UNIVERSITY of York

This is a repository copy of A Non-Canonical Function of Arabidopsis ERECTA Proteins in Gibberellin Signaling.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/200195/</u>

Version: Published Version

Preprint:

Davis, Seth Jon orcid.org/0000-0001-5928-9046 (2021) A Non-Canonical Function of Arabidopsis ERECTA Proteins in Gibberellin Signaling. [Preprint]

https://doi.org/10.1101/2021.12.02.470991

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	A Non-Canonical Function of Arabidopsis ERECTA	
2	Proteins in Gibberellin Signaling	
3		
4	Elzbieta Sarnowska ^{a#} , Szymon Kubala ^{b#} , Pawel Cwiek ^{b#} , Sebastian Sacharowski ^b , Paulina	
5	Oksinska ^b , Jaroslaw Steciuk ^b , Magdalena Zaborowska ^b , Jakub M. Szurmak ^b , Roman Dubianski ^a ,	
6	Anna Maassen ^b , Malgorzata Stachowiak ^a , Bruno Huettel ^c , Monika Ciesla ^b , Klaudia Kogut ^b , Anna	
7	T. Rolicka ^{b,d} , Saleh Alseekh ^{e,f} , Ernest Bucior ^b , Rainer Franzen ^g , Anna Klepacz ^b , Malgorzata A.	
8	Domagalska ^g , Samija Amar ^g , Janusz A. Siedlecki ^a , Alisdair R. Fernie ^{e,f} , Seth J. Davis ^{g,h,i *^} ,	
9	Tomasz J. Sarnowski ^{b*^}	
10		
11	^a Maria Sklodowska- Curie National Research Institute of Oncology, Roentgena 5, Warsaw,	
12	Poland	
13	^b Institute of Biochemistry and Biophysics Polish Academy of Sciences, Pawinskiego 5A	
14	Warsaw, Poland	
15	^c Max Planck Genome Centre Cologne, D-50820 Cologne, Germany	
16	^d Faculty of Biology, University of Warsaw, Warsaw, Poland	
17	^e Max Planck Institute of Molecular Plant Physiology, 14476 Potsdam-Golm, Germany	
18	^f Center for Plant Systems Biology and Biotechnology, 4000 Plovdiv, Bulgaria	
19	^g Max-Planck Institute for Plant Breeding Research; D-50829 Cologne, Germany	
20	^h State Key Laboratory of Crop Stress Biology, School of Life Sciences, Henan University,	
21	13 Kaifeng 475004, China	
22	ⁱ Department of Biology, University of York, York YO10 5DD, UK	
23		
24	Funding information: National Science Centre (Poland) UMO-2011/01/B/NZ1/00053 (TJS),	
25	UMO-2015/16/S/NZ2/00042 (SK), UMO-2011/01/N/NZ1/01525 (ATR), UMO-	
26	2011/01/N/NZ1/01530 (EB), UMO-2017/01/X/NZ2/00282 (AM), UMO-2018/28/T/NZ2/00455	
27	(PC), START 092.2016 fellowship by the Foundation for Polish Science (SS), Deutsche	
28	Forschungsgemeinschaft (DFG) DFG-DA1061/2-1, 111 Project grant D16014, BBSRC-	
29	BB/M000435/1, and Max-Planck Gesellschaft (MPG) core funding (SJD), scholarship of	
30	Ministry of Science and Higher Education (MNiSW) No. 466/STYP/11/2016 (SK), SA and ARF	
31	acknowledge funding of the PlantaSYST project by the European Union's Horizon 2020 research	

32	and innovation programme (SGA-CSA No 664621 and No 739582 under FPA No. 664620), the			
33	equipment used was sponsored in part by the Centre for Preclinical Research and Technology			
34	(CePT), a project co-sponsored by European Regional Development Fund and Innovative			
35	Economy, The National Cohesion Strategy of Poland.			
36				
37	The authors declare no competing interest.			
38	[#] shared first authorship			
39	* senior authors			
40	^ address correspondence to:			
41	Tomasz J. Sarnowski tsarn@ibb.waw.pl			
42	Seth J. Davis seth.davis@york.ac.uk			
43	Corresponding authors: Seth J. Davis and Tomasz J. Sarnowski			
44	Senior authors: Seth J. Davis and Tomasz J. Sarnowski			
45				

3

- 46 Short title: The Role of ERECTA and SWI3B in GA Signaling
- 47 **ONE SENTENCE SUMMARY:** ERECTA leucine-rich receptor-like kinase and SWI3B subunit
- 48 of SWI/SNF chromatin remodeling complex cooperate in direct transcriptional control of *GID1*
- 49 genes in Arabidopsis.
- 50

51 Authors Contributions

- 52 TJS, ES, and SJD planned experiments and wrote the manuscript
- 53 SK and PC participated in the planning of some experiments
- 54 TJS, ES, SK, PC, SS, SA, BH, JAS, and ARF analyzed the data
- 55 ES, PC, SS, SK, PO, JS, MZ, JMS, RD, AM, MS, BH, MC, KN, ATR, EB, RF, AK, MAD, SA,
- 56 and TJS performed experiments
- 57 All authors read, edited, and approved the final manuscript
- 58 Key Words: Arabidopsis, ERECTA, ERECTA-LIKE1, ERECTA-LIKE2, LRR-RLK, SWI/SNF,
- 59 SWI3B, HER2, Chromatin
- 60

61 Abbreviations and Acronyms:

- 62 ERf, ERECTA family; ER, ERECTA; ERL1, ERECTA-LIKE 1; ERL2, ERECT-LIKE 2; LRR-
- 63 RLKs, leucine-rich repeat receptor-like kinases; CRC, chromatin remodeling complex;
- 64 SWI/SNF, Switch/Sucrose Nonfermenting; *GID1*, *GIBBERELLIN INSENSITIVE DWARF 1*; GA,
- 65 gibberellin; PAC, Paclobutrazol; qRT-PCR, quantitative real-time PCR; BFA, Brefeldin A; NLS,
- 66 nuclear localization signal; KDER, the kinase domain of ERECTA; TSS, transcription start site;
- 67 EGFR, epidermal growth factor receptor.
- 68

4

69 Abstract

70 The Arabidopsis ERECTA family (ERf) of leucine-rich repeat receptor-like kinases 71 (LRR-RLKs), comprising ERECTA (ER), ERECTA-LIKE 1 (ERL1) and ERECTA-LIKE 2 72 (ERL2), control epidermal patterning, inflorescence architecture, stomata development, and 73 hormonal signaling. Here we show that the *er/erl1/erl2* triple mutant exhibits impaired 74 gibberellin (GA) biosynthesis and perception alongside broad transcriptional changes. ERf proteins interact in the nucleus, via kinase domains, with the SWI3B subunit of the SWI/SNF 75 76 chromatin remodeling complex (CRCs). The er/erl1/erl2 triple mutant exhibits reduced SWI3B 77 protein level and affected nucleosomal chromatin structure. The ER kinase phosphorylates 78 SWI3B in vitro, and the inactivation of all ERf proteins leads to the decreased phosphorylation of 79 SWI3B protein *in vivo*. Correlation between DELLA overaccumulation and SWI3B proteasomal 80 degradation together with the physical interaction of SWI3B with DELLA proteins explain the 81 lack of RGA accumulation in the GA- and SWI3B-deficient erf mutant plants. Co-localization of 82 ER and SWI3B on GID1 (GIBBERELLIN INSENSITIVE DWARF 1) DELLA target gene 83 promoter regions and abolished SWI3B binding to GID1 promoters in er/erl1/erl2 plants supports the conclusion that ERf-SWI/SNF CRC interaction is important for transcriptional 84 85 control of GA receptors. Thus, the involvement of ERf proteins in transcriptional control of gene expression, and observed similar features for human HER2 (Epidermal Growth Family Receptor-86 87 member), indicate an exciting target for further studies of evolutionarily conserved non-canonical 88 functions of eukaryotic membrane receptors.

5

90 Introduction

91 The ERECTA family (ERf) of leucine-rich-repeat receptor-like kinases (LRR-RLKs) 92 consists of three members: ERECTA (ER), ERECTA-LIKE 1 (ERL1), and ERECTA-LIKE 2 93 (ERL2). ERf proteins carry extra-cellular leucine-rich repeats (LRRs), as well as transmembrane 94 and cytosolic kinase domains (Shpak et al., 2004; Torii et al., 1996, Kosentka et al., 2017). 95 Inactivation of ERECTA leads to inflorescence, pedicels, and siliques compaction, while the 96 individual loss of either ERL1 or ERL2 function has a limited effect on Arabidopsis development 97 (Shpak et al., 2004). ERf proteins are functionally redundant-their simultaneous inactivation 98 results in dramatic growth retardation, severe dwarfism, enlargement of the shoot apical meristem 99 (SAM), clustered stomata, and sterility. ERf regulates stem cell homeostasis via buffering 100 cytokinin responsiveness and auxin perception in SAM and modulating the balance between stem 101 cell proliferation and consumption (Shpak et al., 2004; Griffiths et al., 2006; Torii et al., 2007; 102 Chen et al., 2013; Shpak, 2013; Uchida et al., 2013; Zhang et al., 2021). ERECTA controls the 103 expression of genes associated with gibberellin (GA) metabolism (Uchida et al., 2012a) 104 restricting xylem expansion downstream of the GA pathway (Ragni et al., 2011). It additionally 105 regulates shade avoidance in a GA and auxin-dependent manner (Du et al., 2018) and 106 ethylene-induced hyponastic growth (Van Zanten et al., 2010).

107 Overexpression of ER variant lacking the C-terminal kinase domain (ER ΔK) caused more 108 severe developmental defects than complete inactivation of ERECTA, suggesting an interaction 109 of the kinase domain with important regulatory partners (Shpak, 2003). ERECTA interacts with 110 ERL1 and ERL2 to form receptor complexes recognizing two endodermis-derived peptide 111 hormones (EPFL4 and EPFL6), regulating vascular differentiation and stem elongation. ERf 112 proteins additionally form complexes with the receptor-like protein TOO MANY MOUTHS 113 (TMM), which controls stomatal differentiation by recognition of the secretory peptides 114 EPIDERMAL PATTERNING FACTOR 1 (EPF1), EPF2, and stomagen (Lee et al., 2012; 115 Uchida et al., 2012a; Lee et al., 2015b).

116 ERL2 has been found to undergo endocytosis (Ho et al., 2016), suggesting that ERf 117 proteins may play, as yet uncharacterized, regulatory roles upon internalization, in addition to 118 their functions as ligand-binding membrane receptors. ERECTA signaling, in tandem with the 119 SWR1 chromatin remodeling complex (CRC), controls the expression of the *PACLOBUTRAZOL* 120 *RESISTANCE 1* (*PRE1*) family genes. This observation supports their role in the GA signaling

6

pathway, however, neither direct interaction between ERECTA and SWR1 nor the direct
influence of ERECTA signaling on chromatin structure or SWR1 activity has, as yet, been
demonstrated (Cai et al., 2017; Cai et al., 2021).

124 Here we show that the loss of all ERf proteins in the *er/erl1/erl2* triple mutant (*erf*) results 125 in broad transcriptomic changes affecting hormonal, developmental, and metabolic processes. 126 Inactivation of ERf proteins caused down-regulation of the GA receptor GID1 (GIBBERELLIN 127 INSENSITIVE DWARF 1) genes expression and decreased bioactive GA levels. The ER protein 128 undergoes endocytosis and enters the nucleus. All three ERf proteins interact in the nucleus with 129 the SWI3B core subunit of the SWI/SNF CRCs. The kinase domain of the ER protein exhibits the 130 ability to phosphorylate SWI3B protein. The physical interaction of SWI3B with RGA and 131 RGL1, together with identified correlation between DELLA accumulation and SWI3B 132 proteasomal degradation, provide an explanation as to why GA-deficient erf mutant plants did 133 not overaccumulate RGA. These data collectively suggest cooperation of ERf-signaling with 134 SWI/SNF in the modulation of gene transcription. The ER and SWI3B also co-localized in the 135 promoter regions of GID1 DELLA target genes. In the erf mutant, the binding of SWI3B to GID1 136 promoters was abolished. These results collectively suggest that ERf proteins directly control GA 137 receptor expression by restricting recruitment of the SWI/SNF CRCs to its target *loci*.

139 **Results**

140 Inactivation of *Erf* Proteins Has a Broad Effect on the Arabidopsis Transcriptome 141 including GA Signaling.

- 142 The Arabidopsis *er/erl1/erl2* plants exhibit severe dwarfism, dark green color, defects in
- 143 vascular development, stem elongation, and stomatal differentiation, as well as complete sterility
- 144 (Figure 1 A, Supplemental Figure 1A, (Shpak et al., 2004)).
- 145

bioRxiv preprint doi: https://doi.org/10.1101/2021.12.02.470991; this version posted December 3, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.



146

147 Figure 1. *ERf* inactivation affects Arabidopsis development, causes transcriptomic changes 148 overlapping with the effect of gal-3 mutation and impairs GA biosynthesis and signaling (See 149 also Figures S1, S2 and S3). A, Phenotypic changes conferred by combinations of *erf* mutations. 150 Scale bar= 1 cm. B. Overlapping down-regulated genes in er/erll/erl2 and gal-3 plants. C. 151 Overlapping up-regulated genes in er/erl1/erl2 and ga1-3 plants. D, The er/erl1/erl2 plants 152 exhibit impaired GA response. 14-days old LD (12h day/12 night) grown WT and er/erl1/erl2, 153 sprayed twice a week with water (upper row) or 100µM GA₄₊₇ (lower row). Arrows-er/erl1/erl2 154 plants. Scale bar= 1cm. E, The GA response is retained to various levels in combinations of *erf* 155 mutants. Error bars-SD,* = P < 0.05, Student's *t*-test, *n*= 30 plants. F, The response of various *erf* mutants to 1µM Paclobutrazol treatment. Error bars-SD,* = P < 0.05, Student's t test, n= 30 156 157 plants. G, The er/erl1/erl2 mutant exhibits altered transcription of GID1 GA receptor genes (error 158 bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates were assayed). H, 159 The er/erl1/erl2 mutant displays altered GA biosynthesis and metabolism-related genes 160 expression (error bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates 161 were assayed). I. The *er/erl1/erl2* mutant exhibits dramatically reduced level of bioactive GA_{4+7} 162 gibberellin (error bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates 163 were assayed). J, The er/erl1/erl2 mutant shows decreased level of the DELLA protein RGA. 164

9

165 Given the severe phenotypic alterations of the *er/erl1/erl2* plants, we performed transcript 166 profiling with Affymetrix ATH1 microarrays on RNA samples from aerial parts of the 167 er/erl1/erl2 mutant and WT (wild type) adult plants (representing the most comparable stage of 168 the development) grown for 5 weeks under long-day conditions (16h day/8h night). Data analysis 169 identified 1734 versus 1680 genes showing >1.50-fold decrease and increase, respectively, of 170 transcript levels in er/erl1/erl2 comparing to WT (Supplemental Figure 1B, Supplemental 171 Dataset 1 Sub-tables 1, 2). Gene Ontology (GO) terms of primary metabolism, developmental 172 processes, and response to hormones were enriched among the er/erl1/erl2 down-regulated genes 173 (Supplemental Dataset 1 Sub-table 3). Among these, 27 genes were classified to GA-response 174 (Supplemental Table 1, Supplemental Dataset 1 Sub-tables 1, 3). The up-regulated genes were 175 classified into GO-terms of chloroplast-related metabolic and light-regulated transcription 176 processes, responses to cytokinin, and auxin degradation (Supplemental Dataset 1 Sub-Tables 177 2.4). Several genes acting in leaf epidermal and stomatal cell differentiation showed enhanced 178 transcription in the er/erl1/erl2 mutant (Supplemental Table 2). In conclusion, the inactivation of 179 ERf altered transcriptional regulation of hundreds of targets, including a set of GA-regulated 180 genes.

Phenotypic traits exhibited by double and triple *erf* mutants resemble those of double and triple *gid1abc* (*gibberellin insensitive dwarf 1a, b* and *c*) plants (Figure 1A; (Griffiths et al., 2006)). Inactivation of *GID1abc* genes has a nearly identical effect on the Arabidopsis transcriptome as the severe GA-deficient mutant *ga1-3* (Willige et al., 2007), thus we compared the transcriptomic data available for the *ga1-3* mutant with those caused by inactivation of all *ERf* genes.

187 We identified a large overlap of differentially expressed genes (DEG) in the *er/erl1/erl2* 188 and gal-3 lines. Among 826 genes down-regulated in the gal-3 line, 351 (about 42.5%) also 189 exhibited decreased expression in the *er/erl1/erl2* plants (Figure 1B), while 104 genes (about 190 24.6% of gal-3 up-regulated genes) were up-regulated in both lines (Figure 1C). Only 33 genes 191 were up-regulated in gal-3 but down-regulated in er/erl1/erl2 (Supplemental Figure 1C), and 192 only 64 genes down-regulated in gal-3 but up-regulated in er/erl1/erl2 (Supplemental Figure 193 1D). DEG common to gal-3 and er/erl1/erl2 lines belonged to both DELLA (repressors of GA 194 pathway) -dependent and DELLA-independent classes (Cao et al., 2006), regardless of whether 195 they display co-regulation or contrasting regulation in these lines (Supplemental Figure 1E and

10

196 F). This suggests the involvement of Arabidopsis ERf proteins in the control of GA-related 197 processes. Therefore, we next tested the response of er/erl1/erl2 plants to exogenously supplied 198 bioactive 100 µM GA4+7 and found that spraying of the er/erl1/erl2 mutant grown under LD 199 condition (12h day/12h night) did not lead to increased leaf size by day 14 compared to the 200 remarkable expansion of control WT rosette leaves (Figure 1D, Supplemental Figure 2A). Thus 201 er/erl1/erl2 displayed GA insensitivity. Nonetheless, the GA-treatment resulted in bolting of 202 er/erl1/erl2 plants, only after over two months (Supplemental Figure 2B and C), indicating their 203 residual response to GA.

Although we showed that ERf proteins are involved in the GA response, it remained unclear whether proper GA perception requires all ERf proteins. Thus, we tested the hypocotyl response of single and double *erf* mutants in various combinations to the treatment with 1 μ M GA₄₊₇ or 1 μ M Paclobutrazol (PAC), an inhibitor of GA biosynthesis. The GA response was retained to various levels in all tested mutants (Figure 1E) while the *erl1* and *er/erl1* plants had an impaired response to PAC and *erl1/erl2* displayed a significant reduction of hypocotyl length (Figure 1F).

Upon crossing *er*, *er/erl1*, and *er/erl2* lines with the *ga1-3* mutant, we observed only a discrete enhancement of the *ga1-3* phenotype. However most of the phenotypic changes characteristic for *ga1-3* mutation were retained, indicating that many of the *er*, *erl1*, or *erl2* single or double mutant phenotypes are likely not exclusively a result of GA deficiency (Supplemental Figure 2D and E).

216 We have proven that only parallel inactivation of all ERf proteins causes severe 217 impairment of the GA response. Quantitative real-time PCR (qRT-PCR) measurements of GA 218 response and biosynthesis genes expression revealed a parallel 2.5 to 3-fold reduction in the 219 transcript levels of all three GID1 GA-receptors in the er/erl1/erl2 mutant compared to WT 220 (Figure 1G). The GA-receptor genes GID1A/B have been reported to be direct ChIP targets of 221 RGA, a major DELLA repressor of GA-signaling, which stimulates GID1 transcription (Zentella 222 et al., 2007). The er/erl1/erl2 triple mutant also displayed altered expression of GA biosynthesis 223 genes compared to the WT: a 4-fold reduction of mRNA levels of KAO1 (ent-kaurenoic acid 224 oxidase) and KO (ent-kaurene oxidase), a 2.5-fold increase of mRNA levels of GA-repressed 225 GA3ox1 and GA3ox2 (GIBBERELLIN 3 BETA-HYDROXYLASE 1 and 2), a 2-fold inhibition 226 and 5-fold up-regulation, respectively, of mRNA levels corresponding to the GA20ox1 and

11

227 GA20ox3 genes (Figure 1H). This indicated that the ERf proteins not only influence the 228 expression of GA receptors, but also genes associated with GA biosynthesis. We subsequently 229 found a substantial decrease of bioactive GA₄ as well as GA₁₂ and GA₂₄ intermediates in 230 er/erl1/erl2 mutant (Figure 1I, Supplementary Figure 3). Counterintuitively, the Western blotting 231 using a specific antibody (Willige et al., 2007) detected reduced levels of RGA in the *er/erl1/erl2* 232 mutant plants (Figure 1J). Our results indicate that the parallel inactivation of all ERf proteins 233 results in co-ordinate deregulation of GA biosynthesis and response pathways in Arabidopsis. 234

235 ERECTA (ER) Protein Undergoes Endocytosis and Migrates to the Nucleus.

236 In analogy to some human membrane receptors internalizing to endosomes and migrating 237 to the nucleus (i.e., Giri et al., 2005), the ERL2 member of the ERf undergoes endocytosis (Ho et 238 al., 2016). We next examined, in detail, the cellular localization of ER by creating C-terminal 239 GFP fusions with ER (Figure 2A) after verifying genetic complementation of the er-105 mutation 240 by a 35S::ER-GFP construct (Supplemental Figure 4). 241

12





243 Figure 2. Subcellular localization of ERECTA protein (See also Figures S4 and S5). A, 244 ERECTA is localized in plasma-membrane and endosomes in epidermal cells of 7-days old seedlings. ER-GFP, or free GFP visualized using GFP channel. FM4-64 specifically stains 245 246 plasma-membranes. Scale bar=10µm. B, Root-tip images of approximately two-week-old (14-17 247 days) ER-GFP seedlings showing nuclear localization of ERECTA protein at considerable 248 frequency. C, Root-tip images of 12-day-old ER-GFP seedlings serving as the control for D and 249 E. D, Brefeldin A treatment enhanced the localization of ERECTA protein in Brefeldin A (BFA) 250 bodies. Roots of 12-day-old Arabidopsis seedlings. E, Leptomycin B treatment enhanced the 251 nuclear localization of ERECTA Free GFP was used as a control in C, D, and E, cell nuclei were 252 stained with DAPI, scale bar= 50µm. F, Letomycin B enhances nuclear presence of ER protein. 253 The GFP/DAPI ratio calculated per area for roots of plants expressing ER-GFP protein.

13

255 As observed earlier (Shpak et al., 2005; Uchida et al., 2012a), a pool of the ER-GFP 256 protein was detected in association with plasma-membranes of the leaf epidermis. In addition, a 257 weak localization signal was detected in internal structures, which could represent endosomes 258 (Figure 2A). In guard cell pairs, ER-GFP protein was also detected in circles around the positions 259 of nuclei, which were visualized by propidium iodide staining (Supplemental Figure 5A and 260 Supplemental Movie 1). We also observed with considerable frequency ER protein in the nuclei 261 of roots of 14- to 17-day-old Arabidopsis plants (Figure 2B), however ER was mainly located in 262 the plasma membrane and endosome-like structures (Figure 2C).

263 To verify that the ER-GFP protein indeed undergoes endocytosis, we examined its 264 localization in Arabidopsis seedlings treated with 25µM Brefeldin A (BFA), a compound 265 preventing Golgi-mediated vesicular transport of membrane proteins to the plasma membrane 266 (Miller et al., 1992). We observed accumulation of the ER-GFP protein in BFA bodies within 30-267 40 min after BFA treatment leading to its accumulation at the nuclei periphery 120 min after 268 BFA application (Figure 2D, Supplemental Figure 5B). The 4h long 200 nM leptomycin B 269 treatment (a compound blocking nuclear export by EXPORTINS (Haasen et al., 2002)) resulted 270 in the ER accumulation in the cell nuclei (Figure 2E, F). ER thus appeared to behave similarly to 271 certain human plasma-membrane receptors in migrating into the nucleus (Hung et al., 2008; Chen 272 and Hung, 2015).

273 We noted that ERL1 and ERL2 proteins carry a monopartite nuclear localization signal 274 (NLS) sequence in their kinase domains, while the ERL1 kinase domain carries an additional 275 bipartite NLS identified using cNLS Mapper (Kosugi et al., 2009b). The NLS signal in the ER 276 protein was not recognized, however, all ERf proteins show evolutionally conserved amino acid 277 sequences in this region (Supplemental Figure 6A). Using the NetNES1.1 (la Cour et al., 2004) 278 server, we predicted the existence of specific for AtXPO1/AtCRM1 exportin (Haasen et al., 279 2002) leucine-rich nuclear export signals (NES) in all ERf proteins (Supplemental Figure 6A). 280 The subsequent Western-blotting analysis of nuclear extracts (Supplemental Figure 6B) 281 confirmed the nuclear presence of ER. In addition to the expected full-length forms (140 kDa), 282 we also detected shorter (~75 kDa) versions of the ERECTA protein with the C-terminal GFP tag 283 and smaller products of degradation, including free GFP, suggesting an analogy to the human 284 Epidermal Growth Factor Receptor (EGFR), (Chen and Hung, 2015). The detection of N-285 terminally truncated forms of the ERECTA protein carrying a kinase domain resembled the

14

recently reported fate of the XA21 LRR-RLK immune receptor in rice (Park and Ronald, 2012),
where its C-terminal kinase domain enters the nucleus to interact with the OsWRKY62
transcriptional regulator.

289 To assess whether the ERECTA kinase domain (KDER) is imported into the nucleus, we 290 fused the C-terminal part of ERECTA, harboring the KDER, to a YFP-HA tag (Supplemental 291 Figure 6C) and expressed this construct in the er-105 mutant. The presence of KDER-YFPHA 292 was detected exclusively in cell nuclei (Figure 3A). Furthermore, KDER-YFPHA expression 293 partially restored the *er-105* rosette and cauline leaf phenotype to WT values (Supplemental 294 Figure 7A, B). Still, it failed to genetically complement the defect of stem elongation 295 (Supplemental Figure 7C). Partial genetic complementation of the er-105 mutation and nuclear 296 localization of KDER prove that the KDER has a receptor-domain independent signaling 297 function. Interestingly, by contrast to the full length and kinase domain of ERECTA protein 298 (Figure 3A), truncated ERAkinase form of ERECTA protein fused to YFP-HA (AKDER-YFP-299 HA) did not enter into the nucleus proving the presence of functional NES and NLS sequences in 300 the C-terminal part of ERECTA protein.

bioRxiv preprint doi: https://doi.org/10.1101/2021.12.02.470991; this version posted December 3, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.





304 Figure 3. Nuclear function of ERf proteins (See also Figures S6, S7, S8, S9, and S10). A. Root-305 tip images of approximately two-week-old (14-17 days) plants expressing ER-GFP, KDER-YFP-306 HA (the kinase domain of the ER protein), ER Δ K-YFPHA (truncated ER protein lacking the 307 kinase domain) proteins indicating that the kinase domain is necessary for the nuclear localization 308 of ER protein. WT and GFP expressing plants-negative controls. Panel 2 in the ER-GFP indicates 309 nuclear localization of ER protein appearing at considerable frequency. Cell nuclei were stained 310 with DAPI. Scale bar=25 μ m. B, The *er-105/swi3b3* double mutant shows more retarded growth 311 than either er-105 or swi3b3 plants. Scale bar= 1cm. C, The er/erl1/erl2 triple mutant exhibits 312 reduced SWI3B protein level. D, ER-GFP or free GFP (negative control) pull-down from the 313 nucleus and anti-SWI3B western blotting indicate a specific ER-SWI3B interaction. E, 314 Immunoprecipitation of KDER-YFP-HA from the nucleus indicated that the kinase domain of 315 ER interacts with SWI3B. F, ER, ERL1, and ERL2 kinase domains interact with SWI3B in the 316 nucleus. Bimolecular Fluorescence Complementation assay (BiFC) in epidermis of tobacco 317 leaves. Scale bar = $10\mu m$.

318

302

16

Upon nuclei fractionation (Sarnowski et al., 2002), the ERECTA protein was detected in the nuclear membrane, soluble nuclear-protein fraction, and chromatin, with its major presence within the nuclear matrix. The nuclear fractions contained both full-length and N-terminally truncated ER forms (Supplemental Figure 8) suggest that ER could be involved in either transcriptional regulation or other nuclear functions.

324

325 ERf Proteins Physically Interact with the SWI3B Core Subunit of SWI/SNF CRC

326 The weak swi3b-3 allele (Sáez et al., 2008) carrying, in the er-105 background, a point 327 mutation in the SWI3B gene encoding a core subunit of the SWI/SNF chromatin remodeling 328 complex (CRC) exhibit severe dwarfism, altered leaf shape, delayed flowering and reduced 329 fertility (Figure 3B). We, therefore, introgressed the *swi3b-3* mutation into WT and found that the 330 phenotypic alterations related to swi3b-3 were much weaker (slight reduction of growth rate, 331 leading to decreased plant height, Figure 3B) than the phenotypic traits exhibited by the er-332 105/swi3b-3 as well as single er-105 mutation. The severe phenotypic alterations exhibited by the 333 er-105/swi3b-3 plants indicated the likely existence of a strong genetic interaction between the 334 ERECTA signaling pathway and SWI3B-containing SWI/SNF CRCs. This observation is in line 335 with i) the direct binding of 15 out of 27 potential ERf target genes related to the GA signaling 336 pathway (Supplemental Table 1) by SWI/SNF CRCs (Sacharowski et al., 2015; Archacki et al., 337 2016; Li et al., 2016); *ii*) the unexpected broad transcriptional changes and severe effects on 338 Arabidopsis development and hormonal signaling pathways observed in the *er/erl1/erl2* mutant, 339 and *iii*) the well-recognized function of SWI/SNF CRC in hormonal crosstalk including GA 340 signaling (Sarnowska et al., 2013; Sarnowska et al., 2016).

We next assessed the level of the SWI3B protein in *er/erl1/erl2* plants. We found a significant decrease in the SWI3B protein abundance (Figure 3C), further suggesting that the ERf signaling pathway may influence the proper function of SWI3B-containing SWI/SNF CRCs. Additionally, the SWI3B was found to bind ER-GFP but not free GFP (Figure 3D). Similarly, coimmunoprecipitation indicated that SWI3B interacts with the kinase domain of ER (Figure 3E).

Next, we performed BiFC assays (Hu et al., 2002) in epidermal cells of *Nicotiana benthamiana* and confirmed the SWI3B and ER kinase domain interaction. The YFC-RFP served as a control unrelated protein with broad intracellular localization (Figure 3F, Supplemental Figure 9). We also detected the interaction of SWI3B with the kinase domain of the ERL1 or

17

350 ERL2 (Figure 3F, Supplemental Figure 9), indicating the existence of direct interdependences
351 between the ERf signaling pathway and SWI/SNF-dependent chromatin remodeling.

Moreover, we found similar interactions in the nuclei of human cells for HER2 (Epidermal Growth Factor Receptor- family member), a membrane receptor acting in a noncanonical signaling mode including translocation to the nucleus (Lee et al., 2015a), and BAF155 a SWI3-type subunit of human SWI/SNF CRCs (Supplemental Figure 10). Thus, our data indicate that the phenomenon observed for ERf and SWI3B is not limited to Arabidopsis but rather may be a general feature of SWI/SNF CRCs and membrane receptors.

358

359 *ERECTA* and *SWI3B* Interact Genetically and *er/elr1/erl2* Plants Exhibit Alteration in 360 Chromatin Status.

- 361 The er-105/swi3b-3 double mutant exhibited more severe phenotypic traits than both single er-
- 362 *105* and *swi3b-3* mutant lines (Figure 4A), supporting the observed physical interdependences
- between ER and SWI3B.



365 366 Figure 4. ER and SWI3B interact genetically and affect both GA biosynthesis and response 367 pathways (See also Figures S11 and S12). A, The er-105/swi3b-3 double mutant exhibits more 368 retarded growth than the er-105 and swi3b-3 (three-weeks old plants). Graphical alignment of 369 corresponding leaves. Scale bar= 1 cm. B, The hypersensitivity of 1-week-old *swi3b-3* hypocotyl 370 to GA treatment is abolished by introducing er-105. C, Roots of all tested 1-week-old genotypes similarly respond to PAC treatment (error bars-SD, *P < 0.01,** P < 0.001, ***P<0.0001, 371 372 Student's t-test). D, Hypocotyls of all tested 1-week-old genotypes similarly respond to PAC 373 treatment, right panel- hypocotyl length comparison for PAC treated plants only (error bars-SD. 374 *P < 0.01,** P < 0.001, ***P<0.0001 Student's *t*-test). E, *swi3b-3* weak, point mutant line and 375 er-105/swi3b-3 exhibit elevated SWI3B transcript level, the SWI3B expression is elevated after 376 supplementation with bioactive GA₄₊₇ in all genotypes except *swi3b-3* (error bars-SD, P < 0.05, 377 Student's t-test). F, The examination of GID1 genes indicated that almost all examined lines 378 responded to GA treatment, but the swi3b-3 line was insensitive for GA-induced transcriptional 379 changes (error bars-SD, P < 0.05, Student's *t*-test). G, The examination of GA biosynthesis genes 380 indicated that almost all examined lines responded to GA treatment, but the swi3b-3 line was 381 insensitive for GA-induced transcriptional changes except GA200x2 expression (error bars-SD, P 382 < 0.05, Student's *t*-test).

19

383

384 The treatment with bioactive GA4+7 gibberellins indicated hypersensitivity of swi3b-3 to GA 385 demonstrated by hypocotyl length, while the response of *er-105* was reduced. By contrast, the 386 GA hypersensitivity of *swi3b-3* was abolished by introduced *er-105* mutation (Figure 4B). All 387 tested genotypes were responding to PAC treatment in a similar way (Figure 4C, D). The higher 388 expression of SWI3B was visible in the case of the swi3b-3 mutant, which was even more 389 pronounced in *er-105/swi3b-3*. The expression of *SWI3B* was elevated after supplementation with 390 bioactive GA_{4+7} in all genotypes, except for the *swi3b-3* line (Figure 4E). The examination of 391 GID1 and GA biosynthesis genes indicated that almost all examined lines responded to GA 392 treatment while the swi3b-3 line was insensitive for induced by GA transcriptional changes 393 except for GA200x2 expression (Figure 4F, G). Collectively, our results further indicate that both 394 ERf signaling and SWI3B-containing CRCs play together an important role in the fine-tuning of 395 GA signaling in Arabidopsis.

To verify the biological effect of observed interactions between ERf signaling and SWI3B-SWI/SNF, we analyzed the chromatin status, nuclei shape and chromocenters number in the *er/erl1/erl2* mutant plants. We found that *er/erl1/erl2* plants exhibit increased chromocenter number and altered spindle-like nuclei shape (Supplemental Figure 11A, B). We furthermore screened the effect of inactivation of ERfs on genome-wide nucleosome positioning in chromatin using *micrococcal nuclease* protection assays followed by deep sequencing (MNase-seq) and confirmatory MNase-qPCR in WT and *er/erl1/erl2* plants.

We found that inactivation of ERf proteins has a broad influence on the global nucleosomal chromatin structure in Arabidopsis- *erf* exhibited 41519 nucleosome occupancy changes, 13924 "fuzziness" changes and 4055 nucleosome position changes (Supplemental Figure 12A) and alterations in the presumable regulatory regions upstream of the transcription start site (TSSs) (Supplemental Figure 12B, Supplemental Dataset 1, Sub-Table 9-14).

Among genes with down-regulated expression and altered nucleosome positioning in the er/erl1/erl2 mutant were 14 GA-related genes (*ATBETAFRUCT4, XERICO, PRE1, MYBR1, MYB24, MIF1, HAI2, ZPF6, GA20ox1, CGA1, XTH24, GID1b, RGL1,* and *GIS3*) Interestingly, seven of them (*ATBETAFRUCT4, PRE1, MYBR1, MIF1, XTH24, GID1b,* and *RGL1*) were already observed to be directly targeted by the BRM ATPase of the SWI/SNF CRC (Archacki et al., 2016; Li et al., 2016).

20

An Integrated Gene Browser (IGB) view of *PRE1*, *GID1a,b* promoter regions indicated (Supplemental Figure 12D) various nucleosome alterations on promoter regions of these genes in the *er/erl/erl2* mutant pointing out impaired chromatin remodeling in the absence of functional ERf proteins. The selected changes were confirmed by MNase-qPCR (Supplemental Figure 12E).

419 The Inactivation of ERf proteins Affects SWI3B Protein Phosphorylation

420 We tested the ability of KDER to phosphorylate SWI3B protein. We overexpressed, 421 purified, and subsequently used MBP-His6-KDER and His6-SWI3B (Figure 5A, Supplemental 422 Figure 13A) for non-radioactive in vitro kinase assay. The existence of a strong band 423 corresponding to phosphorylated SWI3B protein and a weaker band of autophosphorylated 424 KDER was indicated (Figure 5B, Supplemental Figure 13B, C). The confirmatory mass-425 spectrometry analysis resulted in the identification of the active phosphorylation sites at KDER 426 and in the SWI3B protein (Supplemental Figure 13D, E). Interestingly three of four KDER-427 dependent phosphorylation sites were located in SWI3B in SWIRM and SANT domains 428 (Supplemental Figure 13E, F), providing a valuable hint that the ERf family proteins may be 429 responsible for the SWI3B phosphorylation.

bioRxiv preprint doi: https://doi.org/10.1101/2021.12.02.470991; this version posted December 3, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

21



431

432 Figure 5. ERf proteins are responsible for the phosphorylation of SWI3B protein, while DELLA 433 proteins control SWI3B protein abundance (See also Figures S13, S14, and S15). A, Coomassie 434 staining of MBP-His6-KDER and His6-SWI3B proteins purified from bacteria. B, Western blot 435 with anti-Thiophosphate ester antibody (ab92570; Abcam) showing in vitro SWI3B 436 phosphorylation by KDER. C. 2D Western blot assay with anti SWI3B antibody indicating in 437 vivo phosphorylation alteration of SWI3B protein in er/erl1/erl2 mutant. D, SWI3B and RGA 438 and RGL1 proteins in the nuclei of living cells. Bimolecular Fluorescence Complementation 439 assay (BiFC) in epidermis of tobacco leaves. Scale bar = $10\mu m$. E, The amounts of SWI3B and 440 RGA proteins in plants are oppositely regulated by PAC treatment. F, The disappearance of 441 SWI3B protein is PAC-dose dependent. G, The PAC-dependent degradation of SWI3B is 442 abolished by the MG132 treatment, a known proteasome inhibitor. H, The gal-3 mutant 443 constitutively accumulating DELLA proteins exhibits the decreased level of SWI3B, which is 444 restored to WT levels upon GA treatment. I, The triple DELLA mutant exhibits a WT-like level 445 of SWI3B protein, and the PAC treatment does not influence SWI3B level in this background. J, 446 Schematic model highlighting ERf and DELLA impact on the SWI3B protein.

22

To verify this possibility the *in vivo* phosphorylation analysis using 2D-IEF-PAGE combined with the Western blot was performed. The alteration in the SWI3B proteins isoelectric point (*pI*) in *er/erl1/erl2* mutant was observed. The bands corresponding to phosphorylated SWI3B form were nearly absent in *er/erl1/erl2* plants, indicating a severe defect in SWI3B phosphorylation (Figure 5C). This data strongly supports the regulatory function of the ERf proteins on the SWI3B subunit of SWI/SNF CRC.

454

Accumulation of DELLA Proteins Correlates with Increased Proteasomal Degradation of SWI3B

457 Our previous study demonstrated that SWI3C, a partner of SWI3B, physically interacts 458 with DELLA proteins (Sarnowska et al., 2013). We also found that the Arabidopsis lines with 459 impaired SWI/SNF CRCs- brm and swi3c exhibit decreased level of bioactive GA₄ gibberellins 460 level (Sarnowska et al., 2013; Archacki et al., 2013), but they do not accumulate RGA DELLA 461 protein similarly as in case of *er/erl1/erl2* plants (Supplemental Figure 14 and Figure 1J). To 462 address this unusual phenomenon, we used a BiFC assay to analyze the interaction between 463 DELLA and SWI3B. The interaction between either RGA or RGL1 protein and SWI3B was 464 found (Figure 5D, Supplemental Figure 15). No YFP signal was detected in control cells.

465 To understand the functional consequences of the detected interactions between SWI3B 466 and DELLA proteins, we analyzed the amounts of SWI3B and RGA proteins in GA or PAC-467 treated plants (Figure 5E). Surprisingly, we observed the PAC-dose-dependent disappearance of 468 SWI3B protein (Figure 5F). To check if the degradation of SWI3B under these conditions 469 depended on the proteasome, we tested the effect of MG-132 on the SWI3B level. MG-132 470 treatment caused increasing SWI3B abundance in PAC treated plants (Figure 5G), suggesting 471 that the degradation of SWI3B observed in parallel to accumulation of DELLAs occurs via the 472 proteasome. We also observed increased degradation of SWI3B in the gal-3 mutant in which 473 DELLA proteins are constitutively accumulated (Figure 5H), but we did not observe enhanced 474 SWI3B degradation in PAC-treated 3xDELLA (Archacki et al., 2013) collectively suggesting 475 that binding of SWI3B by DELLA proteins may be a primary cause of its proteasomal 476 degradation (Figure 5I, J). Thus, the accumulation of DELLA proteins should lead to the same 477 consequences as the elimination of SWI3B protein or SWI3B-containing SWI/SNF CRCs. This 478 conclusion is strongly supported by the lack of RGA protein accumulation in GA deficient brm

23

479 and *swi3c* lines with inactivated other subunits of SWI/SNF CRC (Archacki et al., 2013; 480 Sarnowska et al., 2013). Therefore, it could be indeed expected that GA and SWI3B-deficient 481 *er/erl1/erl2* mutant will also not accumulate RGA protein because, in the case of SWI/SNF CRC 482 impairment, the DELLA accumulation seems to be irrelevant. Collectively, our results provide 483 new insight into the functioning of the ERECTA family proteins and DELLA proteins and their 484 mutual impact on the SWI3B-containing SWI/SNF CRCs.

485

486 Binding of ERECTA and SWI3B to Promoter Regions of the *GID1* Genes

487 The *er/erl1/erl2* mutant displays an impaired response to exogenous GA treatment and a 488 consistently decreased expression of all three GID1 genes. ERf proteins interact with the SWI3B, 489 inactivation of *ERf* proteins results in nucleosomal chromatin structure alterations and decreased 490 abundance of SWI3B and its phosphorylated form in Arabidopsis, and there is an intriguing 491 interdependence between the control of the SWI3B level and DELLA protein accumulation, 492 therefore we examined if ER and the SWI3B participate in transcriptional control of the GID1 493 genes previously reported as targets for DELLA (Rosa et al., 2015). The ChIP analysis on GID1 494 promoters was performed using nuclei purified from 3-week-old seedlings expressing the ER-495 GFP and the SWI3B-HIS-STREP-HA proteins.

The binding of ER-GFP was detected around -130bp upstream of the TSS in the *GID1a* promoter region while SWI3B bound around the TSS of the *GID1a* promoter (Figure 6A). The ER protein was targeted to two regions around -150bp and -350bp from the TSS in the *GID1b* promoter, SWI3B was localized only -350bp upstream TSS (Figure 6B). ER and SWI3B were similarly cross-linked to the -100bp region of the *GID1c* promoter, but SWI3B was also mapped further upstream to -800bp (Figure 6C).

bioRxiv preprint doi: https://doi.org/10.1101/2021.12.02.470991; this version posted December 3, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.



504 Figure 6. ERf proteins enter the nucleus where ERECTA protein binds the GID1 promoters 505 similarly to the SWI3B subunit of SWI/SNF CRC (See also Figures S16 and S17). A, ERECTA 506 protein binds to promoter regions of the GID1a gene in a region targeted by the SWI/SNF 507 complex in three-week-old plants. (error bars refer to SD, P < 0.05, Student's *t*-test, three 508 biological and technical replicates were performed). B, ERECTA and SWI3B core subunit of 509 SWI/SNF CRCs target promoter regions of GID1b gene in three-weeks old plants (error bars 510 refer to SD, P < 0.05, Student's *t*-test, three biological and technical replicates were performed). 511 C, SWI3B binds to the promoter region of the *GID1c* gene in two different regions. One of them 512 is targeted by ERECTA protein in three-week-old plants. D, ERECTA protein binds to promoter 513 regions of the *GID1a* gene in a region targeted by the SWI/SNF complex in five-week-old plants. 514 (error bars refer to SD, P < 0.05, Student's *t*-test, three biological and technical replicates were 515 performed). E, ERECTA targets promoter region of *GID1b* gene in five-week-old plants (error 516 bars refer to SD, P < 0.05, Student's *t*-test, three biological and technical replicates were 517 performed). F, ERECTA binds to the promoter region of the GID1c gene in three-week-old 518 plants. The bottom panel in D-F: the binding of native SWI3B protein to its target sites in GID1a-519 c promoter regions is abolished in 5-week-old er/erl1/erl2 triple mutant plants. G, A model 520 describing the non-canonical nuclear function of ERf proteins in the GA signaling pathway.

522 Inspection of ER and SWI3B binding to the GID1a-c promoter regions in 5-week-old 523 WT, ER-GFP, and the *er/erl1/erl2* mutant demonstrated that ER binds the same GID1 promoter 524 regions as in the case of 3-week-old plants (Figure 6 D-F). The SWI3B binding to the promoter 525 of GID1 genes was abolished by the inactivation of ERf proteins in er/erl1/erl2 plants. The 526 inactivation of ER, ERL1, or ERL2 did not affect the binding of SWI3B to GID1a-c promoter 527 regions in single er105, erl1, and erl2 mutant lines (Supplemental Figure 16). Our study provides 528 evidence that the three ERf proteins have redundant functions regarding proper SWI3B 529 recruitment since only the simultaneous absence of all ERfs proteins abolished SWI3B binding to 530 the *GID1a-c* promoters.

531 Of note, we found a similar binding of HER2 (EGFR-family) receptor and BAF155 532 subunit of SWI/SNF CRCs to the promoter regions of human *BRCA1* and *FBP1* genes 533 (Supplemental Figure 17), indicating that the phenomenon observed for ER may be a general 534 mechanism controlling gene expression that is maintained between kingdoms.

⁵²¹

26

536 Discussion

537 Inactivation of ERf LRR-RLK family members results in various defects in Arabidopsis 538 growth and development. While it is well established that ERf proteins play distinct roles in the 539 control of epidermal patterning, stomatal development, meristem size, inflorescence architecture, 540 and hormonal signaling, the exact mechanisms underlying the regulatory functions of ERf 541 proteins in these processes are largely unknown (*e.g.*, Chen and Shpak, 2014; Chen et al., 2013b; 542 Qi et al., 2004; Van Zanten et al., 2010; Kosentka et al., 2019).

543 Here we show that the inactivation of Arabidopsis ERf proteins has a broad effect on 544 various regulatory processes, including hormonal signaling, and suggest that these sum responses 545 underlie the severe developmental defects exhibited by the er/erl1/erl2 mutant. We demonstrate 546 that parallel inactivation of all *ERf* proteins results in severe deregulation of the GA signaling 547 pathway as evidenced by the impairment of GA perception and GA biosynthesis. When taken 548 together, these findings, alongside the identification of NLS and NES sequences in ERf proteins 549 and our demonstration of their translocation into the nucleus, suggest a novel, non-canonical 550 function of ERf proteins (Figure 6G).

551 Our study also reveals an analogy of this system to the previously described XA21 LRR 552 immune receptor in rice (Park and Ronald, 2012) and to the non-canonical signaling mode of the 553 human Epidermal Growth Factor Receptor (EGFR) family (Lee et al., 2015a). Although it should 554 be stressed that these two classes of plant and animal epidermal receptors carry completely 555 unrelated sequences from one another and from the system we describe here (Supplemental Table 556 3), suggesting that the translocation of the membrane receptors to the nucleus may be a general 557 paradigm maintained between plant and animal kingdoms. In addition to their canonical 558 membrane receptor functions, holoreceptor and truncated forms of EGFRs are imported into 559 nuclei via ER-mediated retrograde transport, although some of them lack known NLSs (Chen and 560 Hung, 2015).

In the nucleus, the EGFR receptors can bind to DNA, interact with various transcription factors. Thereby, nuclear forms of EGFRs are implicated in the control of cell proliferation, DNA replication and repair, and transcription (Chen and Hung, 2015), so their functions extend far beyond the regulation of epidermal patterning.

565 We found here that the Arabidopsis ERECTA LRR-RLK receptor similarly translocates 566 from the plasma membrane into the nucleus. Both intact and N-terminally truncated forms of

27

567 ERECTA were detectable in the nucleus. A truncated ERECTA carrying only the kinase domain 568 localizes exclusively in the nucleus and partially complements the leaf developmental defects 569 caused by the *er-105* mutation implying a ligand-independent non-canonical signaling function of 570 the ERECTA kinase domain.

571 Furthermore, our data show that the ERf proteins interact through their kinase domains 572 with the SWI3B, and ER can phosphorylate the SWI3B core subunit of the SWI/SNF CRC. 573 SWI/SNF plays a pivotal role in the hormonal crosstalk regulation in both humans and plants 574 (Sarnowska et al., 2016). Moreover, we show that analogously as ER protein, the HER2 member 575 of the EGFR family directly interacts with the BAF155 subunit of human SWI/SNF and co-576 localizes with BAF155 on some gene promoters providing evidences that such system is likely 577 maintained between kingdoms.

578 Parallel inactivation of all ERf proteins results in alterations of genome-wide nucleosomal 579 chromatin structure and altered transcriptional activity of a large number of genes. The binding of 580 SWI3B to its target regions in the GID1a-c promoters is retained in er-105, erl1, and erl2 single 581 mutant lines. By contrast, the er/erl1/erl2 mutant plants exhibited a reduction in phosphorylation 582 SWI3B protein level and abolished proper SWI3B binding to GID1 promoter regions together 583 with decreased expression of GID1a-c genes, indicating a strong and direct effect of the ERf 584 signaling pathway on SWI/SNF-dependent chromatin remodeling. The er/erl1/erl2 mutant plants 585 are characterized by a decreased level of endogenous gibberellins; however, they do not 586 accumulate DELLA proteins, similar to the SWI/SNF mutants. Thus, we have demonstrated that 587 the *er/erl1/erl2* mutant plants exhibit severe deregulation of the gibberellin signaling pathway 588 and SWI/SNF-dependent chromatin remodeling. This is, in turn, an attractive explanation of the 589 observed insensitivity of *er/erl1/erl2* mutant to the application of exogenous gibberellin. DELLA 590 proteins are involved in the sequestration of various transcription factors and chromatin 591 remodeling complexes (Phokas and Coates, 2021). In this study, we extend the existing 592 knowledge on DELLA functioning by providing evidence for the existence of the DELLA-593 SWI3B regulatory module and explaining why some GA-deficient mutant lines with impaired 594 SWI/SNF chromatin remodeling complex do not accumulate DELLA proteins (Figure 5G).

595 Collectively our finding that plant ERf proteins play an important, non-canonical nuclear 596 function, *i.e.*, bind directly to chromatin and control the proper recruitment of SWI/SNF CRCs, 597 which are strongly involved in controlling regulatory processes including hormonal crosstalk

28

- 598 (Sarnowska et al., 2016), may be a general paradigm for other classes of plant and mammalian
- 599 membrane receptor kinases.

29

601 Methods

602 Plant Material and Growth Conditions

603 The Arabidopsis thaliana ecotype Columbia was used as wild type (WT) in all 604 experiments. The following Arabidopsis mutants were used for analysis: er-105, er-105/swi3b-3 605 (Sáez et al., 2008), er/erl1/erl2 plants (Shpak et al., 2004) and erf lines in various combinations 606 (Torii et al., 1996), the 35S::GFP Arabidopsis line has been obtained from NASC (N67775). 607 Seeds were sown on soil or plated on ¹/₂ Murashige and Skoog medium (Sigma-Aldrich) 608 containing 0.5% sucrose and 0.8% agar. Plants were grown under long day (LD) condition (12h 609 Day/12h Night or 16h Day/8h Night). For GA response tests, plants were sprayed twice a week 610 with 100 µM GA₄₊₇ or water (control) for a fast response the 2h of GA₄₊₇ treatment was 611 performed.

612 **Construction of Transgenic Lines**

613 Genomic sequences of ERECTA, cDNAs of ERECTA kinase domain and truncated 614 ERECTA lacking kinase domain (ER Δ K) were cloned into binary vector p35S::GW::GFP (F. 615 Turck, Max-Planck-Institut für Züchtungsforschung, in the case of ERECTA), and into pEarley 616 Gate 101 (in the case of the ER kinase domain and $\Delta KDER$; (Earley et al., 2006). Plants were 617 transformed using Agrobacterium tumefaciens GV3101 (pMP90) by floral-dip method (Davis et 618 al., 2009). The STOP codon of the SWI3B genomic sequence was replaced with HIS-STREP-HA 619 using the recombineering method (Bitrián et al., 2011), moved into pCB1 vector (Heidstra et al., 620 2004), and transformed into swi3b-2 Arabidopsis mutant line.

621 **RNA Extraction and qRT-PCR Analysis**

622 Total RNA was isolated from adult (5-week-old) plants using an RNeasy plant kit 623 (Qiagen), treated with a TURBO DNA-free kit (Ambion). Total RNA (2.5µg) was reverse 624 transcribed using a first-strand cDNA synthesis kit (Roche). qRT-PCR assays were performed 625 with SYBR Green Master mix (Bio-Rad) and specific primers for PCR amplification. 626 Housekeeping genes PP2A and UBQ5 (AT1G13320 and AT3G62250, respectively) were used as controls. The relative transcript level of each gene was determined by the $2^{-\Delta\Delta Ct}$ method 627 628 (Schmittgen and Livak, 2008). Each experiment was performed using at least three independent 629 biological replicates. qRT-PCR primers are listed in Supplemental Dataset 3.

630 Transcript Profiling and Gene Ontology Analysis

30

RNA was isolated from adult (5-week-old) WT and *er/erl1/erl2* plants using a Plant RNeasy kit (Qiagen) according to the manufacturer's protocol. Transcriptomes were analyzed using 150ng of total RNA as starting material. Targets were prepared with a cDNA synthesis kit followed by biotin labeling with the IVT labeling kit (GeneChip 39IVT Express; Affymetrix) and hybridized to the ATH1 gene chip for 16h as recommended by the supplier. The raw data were analyzed using GenespringGX according to the manual (guided workflow). GO-TermFinder was used for GO analyses of selected groups of genes (Boyle et al., 2004).

638 Nuclear Fractionation

Nuclei were isolated from 2g of leaves of 3-weeks old Arabidopsis seedlings according to
the method previously described by Gaudino and Pikaard (1997). Subsequent nuclear
fractionation was performed using the high-salt method, with modifications (Sarnowski et al.,
2002).

643 Protein Interaction Study, Confocal Imaging, Subcellular Localization, Brefeldin A and

644 Leptomycin B Treatment, DAPI Staining

Protein interaction was analyzed by performing the immunoprecipitation of ER-GFP or KDER-YFP-HA from nuclei from 4 g of Arabidopsis plants (Saleh et al., 2008). The nuclear extracts were incubated with 25 μ L of GFP Magnetic Trap beads (Chromotek) according to manufactures instructions. The presence of SWI3B protein was determined by western blot analysis using anti-SWI3B antibody (Sarnowski et al., 2002).

The interaction between human proteins was analyzed by immunoprecipitation of HER2
and BAF155 from viscolase treated nuclear extracts prepared, according to Jancewicz et al. 2021.
The presence of HER2 and BAF155 was determined by Western blot analysis using anti HER2
(CST, 12760) and anti BAF155 (CST, 11956) antibodies.

To obtain YFN-ERL1, YFC-ERL1, YFC-ERL2, and YFC-KDER fusions for BiFC (Hu et al., 2002) analysis, cDNAs encoding ERL1 and ERL2 proteins and ERECTA, ERL1, and ERL2 C-terminal kinase domains were PCR amplified and cloned into the binary vectors pYFN43 or pYFC43 (Belda-Palazón et al., 2012). The *in vivo* interactions between proteins were detected by BiFC using Leica TCS SP2 AOBS, a laser scanning confocal microscope (Leica Microsystems). Tobacco (*Nicotiana benthamiana*) epidermal cells were infiltrated using *Agrobacterium*

tumefaciens GV3101 (pMP90) carrying plasmids encoding ERL1, ERL2, or KDER fusions and the p19 helper vector and analyzed by confocal microscopy 3 d later. YFN-RFP and YFC-RFP

31

fusions were used to detect transformed cells in the BiFC assays (Sarnowska et al., 2013); at least
five nuclei were analyzed in three separate experiments.

664 The vesicle trafficking inhibitor BFA (Sigma Aldrich) was used at the 25µM 665 concentration at the following time points 40 min, 90 min, and 120 min. The NES-dependent 666 nuclear export inhibitor Leptomycin B was used at the 200 nM concentration 4h before 667 microscopy observation. Nuclei were stained with 4',6-diamidino-2-phenylinodole (DAPI) at the 668 μ g/mL concentration for 30 min. The observation was carried out on the root tip of about two 669 weeks old plants incubated directly before in $\frac{1}{2}$ MS alone or with the addition of proper 670 compound (BFA or Leptomycin B, respectively). Every time 30 min before the end of 671 incubation, DAPI was added.

672 Chromatin Immunoprecipitation

673 ChIP experiments were performed as described previously (Sacharowski et al., 2015) on 674 three or five-week-old WT, ER-GFP, SWI3B-His-Strep-HA, and er/erl1/erl2 plants cross-linked 675 under vacuum using formaldehyde (final concentration: 1%) and Bis-(sulfosuccinimidyl) 676 glutarate (final concentration: 1mM). For ER-GFP, chromatin immunoprecipitation was 677 performed with GFP-Trap M (Chromotek). For SWI3B ChIP experiments, NiNTA Agarose 678 (Oiagen) or, in the case of anti-SWI3B antibody, the magnetic protein A and G dynabeads 679 (Dynal) were used. ChIP enrichment was determined using qPCR, and relative fold change was calculated using the $2^{-\Delta\Delta Ct}$ method (Schmittgen and Livak, 2008). The TA3 retrotransposon was 680 681 used as negative control (Pastore et al., 2011). Primers used in ChIP experiments are listed in 682 Supplemental Dataset 3.

Chromatin from the SKBR-3 human cell line was immunoprecipitated according to (Komata et al., 2014 and Jancewicz et al., 2021). Recovered chromatin was incubated O/N at 4°C with the following antibodies: anti BAF155 (CST, 11956), antiHER2 (CST, 12760), and Normal Rabbit IgG (CST, 2729, mock control). Results were calculated based on the $2^{-\Delta\Delta Ct}$ (Schmittgen and Livak, 2008). The relative fold enrichment of the analyzed sample represents the fold change with reference to IgG (mock) sample. A set of primers used for ChIP-qPCR analysis is listed in Supplementary Dataset 3.

690 MNase Mapping of Genome-wide Nucleosome Positioning and MNase-qPCR

32

The nuclear extraction, MNase treatment, subsequent NGS analyses, and confirmatory
 MNase-qPCR were performed according to (Sacharowski et al., 2015) on 5 week old plant
 material.

694 Gibberellin Analysis

About 200 mg of frozen materials from 5 week old plants were used to extract and purify the GA as described in Plackett et al. (2012) with minor modifications. GA was quantified using MS/MS analysis using 4000 Triple Quad (AB Sciex Germany GmbH, Darmstadt, Germany) in multiple reaction monitoring (MRM) scan with electrospray ionization (ESI) as described in Salem et al., (2016). The mass spectrometry attached to UPLC system (e.g., Waters Acquity UPLC system, Waters, Machester, UK) separation was achieved on a reversed phase C18-column (100 mm × 2.1 mm 1.8 μm).

702 In vitro Phosphorylation Analysis and in vivo Kinase Assay

SWI3B-6xHis was overexpressed and purified, according to Sarnowski et al. (2002). The
KDER (pDEST-6xHis-MBP vector) was purified using tandem purification using MBP and NiNTA resins. *In vitro* kinase assay was performed according to the method described by (Allen et al., 2007). Phosphorylation was detected by Western blot analysis using an anti-Thiophosphate
ester antibody (ab92570; Abcam) and by the MS/MS analysis.

In vivo phosphorylation analysis was performed using a 2D western blot assay on nuclear 708 709 extracts from 5 weeks old WT and er/erl1/erl2 plants (Saleh et al., 2008). For isoelectrofocusing 710 (IEF), nuclear proteins were prepared according to Kubala et al. (2015). The IEF was performed 711 on the 7cm length gel strips with immobilized pH gradients 3-10 and 3-6 (BioRad). After IEF, 712 the equilibration of immobilized pH gradient was performed according to Wojtyla et al. (2013). 713 The SWI3B protein was detected by Western blotting using an anti SWI3B antibody (Sarnowski 714 et al., 2002). The *in vivo* phosphorylation was identified based on the changes of SWI3B 715 isoelectric point (pl) (Mayer et al., 2015).

716 Accession Numbers

Microarray and MNase-seq data are available in the ArrayExpress database
(www.ebi.ac.uk/arrayexpress) under E-MTAB-5595 and E-MTAB-5830 accession numbers,
respectively.

33

721 Supplemental Data

722 723

724 **Supplemental Figure 1.** Comparative analysis of genes with altered expression in *er/erl1/erl2* 725 and gal-3 mutants. A. Combinations of erf mutants have distinct effects on height of Arabidopsis 726 plants. Error bars refer to SD,* = P < 0.05, Student's t test. B, Genes showing altered expression 727 in the er/erl1/erl2 mutant plants. C, Venn diagrams indicating genes contrastingly up-regulated in 728 er/erl1/erl2 and down-regulated in ga1-3 plants. D, Venn diagrams indicating genes contrastingly 729 down-regulated in *er/erl1/erl2* and up-regulated in *ga1-3* plants. E, ERf proteins control the 730 expression of genes down-regulated in gal-3, which are either DELLA repressed or DELLA 731 independent. F, ERf proteins control the expression of genes up-regulated in gal-3, which are 732 either DELLA activated or DELLA independent.

733

734 Supplemental Figure 2. The *er/erl1/erl2* mutant displays severely impaired response to 735 exogenous GA treatment and slightly enhance gal-3 phenotypic traits. A, The 14 days old er/erl1/erl2 plants treated with GA4+7 did not show rosette expansion indicating defects in GA 736 737 response. Error bars refer to SD,* =P < 0.05, Student's t test, n= 30 plants. B, Two-month-old 738 GA₄₊₇ treated *er/erl1/erl2* plants show accelerated flowering compared to mock treated control. 739 Scale bar= 1cm. n= 30 plants. C, Two-months old er/erl1/erl2 plants. Scale bar= 1cm. n= 30 740 plants. D, Three-weeks old gal-3 plants crossed with erf mutants in various combinations show 741 mostly the phenotypic traits characteristic for gal-3. Scale bar= 1 cm. E, The phenotypic traits of 742 5-weeks old plants carrying erf mutations crossed with gal-3 in various combinations grown in 743 long day conditions. Scale bar= 1 cm.

744

745 Supplemental Figure 3. *er/erl1/erl2* plants exhibit deficiency in gibberellin intermediates.

Left panel: *er/erl1/erl2* mutant exhibits dramatically reduced level of GA_{12} and GA_{24} gibberellin intermediates (error bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates were assayed). Right panel: schematic representation of alteration in GA biosynthesis pathway in *er/erl1/erl2* plants.

750

751 Supplemental Figure 4. 35S::ERECTA-GFP construct complements the *er-105* mutation.

34

Supplemental Figure 5. ERECTA protein is detected in various cell compartments including
endosomes and accumulate in the nuclei periphery after BFA treatment. A, In cell pairs of
stomata, ERECTA-GFP was detected in circles around the positions of nuclei. Scale bar=10µm.
B, Accumulation of ERECTA protein in the BFA bodies after Brefeldin A treatment. Note the
enhanced presence of BFA bodies after 40 min (mid column) and 90 min (right column) BFA
treatment. Scale bar=10µm.

759

760 Supplemental Figure 6. The ERL1 and ERL2 proteins carry defined NLS in their kinase 761 domains. A, The NLS prediction in the kinase domain of ERL1 and ERL2 proteins has been done 762 using cNLS mapper (Kosugi et al., 2009a; Kosugi et al., 2009b). NLS score in range 5-7 means 763 that protein is partially localized in the nucleus and cytoplasm. Bottom panel: The alignment of 764 ERECTA, ERL1, and ERL2 protein sequences (part of kinase domains carrying NLS) using 765 PRALINE indicates high amino-acid sequence conservation between analyzed proteins. 766 Consistency is determined within range 1-10, where 1 means least conserved substitution and 10-767 the most conserved substitution (Simossis et al., 2005). B, Western blot analysis with anti-GFP 768 antibody confirms nuclear localization of ERECTA protein which undergoes proteolytic 769 processing. The samples were standardized by western blotting with anti H3 antibody. C, 770 Schematic presentation of full length and deletion variants of ERECTA protein used for the 771 localization study. ER - ERECTA protein with complete amino acids sequence; $\Delta KDER$ -772 truncated ERECTA protein lacking kinase domain; KDER - the kinase domain of ERECTA 773 protein.

774

775 Supplemental Figure 7. The kinase domain of ERECTA (KDER) has ability to complement the 776 er-105 leaf phenotypic traits. A, Rosette leaves of WT (upper), er-105 (mid), and er-105/KDER-777 YFP (lower panel). Graphical alignment of corresponding leaves indicating partial 778 complementation of er phenotypic traits by KDER. Scale bar= 1cm. B, Cauline leaves of WT 779 (upper), er-105 (mid), and er-105/KDER-YFP (lower panel). Graphical alignment of 780 corresponding laves indicating partial complementation of *er* phenotypic traits by KDER. Scale 781 bar= 1cm. C, The kinase domain of ERECTA cannot restore all (i.e., stem elongation) phenotypic 782 traits of the *er-105* mutant line. Scale bar= 1cm.

35

Supplemental Figure 8. ERECTA protein enters to the nucleus and localizes in various subnuclear fractions.

786

787 **Supplemental Figure 9.** Negative controls for bimolecular fluorescence complementation assay.

788 Negative controls for BiFC interaction analysis of ER, ERL1, ERL2 kinase domains fused to

789 YFC and SWI3B fused to YFN, including the RFP channel. Scale bar 10 μ m.

790

Supplemental Figure 10. Human SWI3-type BAF155 co-precipitates with HER2 EGFR family
 membrane receptor from human cells nuclei.

793

Supplemental Figure 11. *er/elr1/erl2* mutant plants show affected chromatin organization
 demonstrated as altered chromocenters number.

796 Upper panel: exemplary pictures of WT and *er/erl1/erl2* nuclei.

797 Lower panel: calculation of chromocenters (n=20 nuclei for each genotype).

798

799 Supplemental Figure 12. ERf proteins inactivation has a severe impact on genome-wide 800 nucleosome positioning. A, Nucleosome changes identified in the er/erl1/erl2 triple mutant 801 plants. B, Genome-wide nucleosome distribution patterns surrounding the transcription start site 802 (TSS). C, Nucleosome distribution patterns surrounding the TSS of GA-related genes showing 803 altered expression in the er/erl1/erl2 triple mutant plants. D, The alteration of nucleosomal 804 structure on *PRE1*, *GID1a*, and *GID1b loci* misexpressed in the *er/erl1/erl2* triple mutant plants 805 and targeted by the SWI/SNF CRC. Red boxes indicate nucleosome alterations. E, Confirmatory 806 MNase-qPCR for selected genes with altered nucleosomes.

807

808 Supplemental Figure 13. Kinase domain of ERECTA phosphorylates SWI3B protein. A, 809 Western blot with anti His6 antibody for detection of MBP-His6-KDER and His6-SWI3B 810 proteins purified from bacteria. B, Western blot with anti-Thiophosphate ester antibody 811 indicating no phosphorylation of SWI3B protein in the absence of KDER (negative control). C, 812 Western blot with anti-Thiophosphate ester antibody (ab92570; Abcam) showing 813 autophosphorylation of KDER in the absence of SWI3B protein. D, Identification of active 814 phosphorylation sites in KDER by MS/MS analysis. E, Identification of active phosphorylation

36

sites in SWI3B by MS/MS analysis. F, KDER phosphorylates SWI3B at the SWIRM and SANTdomains.

817

818 Supplemental Figure 14. GA-deficient mutant lines with inactivated subunits of SWI/SNF 819 complexes do not accumulate RGA DELLA protein. A, GA-deficient *swi3c* plants are unable to 820 over accumulate RGA protein. B, GA-deficient *brm* plants are unable to over accumulate RGA 821 protein.

822

823 Supplemental Figure 15. Negative controls for bimolecular fluorescence complementation824 assay.

825 Negative controls for BiFC interaction analysis of SWI3B, RGA, and RGL1 fused to YFC, and

SWI3B fused to YFN, including the RFP channel. Scale bar 10 μ m.

827

828 Supplemental Figure 16. Proper SWI3B binding to GID1a-c promoter regions is abolished in 829 the *er/erl1/erl2* whereas occurs in single *er-105*, *erl1* or *erl2* mutant lines. A, SWI3B targets 830 promoter region of GID1a gene in five-weeks old plants WT, er-105, erl1 or erl2 but is abolished 831 in triple *er/erl1/erl2* mutant plants (error bars refer to SD, P < 0.05, Student's t test, three 832 biological and technical replicates were used). B, SWI3B targets promoter region of GID1b gene 833 in five-weeks old plants WT, er-105, erl1 or erl2 but is abolished in triple er/erl1/erl2 mutant 834 plants (error bars refer to SD, P < 0.05, Student's t test, three biological and technical replicates 835 were used). C, SWI3B targets promoter region of GID1c gene in five-weeks old plants WT, er-836 105, erl1 or erl2 but is abolished in triple er/erl1/erl2 mutant plants (error bars refer to SD, P < 837 0.05, Student's t test, three biological and technical replicates were used).

838

Supplemental Figure 17. Human SWI3-type BAF155 targets together with HER2 EGFR family
membrane receptor *FBP1* and *BRCA1* genes *loci*. A, BAF155 subunit of human SWI/SNF
complex binds *Fructose-1,6-Bisphosphatase locus* together with HER2 member of EGFR
membrane receptor family. B, BAF155 subunit of human SWI/SNF complex binds *BRCA1 locus*together with HER2 member of EGFR membrane receptor family.

- 845 Supplemental Table 1. Genes classified to "Response to Gibberellin" GO term and showing
- 846 down-regulated expression level in the *er/erl1/erl2* mutant.
- 847
- 848 Supplemental Table 2. Genes with up-regulated expression in *er/erl1/erl2* mutant plants
- classified to GO-terms of leaf epidermal and stomatal cell differentiation.
- 850
- 851 Supplemental Table 3. Functional analogies between arabidopsis ERf proteins and the human
- EGFR membrane receptors.
- 853
- 854 **Supplemental dataset 1.** Comparative analysis of transcript profiling and MNase-seq data.
- 855 **Sub-table 1.** Transcript profiling using ATH1 microarray analysis to identify genes down-856 regulated in *er/erl1/erl2* mutant line.
- 857 **Sub-table 2.** Transcript profiling using ATH1 microarray analysis to identify genes up-regulated
- 858 in *er/erl1/erl2* mutant line.
- 859 **Sub- table 3.** GO analysis of genes down-regulated in *er/erl1/erl2* mutant line.
- 860 **Sub- table 4.** GO analysis of genes up-regulated in *er/erl1/erl2* mutant line.
- 861 **Sub-table 5.** Comparative transcript profiling analysis for genes down-regulated in *er/erl1/erl2* 862 and *ga1-3* mutants lines.
- 863 **Sub-table 6.** GO analysis of *er/erl1/erl2* and *ga1-3* down-regulated genes.
- 864 **Sub-table 7.** Comparative transcript profiling analysis for genes up-regulated in *er/erl1/erl2* and
- 865 gal-3 mutants lines.
- 866 **Sub-table 8.** GO analysis of genes up-regulated in *ga1-3* and *er/erl1/erl2*.
- 867 **Sub-table 9.** Genes with altered nucleosome positioning in promoter region -3000 to TSS.
- Sub-table 10. GO analysis of genes with altered nucleosome positioning identified in promoter
 region -3000 to TSS.
- 870 Sub-table 11. Genes with altered nucleosome positioning identified in promoter region -3000 to
 871 TSS and down-regulated in *er/erl1/erl2* microarray.
- 872 Sub-table 12. Genes with altered nucleosome positioning identified in promoter region -3000 to
 873 TSS and up-regulated in *er/erl1/erl2* microarray.
- 874 **Sub-table 13.** GO analysis for genes with altered nucleosome positioning identified in promoter
- 875 region -3000 to TSS and down-regulated in *er/erl1/erl2* microarray.
- 876 **Sub-table 14.** GO analysis for genes with altered nucleosome positioning identified in promoter
- 877 region -3000 to TSS and up-regulated in *er/erl1/erl2* microarray.
- 878

38

879 **Supplemental dataset 2.** Comparison of ER, ERL1, and ERL2 protein sequences with 880 highlighted important domains including NLS.

- 881 ClustalW was used to align ER, ERL1, and ERL2 sequences. Under conserved amino acid is
- asterisk mark, highly similar or similar amino-acids are marked :: and . respectively. LRR domain
- is marked as gray background, amino-acids crucial for EPFs or TMM interaction are indicated
- 884 with bold. Transmembrane domain, juxtamembrane domain, and kinase domain are indicated
- with bold green, orange, and blue color fonts, respectively (Kosentka et al., 2017). The predicted
- 886 NLS sequence is marked in a dotted line frame (Kosugi et al., 2009b).
- 887
- 888 **Supplemental Dataset 3.** Primers used in this work.
- 889 Supplemental Movie 1. The ERECTA protein undergoes endocytosis.
- 890 Supplemental Movie 2. The ERECTA protein undergoes endocytosis.

892 Acknowledgements

893 We thank Csaba Koncz for critical comments during manuscript construction, Dorota Zugaj for 894 assistance in plant cultivation, Iga Jancewicz for assistance in leptomycin B assay, Claus 895 Schwechheimer for providing the anti-RGA antibody, Mohammad-Reza Hajirezaei for help with 896 GA the Max Planck-Genome-Center measurements and Cologne 897 (http://mpgc.mpipz.mpg.de/home/) for performing the transcript profiling and MNase-seq 898 analysis described in this study.

40

900 FIGURE LEGENDS

901 Figure 1. ERf inactivation affects Arabidopsis development, causes transcriptomic changes 902 overlapping with the effect of gal-3 mutation and impairs GA biosynthesis and signaling (See 903 also Figures S1, S2 and S3). A, Phenotypic changes conferred by combinations of *erf* mutations. 904 Scale bar= 1 cm. B, Overlapping down-regulated genes in *er/erl1/erl2* and *ga1-3* plants. C, 905 Overlapping up-regulated genes in er/erl1/erl2 and ga1-3 plants. D, The er/erl1/erl2 plants 906 exhibit impaired GA response. 14- days old LD (12h day/12 night) grown WT and er/erl1/erl2, 907 sprayed twice a week with water (upper row) or 100µM GA₄₊₇ (lower row). Arrows-er/erl1/erl2 908 plants. Scale bar= 1cm. E, The GA response is retained to various levels in combinations of erf 909 mutants. Error bars-SD,* =P < 0.05, Student's *t*-test, n= 30 plants. F, The response of various *erf* 910 mutants to 1µM Paclobutrazol treatment. Error bars-SD,* =P < 0.05, Student's t test, n= 30 911 plants. G, The er/erl1/erl2 mutant exhibits altered transcription of GID1 GA receptor genes (error 912 bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates were assayed). H, 913 The er/erl1/erl2 mutant displays altered GA biosynthesis and metabolism-related genes 914 expression (error bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates 915 were assayed). I, The *er/erl1/erl2* mutant exhibits dramatically reduced level of bioactive GA_{4+7} 916 gibberellin (error bars-SD, P < 0.05, Student's *t*-test, three biological and technical replicates 917 were assayed). J, The *er/erl1/erl2* mutant shows decreased level of the DELLA protein RGA.

918

919 Figure 2. Subcellular localization of ERECTA protein (See also Figures S4 and S5). A, 920 ERECTA is localized in plasma-membrane and endosomes in epidermal cells of 7-days old 921 seedlings. ER-GFP, or free GFP visualized using GFP channel. FM4-64 specifically stains 922 plasma-membranes. Scale bar=10µm. B, Root-tip images of approximately two-week-old (14-17 923 days) ER-GFP seedlings showing nuclear localization of ERECTA protein at considerable 924 frequency. C, Root-tip images of 12-day-old ER-GFP seedlings serving as the control for D and 925 E. D, Brefeldin A treatment enhanced the localization of ERECTA protein in Brefeldin A (BFA) 926 bodies. Roots of 12-day-old Arabidopsis seedlings. E, Leptomycin B treatment enhanced the nuclear localization of ERECTA Free GFP was used as a control in C, D, and E, cell nuclei were 927 928 stained with DAPI, scale bar= 50µm. F, Letomycin B enhances nuclear presence of ER protein. 929 The GFP/DAPI ratio calculated per area for roots of plants expressing ER-GFP protein.

41

931 Figure 3. Nuclear function of ERf proteins (See also Figures S6, S7, S8, S9, and S10). A, Root-932 tip images of approximately two-week-old (14-17 days) plants expressing ER-GFP, KDER-YFP-933 HA (the kinase domain of the ER protein), ER Δ K-YFPHA (truncated ER protein lacking the 934 kinase domain) proteins indicating that the kinase domain is necessary for the nuclear localization 935 of ER protein. WT and GFP expressing plants-negative controls. Panel 2 in the ER-GFP indicates 936 nuclear localization of ER protein appearing at considerable frequency. Cell nuclei were stained 937 with DAPI. Scale bar=25 µm. B, The er-105/swi3b3 double mutant shows more retarded growth 938 than either er-105 or swi3b3 plants. Scale bar= 1cm. C, The er/erl1/erl2 triple mutant exhibits 939 reduced SWI3B protein level. D, ER-GFP or free GFP (negative control) pull-down from the 940 nucleus and anti-SWI3B western blotting indicate a specific ER-SWI3B interaction. E, 941 Immunoprecipitation of KDER-YFP-HA from the nucleus indicated that the kinase domain of 942 ER interacts with SWI3B. F, ER, ERL1, and ERL2 kinase domains interact with SWI3B in the 943 nucleus. Bimolecular Fluorescence Complementation assay (BiFC) in epidermis of tobacco 944 leaves. Scale bar = $10\mu m$.

945

946 Figure 4. ER and SWI3B interact genetically and affect both GA biosynthesis and response 947 pathways (See also Figures S11 and S12). A, The er-105/swi3b-3 double mutant exhibits more 948 retarded growth than the er-105 and swi3b-3 (three-weeks old plants). Graphical alignment of 949 corresponding leaves. Scale bar= 1 cm. B, The hypersensitivity of 1-week-old *swi3b-3* hypocotyl 950 to GA treatment is abolished by introducing er-105. C, Roots of all tested 1-week-old genotypes similarly respond to PAC treatment (error bars-SD, *P < 0.01,** P < 0.001, ***P<0.0001, 951 952 Student's t-test). D, Hypocotyls of all tested 1-week-old genotypes similarly respond to PAC 953 treatment, right panel- hypocotyl length comparison for PAC treated plants only (error bars-SD, *P < 0.01,** P < 0.001, ***P<0.0001 Student's *t*-test). E, *swi3b-3* weak, point mutant line and 954 955 er-105/swi3b-3 exhibit elevated SWI3B transcript level, the SWI3B expression is elevated after 956 supplementation with bioactive GA_{4+7} in all genotypes except *swi3b-3* (error bars-SD, P < 0.05, 957 Student's t-test). F, The examination of GID1 genes indicated that almost all examined lines 958 responded to GA treatment, but the *swi3b-3* line was insensitive for GA-induced transcriptional 959 changes (error bars-SD, P < 0.05, Student's *t*-test). G, The examination of GA biosynthesis genes 960 indicated that almost all examined lines responded to GA treatment, but the swi3b-3 line was

42

961 insensitive for GA-induced transcriptional changes except GA20ox2 expression (error bars-SD, P 962 < 0.05, Student's *t*-test).

963

964 Figure 5. ERf proteins are responsible for the phosphorylation of SWI3B protein, while DELLA 965 proteins control SWI3B protein abundance (See also Figures S13, S14, and S15). A, Coomassie 966 staining of MBP-His6-KDER and His6-SWI3B proteins purified from bacteria. B, Western blot 967 with anti-Thiophosphate ester antibody (ab92570; Abcam) showing in vitro SWI3B 968 phosphorylation by KDER. C, 2D Western blot assay with anti SWI3B antibody indicating in 969 vivo phosphorylation alteration of SWI3B protein in er/erl1/erl2 mutant. D, SWI3B and RGA 970 and RGL1 proteins in the nuclei of living cells. Bimolecular Fluorescence Complementation 971 assay (BiFC) in epidermis of tobacco leaves. Scale bar = 10µm. E, The amounts of SWI3B and 972 RGA proteins in plants are oppositely regulated by PAC treatment. F, The disappearance of 973 SWI3B protein is PAC-dose dependent. G, The PAC-dependent degradation of SWI3B is 974 abolished by the MG132 treatment, a known proteasome inhibitor. H, The gal-3 mutant 975 constitutively accumulating DELLA proteins exhibits the decreased level of SWI3B, which is 976 restored to WT levels upon GA treatment. I, The triple DELLA mutant exhibits a WT-like level 977 of SWI3B protein, and the PAC treatment does not influence SWI3B level in this background. J, 978 Schematic model highlighting ERf and DELLA impact on the SWI3B protein.

979

980 Figure 6. ERf proteins enter the nucleus where ERECTA protein binds the GID1 promoters 981 similarly to the SWI3B subunit of SWI/SNF CRC (See also Figures S16 and S17). A, ERECTA 982 protein binds to promoter regions of the GID1a gene in a region targeted by the SWI/SNF 983 complex in three-week-old plants. (error bars refer to SD, P < 0.05, Student's *t*-test, three 984 biological and technical replicates were performed). B, ERECTA and SWI3B core subunit of 985 SWI/SNF CRCs target promoter regions of GID1b gene in three-weeks old plants (error bars 986 refer to SD, P < 0.05, Student's *t*-test, three biological and technical replicates were performed). 987 C, SWI3B binds to the promoter region of the GID1c gene in two different regions. One of them 988 is targeted by ERECTA protein in three-week-old plants. D, ERECTA protein binds to promoter 989 regions of the *GID1a* gene in a region targeted by the SWI/SNF complex in five-week-old plants. 990 (error bars refer to SD, P < 0.05, Student's *t*-test, three biological and technical replicates were 991 performed). E, ERECTA targets promoter region of GID1b gene in five-week-old plants (error

bars refer to SD, P < 0.05, Student's *t*-test, three biological and technical replicates were performed). F, ERECTA binds to the promoter region of the *GID1c* gene in three-week-old plants. The bottom panel in D-F: the binding of native SWI3B protein to its target sites in *GID1ac* promoter regions is abolished in 5-week-old *er/erl1/erl2* triple mutant plants. G, A model describing the non-canonical nuclear function of ERf proteins in the GA signaling pathway.

000	Defenences	
998	Kelerences	
1000	Allen II I i M Prinkwerth CS Deulsen II. Weng D Hühner A. Chey W. H. Devis D.	
1000	Anen JJ, El W, Dinkworth CS, Fauson JE, Wang D, Hubber A, Chou W-H, Davis KJ,	
1001	Net Methods 2007 46 4: 511, 516	
1002	Archaeki P. Buszawicz D. Sarnawski TI. Sarnawska F. Dalieka AT. Tahga T. Farnia AD	
1003	Archacki n, Duszewicz D, Sarhowski 1J, Sarhowska E, Kolicka A1, 10nge I, Fernie AK,	
1004	SWI/SNE Chrometin Demodeling Complex Acts as a Desitive Degulator of Cibbarellin	
1005	Mulieted Decrements in Archidencia DLeS One 8: e59599	
1000	Mediated Responses in Arabidopsis. PLos One 8: e58588	
1007	Archacki R, Yatusevich R, Buszewicz D, Krzyczmonik K, Patryn J, Iwanicka-Nowicka R,	
1008	Biecek P, Wilczynski B, Koblowska M, Jerzmanowski A, et al (2016) Arabidopsis	
1009	SWI/SNF chromatin remodeling complex binds both promoters and terminators to regulate	
1010	gene expression. Nucleic Acids Res 45: 3116–3129	
1011	Belda-Palazón B, Ruiz L, Martí E, Tárraga S, Tiburcio AF, Culiáñez F, Farràs R, Carrasco	
1012	P, Ferrando A (2012) Aminopropyltransferases Involved in Polyamine Biosynthesis	
1013	Localize Preferentially in the Nucleus of Plant Cells. PLoS One. doi:	
1014	10.1371/journal.pone.0046907	
1015	Bitrián M, Roodbarkelari F, Horváth M, Koncz C (2011) BAC-recombineering for studying	
1016	plant gene regulation: Developmental control and cellular localization of SnRK1 kinase	
1017	subunits. Plant J 65 : 829–842	
1018	Bogdan S, Klämbt C (2001) Epidermal growth factor receptor signaling. Curr Biol 11: R292–	
1019	R295	
1020	Boyle EI, Weng S, Gollub J, Jin H, Botstein D, Cherry JM, Sherlock G (2004)	
1021	GO::TermFinder - Open source software for accessing Gene Ontology information and	
1022	finding significantly enriched Gene Ontology terms associated with a list of genes.	
1023	Bioinformatics 20 : 3710–3715	
1024	Cai H, Huang Y, Chen F, Liu L, Chai M, Zhang M, Yan M, Aslam M, He Q, Qin Y (2021)	
1025	ERECTA signaling regulates plant immune responses via chromatin-mediated promotion of	
1026	WRKY33 binding to target genes. New Phytol 230: 737–756	
1027	Cai H, Zhao L, Wang L, Zhang M, Su Z, Cheng Y, Zhao H, Qin Y (2017) ERECTA	
1028	signaling controls Arabidopsis inflorescence architecture through chromatin-mediated	

- activation of PRE1 expression. New Phytol **214**: 1579–1596
- 1030 Cao D, Cheng H, Wu W, Soo HM, Peng J (2006) Gibberellin mobilizes distinct DELLA-
- 1031 dependent transcriptomes to regulate seed germination and floral development in
- 1032 Arabidopsis. PLANT Physiol 142: 509–525
- Casalini P, Iorio M V., Galmozzi E, Ménard S (2004) Role of HER receptors family in
 development and differentiation. J Cell Physiol 200: 343–350
- 1035 Chen M-K, Wilson RL, Palme K, Ditengou FA, Shpak ED (2013) ERECTA Family Genes
- 1036 Regulate Auxin Transport in the Shoot Apical Meristem and Forming Leaf Primordia. Plant
 1037 Physiol 162: 1978–1991
- 1038 Chen MK, Hung MC (2015) Proteolytic cleavage, trafficking, and functions of nuclear receptor
 1039 tyrosine kinases. FEBS J 282: 3693–3721
- 1040 Chen MK, Shpak ED (2014) ERECTA family genes regulate development of cotyledons during
 1041 embryogenesis. FEBS Lett 588: 3912–3917
- 1042 Chenlong Li, Gu L, Gao Lei, Chen C, Wei CQ, Qiu Q, Chien CW, Wang S, Jiang L, Ai LF,
- 1043 Chen CY, Yang S, Nguyen V, Qi Y, Snyder MP, Burlingame AL, Kohalmi SE, Huang
- 1044 S, Cao X, Wang ZY, Wu K, Chen X CY (2016) Concerted genomic targeting of H3K27
- 1045 demethylase REF6 and chromatin-remodeling ATPase BRM in Arabidopsis. Nat Genet 48:1046 687–693
- 1047 la Cour T, Kiemer L, Mølgaard A, Gupta R, Skriver K, Brunak S (2004) Analysis and
 1048 prediction of leucine-rich nuclear export signals. Protein Eng Des Sel 17: 527–536
- 1049 Craft N, Shostak Y, Carey M, Sawyers CL (1999) A mechanism for hormone-independent
 1050 prostate cancer through modulation of androgen receptor signaling by the HER-2/neu
 1051 prostate cancer through modulation of androgen receptor signaling by the HER-2/neu
- 1051 tyrosine kinase. Nat Med 5: 280–5
- Davis AM, Hall A, Millar AJ, Darrah C, Davis SJ (2009) Protocol: Streamlined sub-protocols
 for floral-dip transformation and selection of transformants in Arabidopsis thaliana. Plant
 Methods 5: 3
- 1055 Du J, Jiang H, Sun X, Li Y, Liu Y, Sun M, Fan Z, Cao Q, Feng L, Shang J, et al (2018)
- 1056 Auxin and Gibberellins Are Required for the Receptor-Like Kinase ERECTA Regulated
- 1057 Hypocotyl Elongation in Shade Avoidance in Arabidopsis. Front Plant Sci. doi:
- 1058 10.3389/fpls.2018.00124
- 1059 Earley KW, Haag JR, Pontes O, Opper K, Juehne T, Song K, Pikaard CS (2006) Gateway-

- 1060 compatible vectors for plant functional genomics and proteomics. Plant J. doi:
- 1061 10.1111/j.1365-313X.2005.02617.x
- Gaudino RJ, Pikaard CS (1997) Cytokinin induction of RNA polymerase I transcription in
 Arabidopsis thaliana. J Biol Chem 272: 6799–6804
- 1064 Giri DK, Ali-Seyed M, Li L-Y, Lee D-F, Ling P, Bartholomeusz G, Wang S-C, Hung M-C
- 1065 (2005) Endosomal transport of ErbB-2: mechanism for nuclear entry of the cell surface
 1066 receptor. Mol Cell Biol 25: 11005–11018
- 1067 Griffiths J, Murase K, Rieu I, Zentella R, Zhang Z-L, Powers SJ, Gong F, Phillips AL,
- Hedden P, Sun T, et al (2006) Genetic characterization and functional analysis of the GID1
 gibberellin receptors in Arabidopsis. Plant Cell 18: 3399–3414
- 1070 Haasen D, Köhler C, Neuhaus G, Merkle T (2002) Nuclear export of proteins in plants:
- 1071 AtXPO1 is the export receptor for leucine-rich nuclear export signals in Arabidopsis
- 1072 thaliana. Plant J **20**: 695–705
- Heidstra R, Welch D, Scheres B (2004) Mosaic analyses using marked activation and deletion
 clones dissect Arabidopsis SCARECROW action in asymmetric cell division. Genes Dev
 18: 1964–1969
- Ho CMK, Paciorek T, Abrash E, Bergmann DC (2016) Modulators of Stomatal Lineage
 Signal Transduction Alter Membrane Contact Sites and Reveal Specialization among
 ERECTA Kinases. Dev Cell 38: 345–357
- Hsu JL, Hung MC (2016) The role of HER2, EGFR, and other receptor tyrosine kinases in
 breast cancer. Cancer Metastasis Rev 35: 575–588
- Hu CD, Chinenov Y, Kerppola TK (2002) Visualization of interactions among bZIP and Rel
 family proteins in living cells using bimolecular fluorescence complementation. Mol Cell 9:
 789–798
- 1084 Hung LY, Tseng JT, Lee YC, Xia W, Wang YN, Wu ML, Chuang YH, Lai CH, Chang WC
- 1085 (2008) Nuclear epidermal growth factor receptor (EGFR) interacts with signal transducer
- 1086and activator of transcription 5 (STAT5) in activating Aurora-A gene expression. Nucleic1087Acids Res 36: 4337–4351
- 1088 James Cao H, Lin H-Y, Luidens MK, Davis FB, Davis PJ (2009) Cytoplasm-To-Nucleus
- 1089 Shuttling Of Thyroid Hormone Receptor- β 1 (Tr β 1) Is Directed From A Plasma Membrane
- 1090 Integrin Receptor By Thyroid Hormone. Endocr Res **34**: 31–42

1091	Karachaliou N, Pilotto S, Lazzari C, Bria E, de Marinis F, Rosell R (2016) Cellular and	
1092	molecular biology of small cell lung cancer: an overview. Transl Lung Cancer Res 5: 2–15	
1093	Komata M, Katou Y, Tanaka H, Nakato R, Shirahige K, Bando M (2014) Chromatin	
1094	immunoprecipitation protocol for mammalian cells. Methods Mol Biol 1164: 33–38	
1095	Kosentka PZ, Zhang L, Simon YA, Satpathy B, Maradiaga R, Mitoubsi O, Shpak ED	
1096	(2017) Identification of critical functional residues of receptor-like kinase ERECTA. J Exp	
1097	Bot 68 : 1507–1518	
1098	Kosugi S, Hasebe M, Matsumura N, Takashima H, Miyamoto-Sato E, Tomita M,	
1099	Yanagawa H (2009a) Six classes of nuclear localization signals specific to different binding	
1100	grooves of importin alpha. J Biol Chem 284: 478–485	
1101	Kosugi S, Hasebe M, Tomita M, Yanagawa H (2009b) Systematic identification of cell cycle-	
1102	dependent yeast nucleocytoplasmic shuttling proteins by prediction of composite motifs.	
1103	Proc Natl Acad Sci U S A 106: 10171–10176	
1104	Kubala S, Garnczarska M, Wojtyla Ł, Clippe A, Kosmala A, Zmieńko A, Lutts S, Quinet M	
1105	(2015) Deciphering priming-induced improvement of rapeseed (Brassica napus L.)	
1106	germination through an integrated transcriptomic and proteomic approach. Plant Sci 231:	
1107	94–113	
1108	Lee H-H, Wang Y-N, Hung M-C (2015a) Non-canonical signaling mode of the epidermal	
1109	growth factor receptor family. Am J Cancer Res 5: 2944–2958	
1110	Lee JS, Hnilova M, Maes M, Lin Y-CL, Putarjunan A, Han S-K, Avila J, Torii KU (2015b)	
1111	Competitive binding of antagonistic peptides fine-tunes stomatal patterning. Nature 522:	
1112	439–443	
1113	Lee JS, Kuroha T, Hnilova M, Khatayevich D, Kanaoka MM, Mcabee JM, Sarikaya M,	
1114	Tamerler C, Torii KU (2012) Direct interaction of ligand-receptor pairs specifying	
1115	stomatal patterning. Genes Dev 26: 126–136	
1116	Lo HW, Hsu SC, Ali-Seyed M, Gunduz M, Xia W, Wei Y, Bartholomeusz G, Shih JY, Hung	
1117	MC (2005) Nuclear interaction of EGFR and STAT3 in the activation of the iNOS/NO	
1118	pathway. Cancer Cell 7: 575–589	
1119	Marti U, Burwen SJ, Wells A, Barker ME, Huling S, Feren AM, Jones AL (1991)	
1120	Localization of epidermal growth factor receptor in hepatocyte nuclei. Hepatology 13: 15-	
1121	20	

1122	Mayer K, Albrecht S, Schaller A (2015) Targeted analysis of protein phosphorylation by 2D	
1123	electrophoresis. Methods Mol Biol 1306: 167–176	
1124	Miettinen PJ, Berger JE, Meneses J, Phung Y, Pedersen RA, Werb Z, Derynck R (1995)	
1125	Epithelial immaturity and multiorgan failure in mice lacking epidermal growth factor	
1126	receptor. Nature 376 : 337–341	
1127	Migliaccio A, Di Domenico M, Castoria G, Nanayakkara M, Lombardi M, De Falco A,	
1128	Bilancio A, Varricchio L, Ciociola A, Auricchio F (2005) Steroid receptor regulation of	
1129	epidermal growth factor signaling through Src in breast and prostate cancer cells: Steroid	
1130	antagonist action. Cancer Res 65: 10585–10593	
1131	Miller SG, Carnell L, Moore HPH (1992) Post-Golgi membrane traffic: Brefeldin A inhibits	
1132	export from distal Golgi compartments to the cell surface but not recycling. J Cell Biol 118:	
1133	267–283	
1134	Park CJ, Ronald PC (2012) Cleavage and nuclear localization of the rice XA21 immune	
1135	receptor. Nat Commun 3 : 920	
1136	Pastore JJ, Limpuangthip A, Yamaguchi N, Wu MF, Sang Y, Han SK, Malaspina L,	
1137	Chavdaroff N, Yamaguchi A, Wagner D (2011) LATE MERISTEM IDENTITY2 acts	
1138	together with LEAFY to activate APETALA1. Development 138: 3189–3198	
1139	Phokas A, Coates JC (2021) Evolution of DELLA function and signaling in land plants. Evol	
1140	Dev 23 : 137–154	
1141	Pignon J-C, Koopmansch B, Nolens G, Delacroix L, Waltregny D, Winkler R (2009)	
1142	Androgen receptor controls EGFR and ERBB2 gene expression at different levels in	
1143	prostate cancer cell lines. Cancer Res 69: 2941–2949	
1144	Plackett ARG, Powers SJ, Fernandez-Garcia N, Urbanova T, Takebayashi Y, Seo M,	
1145	Jikumaru Y, Benlloch R, Nilsson O, Ruiz-Rivero O, et al (2012) Analysis of the	
1146	Developmental Roles of the Arabidopsis Gibberellin 20-Oxidases Demonstrates That	
1147	GA20ox1, -2, and -3 Are the Dominant Paralogs. Plant Cell 24: 941–960	
1148	Prigent SA, Gullick WJ (1994) Identification of c-erbB-3 binding sites for phosphatidylinositol	
1149	3'-kinase and SHC using an EGF receptor/c-erbB-3 chimera. EMBO J 13: 2831–2841	
1150	Qi Y, Sun Y, Xu L, Xu Y, Huang H (2004) ERECTA is required for protection against heat-	
1151	stress in the AS1/AS2 pathway to regulate adaxial-abaxial leaf polarity in Arabidopsis.	
1152	Planta 219 : 270–276	

1153	Ragni L, Nieminen K, Pacheco-Villalobos D, Sibout R, Schwechheimer C, Hardtke CS	
1154	(2011) Mobile Gibberellin Directly Stimulates Arabidopsis Hypocotyl Xylem Expansion.	
1155	Plant Cell 23 : 1322–1336	
1156	Rosa NM la, Pfeiffer A, Hill K, Locascio A, Bhalerao RP, Miskolczi P, Grønlund AL,	
1157	Wanchoo-Kohli A, Thomas SG, Bennett MJ, et al (2015) Genome Wide Binding Site	
1158	Analysis Reveals Transcriptional Coactivation of Cytokinin-Responsive Genes by DELLA	
1159	Proteins. PLOS Genet 11: e1005337	
1160	Sacharowski SP, Gratkowska DM, Sarnowska EA, Kondrak P, Jancewicz I, Porri A,	
1161	Bucior E, Rolicka AT, Franzen R, Kowalczyk J, et al (2015) SWP73 subunits of	
1162	arabidopsis SWI/SNF chromatin remodeling complexes play distinct roles in leaf and flower	
1163	development. Plant Cell 27: 1889–1906	
1164	Sáez A, Rodrigues A, Santiago J, Rubio S, Rodriguez PL (2008) HAB1-SWI3B interaction	
1165	reveals a link between abscisic acid signaling and putative SWI/SNF chromatin-remodeling	
1166	complexes in Arabidopsis. Plant Cell 20 : 2972–88	
1167	Saleh A, Alvarez-Venegas R, Avramova Z (2008) An efficient chromatin immunoprecipitation	
1168	(ChIP) protocol for studying histone modifications in Arabidopsis plants. Nat Protoc 3 :	
1169	1018–1025	
1170	Salem MA, Jüppner J, Bajdzienko K, Giavalisco P (2016) Protocol: a fast, comprehensive and	
1171	reproducible one-step extraction method for the rapid preparation of polar and semi-polar	
1172	metabolites, lipids, proteins, starch and cell wall polymers from a single sample. Plant	
1173	Methods 2016 121 12 : 1–15	
1174	Sarnowska E, Gratkowska DM, Sacharowski SP, Cwiek P, Tohge T, Fernie AR, Siedlecki	
1175	JA, Koncz C, Sarnowski TJ (2016) The Role of SWI/SNF Chromatin Remodeling	
1176	Complexes in Hormone Crosstalk. Trends Plant Sci 21: 594–608	
1177	Sarnowska EA, Rolicka AT, Bucior E, Cwiek P, Tohge T, Fernie AR, Jikumaru Y, Kamiya	
1178	Y, Franzen R, Schmelzer E, et al (2013) DELLA-Interacting SWI3C Core Subunit of	
1179	Switch/Sucrose Nonfermenting Chromatin Remodeling Complex Modulates Gibberellin	
1180	Responses and Hormonal Cross Talk in Arabidopsis. Plant Physiol 163: 305–317	
1181	Sarnowski TJ, Swiezewski S, Pawlikowska K, Kaczanowski S, Jerzmanowski A (2002)	
1182	AtSWI3B, an Arabidopsis homolog of SWI3, a core subunit of yeast Swi/Snf chromatin	
1183	remodeling complex, interacts with FCA, a regulator of flowering time. Nucleic Acids Res	

50

- **30**: 3412–3421
- Schmittgen TD, Livak KJ (2008) Analyzing real-time PCR data by the comparative CT method.
 Nat Protoc 3: 1101–1108
- 1187 Shpak ED (2003) Dominant-Negative Receptor Uncovers Redundancy in the Arabidopsis
- ERECTA Leucine-Rich Repeat Receptor-Like Kinase Signaling Pathway That Regulates
 Organ Shape. Plant Cell Online 15: 1095–1110
- Shpak ED (2013) Diverse roles of ERECTA family genes in plant development. J Integr Plant
 Biol 55: 1238–1250
- 1192 Shpak ED, Berthiaume CT, Hill EJ, Torii KU (2004) Synergistic interaction of three
- 1193 ERECTA-family receptor-like kinases controls Arabidopsis organ growth and flower
- development by promoting cell proliferation. Development **131**: 1491–1501
- 1195 Shpak ED, McAbee JM, Pillitteri LJ, Torii KU (2005) Stomatal Patterning and Differentiation
- by Synergistic Interactions of Receptor Kinases. Science (80-) **309**: 290–293
- Simossis VA, Kleinjung J, Heringa J (2005) Homology-extended sequence alignment. Nucleic
 Acids Res 33: 816–824
- 1199 Stewart M, Turley H, Cook N, Pezzella F, Pillai G, Ogilvie D, Cartlidge S, Paterson D,
- 1200 **Copley C, Kendrew J, et al** (2003) The angiogenic receptor KDR is widely distributed in
- 1201 human tissues and tumours and relocates intracellularly on phosphorylation. An
- immunohistochemical study. Histopathology **43**: 33–39
- Tameshige T, Okamoto S, Tasaka M, Torii KU (2016) Impact of erecta mutation on leaf
 serration differs between Arabidopsis accessions. Plant Signal Behav 11 (12): 2478–2485
- Torii KU, Mitsukawa N, Oosumi T, Matsuura Y, Yokoyama R, Whittier RF, Komeda Y
 (1996) The Arabidopsis ERECTA gene encodes a putative receptor protein kinase with
 extracellular leucine-rich repeats. Plant Cell 8: 735–746
- 1207 extracellular leucine-rich repeats. Plant Cell 8: 735–746
- Torii KU, Pillitteri LJ, Sloan DB, Bogenschutz NL (2007) Termination of asymmetric cell
 division and differentiation of stomata. Nature 445: 501–505
- 1210 Uchida N, Lee JS, Horst RJ, Lai H-H, Kajita R, Kakimoto T, Tasaka M, Torii KU (2012a)
- 1211 Regulation of inflorescence architecture by intertissue layer ligand-receptor communication
- between endodermis and phloem. Proc Natl Acad Sci **109**: 6337–6342

1213 Uchida N, Shimada M, Tasaka M (2013) ERECTA-family receptor kinases regulate stem cell

1214 homeostasis via buffering its cytokinin responsiveness in the shoot apical meristem. Plant

215	Cell Physiol 54 : 343–351
-----	----------------------------------

- 1216 Uchida N, Shimada M, Tasaka M (2012b) Modulation of the balance between stem cell
- 1217 proliferation and consumption by ERECTA-family genes. Plant Signal Behav 7: 1506–1508
- 1218 Willige BC, Ghosh S, Nill C, Zourelidou M, Dohmann EMN, Maier A, Schwechheimer C
- 1219 (2007) The DELLA domain of GA INSENSITIVE mediates the interaction with the GA
- 1220 INSENSITIVE DWARF1A gibberellin receptor of Arabidopsis. Plant Cell **19**: 1209–1220
- 1221 Wojtyla Ł, Rucińska-Sobkowiak R, Kubala S, Garnczarska M (2013) Lupine embryo axes
- 1222 under salinity stress. I. Ultrastructural response. Acta Physiol Plant **35**: 2219–2228
- 1223 **Wong RWC** (2003) Transgenic and knock-out mice for deciphering the roles of EGFR ligands.
- 1224 Cell Mol Life Sci **60**: 113–118
- 1225 Woodward AW, Bartel B (2005) A receptor for auxin. Plant Cell 17: 2425–2429
- Yarden Y, Sliwkowski MX (2001) Untangling the ErbB signalling network. Nat Rev Mol Cell
 Biol 2: 127–137
- Van Zanten M, Basten Snoek L, Van Eck-Stouten E, Proveniers MCG, Torii KU, Voesenek
 LACJ, Peeters AJM, Millenaar FF (2010) Ethylene-induced hyponastic growth in
 Arabidopsis thaliana is controlled by ERECTA. Plant J 61: 83–95
- 1231 Zentella R, Zhang Z-L, Park M, Thomas SG, Endo A, Murase K, Fleet CM, Jikumaru Y,
- 1232 Nambara E, Kamiya Y, et al (2007) Global analysis of della direct targets in early
- 1233 gibberellin signaling in Arabidopsis. Plant Cell **19**: 3037–3057
- 1234 Zhang L, DeGennaro D, Lin G, Chai J, Shpak ED (2021) ERECTA family signaling
- 1235 constrains CLAVATA3 and WUSCHEL to the center of the shoot apical meristem.
- 1236 Development. doi: 10.1242/DEV.189753
- 1237