site-directed mutagenesis. The MBP-fusion 459 proteins were expressed and purified from *E. coli* BL21 (DE3) using Amylose Resin 460 461 (New England Biolabs) according to the manufacturer's instructions. GST-SRK2E and GST-SRK2E K50N proteins were expressed using pGEX6p-2 (GE healthcare, 462 463 IL) and purified using Glutathione Sepharose 4B resin (GE healthcare, IL). Both MBP-tagged and GST-tagged recombinant proteins were further purified with a 464 Nanosep<sup>®</sup> 30-kDa size-exclusion column (PALL, NY). In addition, to identify 465 phosphorylation sites in Raf36, six types of 30-amino-acid peptides with mutations 466 467 were designed as Raf36 peptides (134-163) #1- #6 as shown in Fig. 3C. Similarly,

to confirm Raf22-dependent phosphorylation, six phosphopeptides and OLE1 as a
positive control, were designed as 31-amino-acid peptides including the putative
phosphorylation site(s). These peptides were expressed and purified from *E. coli*BL21 (DE3) using pGEX4T-3 vector (GE healthcare, IL).

472

# 473 *In vitro* phosphorylation assay

In vitro phosphorylation assays were performed as described previously with some modifications (Umezawa et al., 2009). The recombinant proteins of SnRK2, Raf or substrates were mixed with indicated pair(s) and incubated in 50 mM Tris-HCI (pH 7.5), 5 mM MgCl<sub>2</sub> or 5 mM MnCl<sub>2</sub>, 50  $\mu$ M ATP and 0.037 MBq of [ $\gamma$ -<sup>32</sup>P] ATP (PerkinElmer, MA) at 30 °C for 30 min. Samples were subsequently separated by SDS-PAGE, and phosphorylation levels were detected by autoradiography with BAS-5000 (Fujifilm, Japan).

481

# 482 Water loss analysis

To measure leaf water loss, 2-week-old seedlings were transferred from GM agar medium to soil, and the plants were grown under a 16 h/8 h (light/dark) photoperiod at 22 °C for another 2 weeks. The fully expanded rosette leaves were detached from 4- to 5-week-old plants and placed on weighing dishes. These dishes were kept under the same conditions used for seedling growth on soil, and then their fresh weights were monitored at the indicated times with three replicates per

time-point. One replicate consists of 5 individual leaves. Water loss was calculated
as a percentage of relative weight at the indicated times versus initial fresh weight.

491

# 492 **RNA extraction and qRT-PCR**

For quantitative reverse transcription PCR (qRT-PCR) analysis, total RNA was 493 494 extracted by LiCl precipitation from 1-week-old seedlings treated with 50 µM ABA for indicated periods. 1 µg of total RNA treated with RNase-free DNase I (Nippon 495 Gene, Japan) was used for reverse transcription with ReverTra Ace<sup>®</sup> reverse 496 transcriptase (TOYOBO, Japan). gRT-PCR analysis was performed using GoTag<sup>®</sup> 497 498 gPCR Master Mix (Promega, WI) with Light Cycler 96 (Roche Life Science, CA). For normalization, GAPDH was used as an internal control. The gene-specific primers 499 500 used for qRT-PCR analysis were shown in Dataset S5.

501

#### 502 **Phosphoproteomic analysis.**

Following imbibition with 50 µM ABA, 2-week-old Arabidopsis seedling of wild-type (Col-0) and *raf22raf36-1* were used for phosphoproteomic analysis with three biological replicates. Total crude protein was extracted from grounded sample. The phosphoproteomic analyses were performed as previously described with 400 µg of total crude protein (Ishikawa et al., 2019a, 2019b; Nakagami et al., 2010; Sugiyama et al., 2007; Umezawa et al., 2013). Enriched phosphopeptides by using HAMMOC method (Sugiyama et al., 2007) were analyzed with a LC-MS/MS system,

510 TripleTOF 5600 (AB-SCIEX). Peptides and proteins were identified using the 511 database (TAIR 10) with Mascot (Matrix Science, version 2.4.0). The false 512 discovery score was calculated using Benjamini-Hochberg method and set to 5 %. 513 Each phosphorylation site was assessed by the site localization probability score 514 calculated with the mascot delta score and defined confident as > 0.75 (Ishikawa et 515 al., 2019a). Skyline version 4.2 (Maccoss lab software) was used for quantification 516 of phosphopeptides on the basis of LC-MS peak area. All raw data files were 517 deposited in the Japan Proteome Standard Repository Database (jPOST; 518 https://repository.jpostdb.org/preview/20386920535e1580015868a, JPST000630. 519 access key; 3457). Each phosphoproteomic sample was compared by principal component analysis by using all identified phosphopeptides and their 520 phosphorylation level. The motif analysis was conducted using the Motif-X 521 522 algorithm. DAVID (https://david.ncifcrf.gov) and REViGO (http://revigo.irb.hr) were 523 used for GO analysis.

524

#### 525 Accession numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries
under the following accession numbers: *SRK2D*, *AT3G50500*; *SRK2E*, *AT4G33950*, *SRK2I*, *AT5G66880*, *Raf36*, *AT5G58950*; *Raf22*, *AT2G24360*; *Raf43*, *AT3G46930*; *Raf28*, *AT4G31170*; *RD29B*, *AT5G52300*, *RAB18*, *AT5G66400* and *OLE1*, *AT4G25140*.

# 532 Supplemental Information

- **Supplemental Figure 1.** AlphaScreen<sup>®</sup> assay for screening of SnRK2-interacting
- 534 MAPKKKs.
- **Supplemental Figure 2.** A phylogenetic tree of Arabidopsis Raf-like kinases.
- **Supplemental Figure 3.** Isolation and characterization of *raf36-1* and *raf36-2*
- 537 T-DNA insertion lines.
- **Supplemental Figure 4.** *in vitro* phosphorylation assays determining preference for

 $Mg^{2+}$  or  $Mn^{2+}$ .

- **Supplemental Figure 5.** The characterization of Raf22 in ABA response.
- **Supplemental Figure 6.** Raf22 protein kinase activity is required for its function in

542 ABA signaling.

- **Supplemental Figure 7.** An overview of phosphoproteomic analysis.
- **Supplemental Figure 8.** GO analysis of phosphopeptides in wild-type and *raf22raf36-1*.
- **Supplemental Figure 9.** GO analysis of phosphopeptides in *raf22raf36-1* in normal
- 547 condition.
- **Supplemental Figure 10.** Motif analysis of phosphopeptides in wild-type and *raf22raf36-1*.
- **Supplemental Figure 11.** Salt tolerance of *raf22raf36-1* plants.
- **Supplemental Figure 12.** Dwarf phenotype of *raf36* and *raf22raf36-1* plants under

552 normal condition.

553 **Supplemental Dataset 1.** List of phosphopeptides detected in this study.

554 **Supplemental Dataset 2.** List of responded phosphopeptides during ABA 555 treatment.

556 **Supplemental Dataset 3.** List of GO terms for phosphopeptides in this study.

557 **Supplemental Dataset 4.** Classification of phosphopeptides by motif groups.

558 **Supplemental Dataset 5.** Primer sequences used for quantitative RT-PCR 559 (qRT-PCR).

560

# 561 Acknowledgments

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570

# 571 **Conflict of interest**

572 The authors declare no conflict of interest.

573

# 574 **Author contributions**

- 575 Yo.K. and T.U. designed research; Yo.K., M.H., S.I., F.M. and S.K. performed 576 research; F.T., M.N., K.I., Yu.K., Y.T., D.T. and K.S. contributed new 577 reagents/analysis tools; Yo.K., S.I., F.M. and S.K. analyzed data; and Yo.K., S.C.P. 578 and T.U. wrote the paper.
- 579

### 580 Figure Legends

# 581 Figure 1. Raf36 interacts with subclass III SnRK2s.

(A) AlphaScreen<sup>®</sup> assay shows interaction of Raf36 and subclass III SnRK2s. Bars 582 583 indicate means  $\pm$  standard error (n=3), and asterisks indicate significant differences by Student's *t* test (P < 0.05). (B) Yeast two-hybrid (Y2H) assay shows interaction 584 585 between Raf36 and subclass III SnRK2s. Yeast cells expressing GAL4AD:Raf36 and GAL4BD:SnRK2s fusion proteins were incubated on SD media supplemented 586 587 with or without 3-amino-1,2,4-triazole (3-AT) and lacking combinations of amino 588 acids leucine (L), tryptophan (W) and histidine (H), as follows (in order from low to 589 high stringency): -LW, -LWH, -LWH +10 mM 3-AT, -LWH +50 mM 3-AT. Photographs 590 were taken at 10 days (SRK2D and SRK2E) or 12 days (SRK2I) after incubation. 591 (C) Subcellular localization of Raf36-GFP in leaf mesophyll cells. Chl indicates 592 chlorophyll autofluorescence. Scale bar, 20 µm. (D) BiFC assays for Raf36 and subclass III SnRK2s. SnRK2 and Raf36 were transiently expressed in N. 593

*benthamiana* leaves by Agrobacterium infiltration. Empty vector constructs were used as negative controls. nEYFP and cEYFP represent the N- and C-terminal fragments of the EYFP, respectively. BF indicates bright field images. Scale bar, 50 µm. (E) Y2H assay for truncated versions of Raf36 and SRK2E. Yeast cells co-expressing GAL4AD:Raf36, Raf36 N or Raf36 KD+C and GAL4BD:SRK2E fusion proteins were incubated on SD media lacking L, W, H, and adenine (A), as follows (in order from low to high stringency): -LW, -LWH, -LWHA.

601

# Figure 2. Raf36 negatively regulates SnRK2-dependent ABA response phenotypes in Arabidopsis seedlings.

(A) Abundance of *Raf36* mRNA transcripts measured by quantitative RT-PCR. Total 604 RNA was extracted from 1-week-old wild-type (WT) Col-0 Arabidopsis seedlings 605 606 treated with 50 µM ABA for indicated periods. Bars indicate means ± standard error (n=3), and asterisks indicate significant differences by Student's t test (\*P < 0.05, 607 608 \*\*P < 0.01). (B and C) Quantification of the cotyledon greening rates of WT (Col-0), 609 raf36-1 and raf36-2 on GM agar medium with or without 0.5 µM ABA. Data are 610 means ± standard error (n=3). Each replicate contains 36 seeds. Photographs were 611 taken 6 days after vernalization. (D) Functional complementation of raf36-1 by 612 *CaMV35S:Raf36-GFP*. Shown in photograph of seedlings grown for 7 days on GM agar medium in the presence or absence of 0.5 µM ABA. (E and F) Quantification of 613 614 the cotyledon greening rates of WT (Col-0), raf36-1, srk2dsrk2e and *raf36-1srk2dsrk2e* on GM agar medium in the presence or absence of 0.5 μM ABA.
Data are means ± standard error (n=3). Each replicate contains 36 seeds.
Photographs were taken 6 days after vernalization.

618

# 619 Figure 3. Subclass III SnRK2s directly phosphorylate Raf36.

620 (A) In vitro phosphorylation assay using kinase-dead forms of GST-SRK2E (SRK2E) 621 K50N) or MBP-Raf36 (Raf36 K234N). Each kinase-dead form was co-incubated 622 with an active GST-SRK2E or MBP-Raf36 kinase as indicated. Assays were performed in the presence of 5 mM  $Mn^{2+}$  (left 3 lanes) or 5 mM  $Mq^{2+}$  (right 3 lanes) 623 with [y-<sup>32</sup>P] ATP. (B) In vitro phosphorylation assay using truncated forms of 624 625 MBP-tagged Raf36. Each MBP-Raf36 protein was incubated with MBP-SRK2E in the presence of 5 mM Mg<sup>2+</sup> with  $[y-^{32}P]$  ATP. N: N-terminal region, KD: kinase 626 627 domain, C: C-terminal region. (C) Schematic representation of six Raf36 (134-163) peptides tested as SRK2E substrates. Ser141, Ser145, Ser150 and Ser155 are 628 629 labeled in blue, with alanine substitutions shown in red. Ser157, labeled in green, was replaced with alanine in Raf36 (134-163) peptides #1- #5. (D) In vitro 630 631 phosphorylation of Raf36 peptides by MBP-SRK2E. (E) In vitro phosphorylation of 632 GST-Raf36 (134-163) peptide #3 by MBP-SRK2D or MBP-SRK2I. Autoradiography (<sup>32</sup>P) and CBB staining (CBB) show protein phosphorylation and loading, 633 respectively. 634

635

#### 636 Figure 4. Raf22, a C6 Raf-like kinase, functions redundantly with Raf36.

637 (A) Phylogenetic tree of subfamily C5 and C6 Raf-like kinases in Arabidopsis. (B) In 638 vitro phosphorylation of C5/C6 Raf kinases by GST-tagged SRK2E. MBP-tagged 639 kinase-dead forms of Raf43 (Raf43 K228N), Raf22 (Raf22 K157N) or Raf28 (Raf28 640 K158N) were used as substrates. (C) In vitro phosphorylation of kinase-dead Raf22 641 (K157N) and Raf22 (S81A K157N) proteins by GST-SRK2E. (D) BiFC assay of 642 Raf22 and SRK2E in N. benthamiana leaves. nEYFP and cEYFP represent the N-643 and C-terminal fragments of the EYFP, respectively. BF indicates bright field images. 644 Scale bar, 50 µm. (E and F) Quantification of the cotyledon greening rates of wild-type (Col-0), raf36-1, raf22 and raf22raf36-1 on GM agar medium with or 645 646 without 0.5 µM ABA. Data are means ± standard error (n=4). Each replicate contains 36 seeds. Photographs were taken 9 days after vernalization. (G and H) 647 Relative gene expression of ABA-responsive genes. Total RNA was extracted from 648 649 1-week-old plants including wild-type, raf36-1, raf22 and raf22raf36-1 treated with 650 50  $\mu$ M ABA for indicated periods. Bars indicate means ± standard error (n=3) and asterisks indicate significant differences by Student's *t* test (\*P < 0.05, \*\*P < 0.01). 651

652

# Figure 5. Phosphoproteomic analysis of wild-type and *raf22raf36-1* identifies ABA signaling components downstream of Raf kinases.

(A) Functional complementation of *raf36-1* by *CaMV35S:Raf36-GFP* or *Raf36 K234N-GFP*. Shown in photograph of seedlings grown for 7 days on GM agar

medium in the presence or absence of 0.5 µM ABA. (B) Principal component 657 658 analysis of phosphoproteomic profiles of wild-type (WT) and raf22raf36-1. (C) Venn diagram of up- or down-regulated phosphopeptides in WT seedlings after 50 µM 659 660 ABA treatment, and up- or down-regulated phosphopeptides in raf22raf36-1 compared to WT (P < 0.05). (D) In vitro phosphorylation of GST-tagged 661 phosphopeptides from proteins AT1G21630.1 (lane 3), AT1G20760.1 (lane 4), 662 AT4G33050.3 (lane 5), AT1G60200.1 (lane 6), AT3G01540.2 (lane 663 7). AT1G20440.1 (lane 8) by MBP-Raf22. GST (lane 1) and GST-OLE1 fragment 664 (143-173 aa, lane 2) were included as negative and positive controls, respectively. 665 The autoradiography (<sup>32</sup>P) and CBB staining (CBB) show protein phosphorylation 666 and loading, respectively. 667

668

669 Supplemental Figure 1. AlphaScreen<sup>®</sup> assay for screening of 670 SnRK2-interacting MAPKKKs.

Interaction of SnRK2 with MAPKKKs was tested by AlphaScreen<sup>®</sup> assay. N-terminal biotin-tagged SRK2I protein and C-terminal FLAG-tagged MAPKKK proteins were synthesized by *in vitro* translation system in wheat germ extracts. After incubating in a reaction buffer including AlphaScreen<sup>®</sup> acceptor and donor beads for 12 h, the AlphaScreen<sup>®</sup> luminescence intensity was analyzed with a multi-mode plate reader (TECAN M1000pro).

677

#### 678 Supplemental Figure 2. A phylogenetic tree of Arabidopsis Raf-like kinases.

Amino acid sequences of predicted kinase domains from Raf-like kinases were aligned using ClustalW. The phylogenetic tree was generated using MEGA-X software with the neighbor-joining method. The Raf-like kinases were classified as B1- B4 and C1- C7 subfamilies, according to Ichimura et al., 2002. The C5 subfamily is shown in bold red letters.

684

685 Supplemental Figure 3. Isolation and characterization of *raf36-1* and *raf36-2* 

# 686 **T-DNA insertion lines.**

(A) Schematic depiction of *Raf36* genomic DNA with T-DNA insertions. Black boxes 687 and lines indicate exons and introns, respectively. (B) RT-PCR analysis of Raf36 688 transcript levels in wild-type (Col-0), raf36-1 and raf36-2 seedlings. GAPDH was 689 690 used as a positive control. (C) RT-PCR analysis of Raf36 transcript levels in raf36-1 and complementation lines (comp#3 and comp#4). GAPDH was used as a positive 691 692 control. (D) Germination rates of wild-type (Col-0), raf36-1 and raf36-2 on GM agar medium in the presence or absence of 0.5 µM ABA. Data are means ± standard 693 694 error (n=3). Each replicate contains 36 seeds.

695

Supplemental Figure 4. *in vitro* phosphorylation assays determining
 preference for Mg<sup>2+</sup> or Mn<sup>2+</sup>.

(A) Effects of Mg<sup>2+</sup> or Mn<sup>2+</sup> on kinase activity of Raf36. MBP-tagged Raf36 and

 $\alpha$ -casein were incubated with [y-<sup>32</sup>P] ATP in the presence of 5 mM Mg<sup>2+</sup> (left) or 5 699 mM Mn<sup>2+</sup> (right). Autophosphorylation (upper) and substrate phosphorylation 700 (lower) signals were visualized by autoradiography. Coomassie Brilliant Blue (CBB) 701 staining shows protein loading. (B) Effects of Mg<sup>2+</sup> or Mn<sup>2+</sup> on kinase activity of 702 703 SRK2E. MBP-tagged SRK2E and histone as substrate were co-incubated with  $[\gamma^{-32}P]$  ATP in the presence of 5 mM Mg<sup>2+</sup> (left) or 5 mM Mn<sup>2+</sup> (right). 704 705 Autophosphorylation (upper) and substrate phosphorylation (lower) signals were 706 visualized by autoradiography. Coomassie Brilliant Blue (CBB) staining shows 707 protein loading. (C) identify In vitro phosphorylation assav to SnRK2-phosphorylation site(s) in the N-terminal region of Raf36. 708

709

### 710 Supplemental Figure 5. The characterization of Raf22 in ABA response.

711 (A) gRT-PCR analysis of Raf22 mRNA transcript abundance. Total RNA was 712 extracted from 1-week-old wild-type seedlings treated with 50 µM ABA for indicated 713 periods. Bars indicate means ± standard error (n=3). Asterisks showed significant 714 differences by Student's *t* test (\*P < 0.05, \*\*P < 0.01). (B) BiFC assays of Raf22 and 715 SnRK2s. SRK2D or SRK2I were transiently expressed with Raf22 in N. 716 benthamiana leaves by Agrobacterium infiltration. nEYFP and cEYFP represent the 717 N- and C-terminal fragments of the EYFP, respectively. BF indicates bright field 718 images. Scale bar, 50 µm. (C) Germination rates of wild-type (Col-0), raf36-1, raf22 719 and raf22raf36-1 on GM agar medium with/without 0.5  $\mu$ M ABA. Data are means ±

720	standard error (n=3). Each replicate contains 36 seeds. (D) Water loss from
721	detached leaves of wild-type (Col-0), raf36-1, raf22 and raf22raf36-1 plants. The
722	<i>srk2e</i> mutant was included as a positive control. Data are means ± standard error
723	(n=3). Each replicate consists of five individual leaves.
724	
725	Supplemental Figure 6. Raf22 protein kinase activity is required for its
726	function in ABA signaling.
727	Functional complementation of raf22 mutant by introducing Raf22-GFP or Raf22
728	K157N-GFP. Photographs were taken 7 days after vernalization.
729	
730	Supplemental Figure 7. An overview of phosphoproteomic analysis.
731	(A) Distribution of the number of phosphosites per peptide. (B) Distribution of
732	phosphorylated residues in each peptide. $pS$ , $pT$ and $pY$ indicate phospho-serine,
733	phospho-threonine and phospho-tyrosine, respectively.
734	
735	Supplemental Figure 8. GO analysis of phosphopeptides in wild-type and
736	raf22raf36-1.
737	Each graph represents GO terms for upregulated (A) and downregulated
738	phosphopeptides (B) in ABA-treatment wild-type (WT) seedlings, or GO terms of
739	up- (C) or down-regulated phosphopeptides (D) in raf22raf36-1 as compared with
740	WT. GO terms were evaluated by DAVID program and visualized with REVIGO ( $P <$

0.05). Circle color and size show *P* value and frequency (%), respectively.

742

# 743 Supplemental Figure 9. GO analysis of phosphopeptides in *raf22raf36-1* in

- 744 normal condition.
- Each graph represents GO terms for phosphopeptides up- (A) or down-regulated
- (B) in *raf22raf36-1* under normal condition (P < 0.05). GO terms were evaluated by
- 747 DAVID program and visualized with REVIGO.
- 748

749 Supplemental Figure 10. Motif analysis of phosphopeptides in wild-type and

- 750 *raf22raf36-1*.
- 751 Phosphorylation motifs in up- or down-regulated phosphopeptides in response to
- ABA in wild-type seedlings, and motifs in up- or down-regulated phosphopeptides in
- *raf22raf36-1* in comparison with wild-type.

754

- 755 Supplemental Figure 11. Salt tolerance of *raf22raf36-1* plants.
- Wild-type (Col-0), *raf36-1*, *raf22* and *raf22raf36-1* seeds were germinated on GM agar medium with or without 150 mM NaCl. Photographs were taken 19 days after vernalization.

759

Supplemental Figure 12. Dwarf phenotype of *raf36* and *raf22raf36-1* plants
 under normal condition.

- 762 Wild-type, raf22, raf36 and raf22raf36-1 plants grown at 22°C under 16/8 h
- 763 photoperiod for 29 days. Scale bar, 3 cm.



#### Figure 1. Raf36 interacts with subclass III SnRK2s.

(A) AlphaScreen<sup>®</sup> assay shows interaction of Raf36 and subclass III SnRK2s. Bars indicate means  $\pm$ standard error (n=3), and asterisks indicate significant differences by Student's t test (P < 0.05). (B) Yeast two-hybrid (Y2H) assay shows interaction between Raf36 and subclass III SnRK2s. Yeast cells expressing GAL4AD:Raf36 and GAL4BD:SnRK2s fusion proteins were incubated on SD media supplemented with or without 3-amino-1,2,4-triazole (3-AT) and lacking combinations of amino acids leucine (L), tryptophan (W) and histidine (H), as follows (in order from low to high stringency): -LW, -LWH, -LWH +10 mM 3-AT, -LWH +50 mM 3-AT. Photographs were taken at 10 days (SRK2D and SRK2E) or 12 days (SRK2I) after incubation. (C) Subcellular localization of Raf36-GFP in leaf mesophyll cells. Chl indicates chlorophyll autofluorescence. Scale bar, 20 µm. (D) BiFC assays for Raf36 and subclass III SnRK2s. SnRK2 and Raf36 were transiently expressed in N. benthamiana leaves by Agrobacterium infiltration. Empty vector constructs were used as negative controls. nEYFP and cEYFP represent the N- and C-terminal fragments of the EYFP, respectively. BF indicates bright field images. Scale bar, 50 µm. (E) Y2H assay for truncated versions of Raf36 and SRK2E. Yeast cells co-expressing GAL4AD:Raf36, Raf36 N or Raf36 KD+C and GAL4BD:SRK2E fusion proteins were incubated on SD media lacking L, W, H, and adenine (A), as follows (in order from low to high stringency): -LW, -LWH, -LWHA.



#### Figure 2. Raf36 negatively regulates SnRK2-dependent ABA response phenotypes in Arabidopsis seedlings.

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#### Figure 3. Subclass III SnRK2s directly phosphorylate Raf36.

(A) *In vitro* phosphorylation assay using kinase-dead forms of GST-SRK2E (SRK2E K50N) or MBP-Raf36 (Raf36 K234N). Each kinase-dead form was co-incubated with an active GST-SRK2E or MBP-Raf36 kinase as indicated. Assays were performed in the presence of 5 mM Mn<sup>2+</sup> (left 3 lanes) or 5 mM Mg<sup>2+</sup> (right 3 lanes) with [ $\gamma$ -<sup>32</sup>P] ATP. (B) *In vitro* phosphorylation assay using truncated forms of MBP-tagged Raf36. Each MBP-Raf36 protein was incubated with MBP-SRK2E in the presence of 5 mM Mg<sup>2+</sup> with [ $\gamma$ -<sup>32</sup>P] ATP. N: N-terminal region, KD: kinase domain, C: C-terminal region. (C) Schematic representation of six Raf36 (134-163) peptides tested as SRK2E substrates. Ser141, Ser145, Ser150 and Ser155 are labeled in blue, with alanine substitutions shown in red. Ser157, labeled in green, was replaced with alanine in Raf36 (134-163) peptides #1- #5. (D) *In vitro* phosphorylation of Raf36 peptides by MBP-SRK2E. (E) *In vitro* phosphorylation of GST-Raf36 (134-163) peptide #3 by MBP-SRK2D or MBP-SRK2I. Autoradiography (<sup>32</sup>P) and CBB staining (CBB) show protein phosphorylation and loading, respectively.



#### Figure 4. Raf22, a C6 Raf-like kinase, functions redundantly with Raf36.

(A) Phylogenetic tree of subfamily C5 and C6 Raf-like kinases in Arabidopsis. (B) *In vitro* phosphorylation of C5/C6 Raf kinases by GST-tagged SRK2E. MBP-tagged kinase-dead forms of Raf43 (Raf43 K228N), Raf22 (Raf22 K157N) or Raf28 (Raf28 K158N) were used as substrates. (C) *In vitro* phosphorylation of kinase-dead Raf22 (K157N) and Raf22 (S81A K157N) proteins by GST-SRK2E. (D) BiFC assay of Raf22 and SRK2E in *N. benthamiana* leaves. nEYFP and cEYFP represent the N- and C-terminal fragments of the EYFP, respectively. BF indicates bright field images. Scale bar, 50 µm. (E and F) Quantification of the cotyledon greening rates of wild-type (Col-0), *raf36-1*, *raf22* and *raf22raf36-1* on GM agar medium with or without 0.5 µM ABA. Data are means  $\pm$  standard error (n=4). Each replicate contains 36 seeds. Photographs were taken 9 days after vernalization. (G and H) Relative gene expression of ABA-responsive genes. Total RNA was extracted from 1-week-old plants including wild-type, *raf36-1*, *raf22* and *raf22raf36-1* treated with 50 µM ABA for indicated periods. Bars indicate means  $\pm$  standard error (n=3) and asterisks indicate significant differences by Student's *t* test (\**P* < 0.05, \*\**P* < 0.01).



# Figure 5. Phosphoproteomic analysis of wild-type and *raf22raf36-1* identifies ABA signaling components downstream of Raf kinases.

(A) Functional complementation of *raf36-1* by *CaMV35S:Raf36-GFP* or *Raf36 K234N-GFP*. Shown in photograph of seedlings grown for 7 days on GM agar medium in the presence or absence of 0.5  $\mu$ M ABA. (B) Principal component analysis of phosphoproteomic profiles of wild-type (WT) and *raf22raf36-1*. (C) Venn diagram of up- or down-regulated phosphopeptides in WT seedlings after 50  $\mu$ M ABA treatment, and up- or down-regulated phosphopeptides in *raf22raf36-1* compared to WT (*P* < 0.05). (D) *In vitro* phosphorylation of GST-tagged phosphopeptides from proteins AT1G21630.1 (lane 3), AT1G20760.1 (lane 4), AT4G33050.3 (lane 5), AT1G60200.1 (lane 6), AT3G01540.2 (lane 7), AT1G20440.1 (lane 8) by MBP-Raf22. GST (lane 1) and GST-OLE1 fragment (143-173 aa, lane 2) were included as negative and positive controls, respectively. The autoradiography (<sup>32</sup>P) and CBB staining (CBB) show protein phosphorylation and loading, respectively.

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