

This is a repository copy of A molecular signal integration network underpinning Arabidopsis thaliana seed germination.

White Rose Research Online URL for this paper: https://eprints.whiterose.ac.uk/162909/

Version: Published Version

Article:

Xu, Hao, Lantzouni, Ourania, Bruggink, Tonko et al. (5 more authors) (2020) A molecular signal integration network underpinning Arabidopsis thaliana seed germination. Current Biology. pp. 3703-3712. ISSN 0960-9822

https://doi.org/10.1016/j.cub.2020.07.012

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.



A Molecular Signal Integration Network Underpinning *Arabidopsis* Seed Germination

Highlights

- The network linking integration of environmental signals to seed growth is mapped
- EXPANSIN gene expression is redundantly regulated and promotes GA-mediated germination
- The TCP14 transcription factor directly regulates EXPANSIN9 expression
- The tcp14/15 germination phenotype is complemented by EXPANSIN9 expression

Authors

Hao Xu, Ourania Lantzouni, Tonko Bruggink, ..., Katherine Denby, Claus Schwechheimer, George W. Bassel

Correspondence

george.bassel@warwick.ac.uk

In Brief

Seeds use a range of environmental cues to decide when to commence germination. Xu et al. report the molecular interaction network used to integrate inputs into growth-promoting *EXPANSIN* gene expression. Functional validation defines *EXPANSIN*s as downstream factors driving germination and TCP14/15 as transcription factors mediating their expression.







Article

A Molecular Signal Integration Network Underpinning *Arabidopsis* Seed Germination

Hao Xu,¹ Ourania Lantzouni,² Tonko Bruggink,⁴ Rene Benjamins,⁴ Frank Lanfermeijer,⁴ Katherine Denby,³ Claus Schwechheimer,² and George W. Bassel^{1,5,*}

¹School of Life Sciences, University of Warwick, Coventry CV4 7AL, UK

²Plant Systems Biology, Technical University of Munich, 85354 Freising, Germany

³Department of Biology, University of York, York YO10 5DD, UK

⁴Applied Physiology, Syngenta Seeds B.V., P.O. Box 2, NL-1600AA Enkhuizen, the Netherlands

⁵Lead Contact

*Correspondence: george.bassel@warwick.ac.uk https://doi.org/10.1016/j.cub.2020.07.012

SUMMARY

Seed dormancy is an adaptive trait defining where and when plants are established. Diverse signals from the environment are used to decide when to initiate seed germination, a process driven by the expansion of cells within the embryo. How these signals are integrated and transduced into the biomechanical changes that drive embryo growth remains poorly understood. Using Arabidopsis seeds, we demonstrate that cell-wallloosening EXPANSIN (EXPA) genes promote gibberellic acid (GA)-mediated germination, identifying EXPAs as downstream molecular targets of this developmental phase transition. Molecular interaction screening identified transcription factors (TFs) that bind to both EXPA promoter fragments and DELLA GA-response regulators. A subset of these TFs is targeted each by nitric oxide (NO) and the phytochrome-interacting TF PIL5. This molecular interaction network therefore directly links the perception of an external environmental signal (light) and internal hormonal signals (GA and NO) with downstream germination-driving EXPA gene expression. Experimental validation of this network established that many of these TFs mediate GA-regulated germination, including TCP14/15, RAP2.2/2.3/2.12, and ZML1. The reduced germination phenotype of the tcp14 tcp15 mutant seed was partially rescued through ectopic expression of their direct target EXPA9. The GA-mediated control of germination by TCP14/15 is regulated through EXPA-mediated control of cell wall loosening, providing a mechanistic explanation for this phenotype and a previously undescribed role for TCPs in the control of cell expansion. This network reveals the paths of signal integration that culminate in seed germination and provides a resource to uncover links between the genetic and biomechanical bases of plant growth.

INTRODUCTION

The development of plants is highly plastic, being capable of change in response to their environment [1]. Rather than making use of individual signals from the environment, complex combinations of inputs are perceived and integrated in order to make developmental decisions. An example of complex signal integration is observed in seed dormancy. This adaptive trait enables plants to move [2] by imposing a growth arrest upon the enclosed embryo [3]. The decision to break dormancy and restart growth is influenced by a variety of signals from the environment including temperature, light quality, smoke, and others [4]. These multiple signals must be perceived, integrated, and turned into a single decision to restart embryo growth. How these diverse environmental inputs are integrated and transduced to stimulate embryo growth remains unknown.

The seed-to-seedling transition in *Arabidopsis* is principally driven by cell expansion [5, 6]. Immediately preceding this induction of cell growth is the expression of diverse gene families that encode proteins that modify the cell wall [7–9]. These include

expansins (*EXPAs*) [10], xyloglucan endo-transglycosylases (XETs) [11], pectin methylesterases (PMEs) [12], and polygalacturonases (PGs) [13]. The functional contribution of each individual class of cell-wall-modifying protein toward the promotion of germination remains unclear.

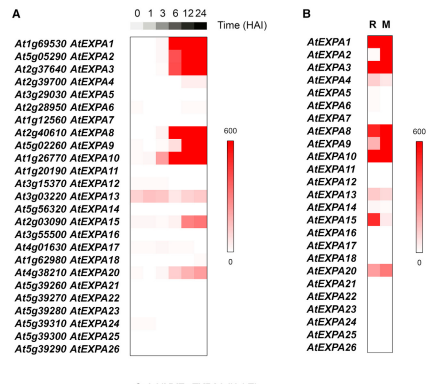
Both germination and cell-expansion-associated gene expression are promoted by the hormone gibberellic acid (GA). The action of this hormone has been demonstrated to occur in multiple sites within seeds, promoting cell expansion in both the embryo [6] and the endosperm [14] in *Arabidopsis*. A necessity for GA in the seed-to-seedling transition is not present because embryos from GA-deficient genotypes grow into stunted seedlings, whereas intact seeds in this species depend on the hormone for germination to occur [15]. Evidence of bidirectional communication between the embryo and endosperm is proposed to coordinate growth across these tissues [16, 17].

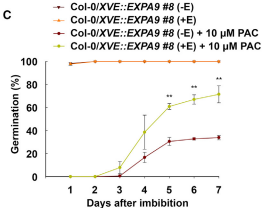
GA responses are repressed by DELLA proteins [8, 18]. DEL-LAs have been proposed to act by physically interacting and inhibiting the activity of transcription factors (TFs) [19]. DELLAs also integrate different signals from the environment, including





Current Biology Article





light and temperature, to control plant growth and development [20]. The mechanistic basis as to how signals from the environment are transduced via GA and DELLAs into the gene expression driving cell expansion and germination remains unknown.

Interaction mapping provides a powerful means to uncover relationships between molecular entities as well as system-level properties, and to identify previously uncharacterized regulators of developmental processes. Systematic mapping of protein-protein interactions in plants has identified regulatory hubs as targets of pathogen effectors [21], established protein-DNA interactions as identified novel regulators of vascular development [22], uncovered the architecture of nitrogen assimilation [23], and described the cistrome-binding landscape of TFs in *Arabidopsis* [24].

This study makes use of interaction mapping to uncover the molecular network used by seeds to integrate multiple signals from the environment and transduce this into growth-promoting gene expression and germination. This establishes direct

Figure 1. EXPA Expression and Function during Seed Germination

(A) Heatmap showing expression of the *EXPA* gene family in germinating seeds at different time points after imbibition (0, 1, 3, 6, 12, and 24 h) [7].

(B) Expression of the *EXPA* gene family in the radicle (R) and micropylar endosperm (M) at 16 h after imbibition [25].

(C) Germination of XVE::EXPA9 seeds on water and in the presence of 10 μM GA-synthesis inhibitor PAC. Ectopic expression of EXPA9 is induced after the application of 30 μM β -estradiol (+E). Data are expressed as mean value \pm standard deviation (SD) (n \geq 50 seeds per biological replicate). See also Figure S1.

molecular links between the perception of environmental signals and the downstream gene expression driving a developmental phase transition in plants.

RESULTS

EXPA Expression Promotes GA-Mediated Seed Germination

A range of gene families associated with cell-wall modification is induced during *Arabidopsis* seed germination [7, 25, 26]. In light of a clearly demonstrated role in promoting plant cell expansion [10] and the implication that they participate in the germination process [14, 26, 27], we investigated the role of *EXPA* genes in GA-mediated embryo growth.

In *Arabidopsis*, the α -*EXPA* gene family consists of 26 members [28]. Publicly available gene expression data indicate 8 of these are induced during seed germination (Figure 1A) [7]. Of these 8 induced genes, only *EXPA2* is specific to the endosperm and not present in the embryo (Figure 1B)

[14, 25, 26]. The subset of 7 genes including *EXPA1*, *EXPA3*, *EXPA8*, *EXPA9*, *EXPA10*, *EXPA15*, and *EXPA20* represents embryo-induced *EXPA* family members.

The induction of *EXPA* genes during seed germination in response to GA has been reported previously [8, 26, 29]. The functional role this *EXPA* gene expression plays in the control of germination, however, remains poorly defined. We examined this by creating an inducible construct consisting of *XVE::EXPA9*, enabling high-level expression in response to estrogen application [30] (Figure S1B).

Seed germination is reduced in the presence of the GA-synthesis inhibitor paclobutrazol (PAC) (Figure 1C) [31]. Ectopic induction of *EXPA9*, however, partially restored this GA-limited germination phenotype (Figures 1C and S1A). This demonstrates *EXPA* expression supports embryo growth under GA-limiting conditions, and presents this gene family as downstream molecular targets promoting the seed-to-seedling transition.

Article



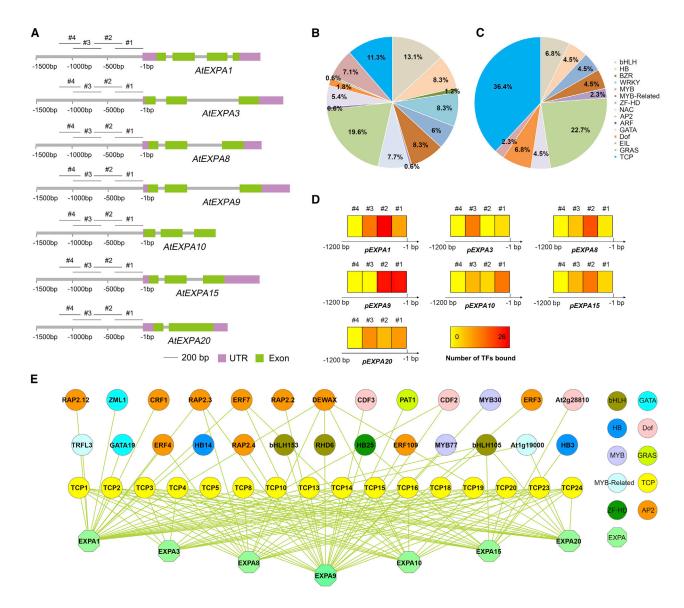


Figure 2. Identification of EXPA-Binding Transcription Factors

(A) Schematics of genomic regions of embryo-induced EXPA genes. 3' UTR regions are colored in purple, and exons are colored in green. The numbers below the gray lines indicate relative positions upstream of the transcription start site (TSS). Promoter bait fragments used in the Y1h assay are labeled with the numbers 1 to 4, consisting of ~300- to ~400-bp sequences with 100-bp overlaps between adjacent fragments.

- (B) Pie chart showing the categorization of transcription factor (TF) families included in the Y1h library screened (Data S1B).
- (C) Pie chart showing the percentage of TFs by family that bound to EXPA promoters in yeast (Data S1C).
- (D) Heatmaps showing the number of TFs that bound to the overlapping promoter fragments of EXPA genes.
- (E) Interaction between TFs and EXPA promoter fragments in Y1h assays. Node color indicates TF family after the legend on the right. See also Figures S2–S4.

Identification of TFs that Bind Germination *EXPA* Promoter Sequences

With the (1) identification of *EXPA* gene expression as a promoter of germination under GA-limiting conditions, and (2) identification of 7 *EXPA* genes being induced in *Arabidopsis* embryos during seed germination, we sought to identify the molecular factors that regulate the expression of these germination-promoting genes.

To identify TFs that bind the 7 EXPA promoter sequences, a yeast-1-hybrid (Y1h) assay was used. To generate the EXPA

promoter bait constructs, 1.2-kb sequence fragments upstream of the transcription start site were cloned and divided into four bait fragments of 300~400 bp with 100-bp-overlapping regions (Figure 2A).

To obtain a suitable TF library for screening, TFs expressed at 7 h after imbibition (HAI) and 12 HAI in the germinating embryo axis, according to publicly available expression data, were selected for subsequent analyses [25] (Data S1A). The selection of the respective time window was guided by the temporal induction of *EXPA* gene expression, reaching high levels by 12



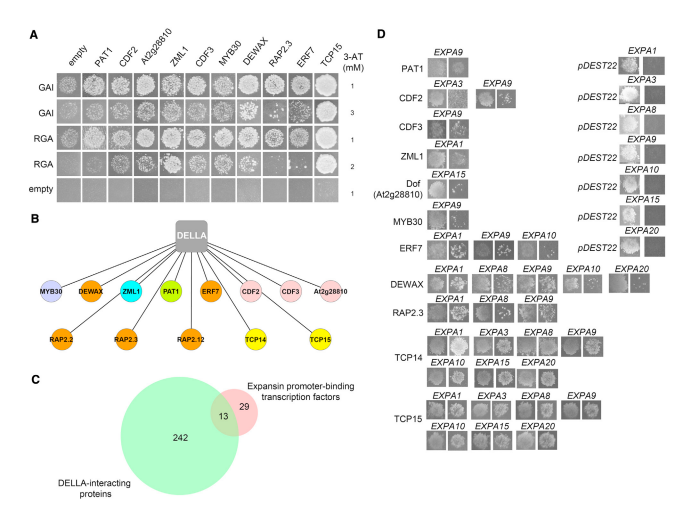


Figure 3. Interaction between EXPA-Binding Transcription Factors and DELLA

Molecular interactions between TFs and DELLA, and TFs with EXPA promoter sequences.

- (A) Interaction between TFs and DELLA proteins in Y2h assays.
- (B) Network illustrating the interaction between DELLA and EXPA promoter-binding TFs.
- (C) Venn diagram showing the overlap between DELLA-interacting proteins and TFs that bind the promoter sequences of EXPA genes.
- (D) Y1h colonies showing interaction between TFs that bind to *EXPA* promoter fragments and also interact with DELLA. Left: yeast growth on standard deviation (SD)-Leu-Trp media. Right: yeast growth on SD-Leu-Trp-His + 3-amino-1,2,4-triazole (3-AT). See also Data S2.

HAI (Figure 1A). A total of 168 of the 255 (66%) seed-expressed TFs were covered in a prey library (Data S1B) [32], representing transcripts from 15 TF families (Figure 2B).

Screening of the 28 genomic bait fragments representing the *EXPA* promoter sequences against 168 prey TFs in the Y1h assay resulted in a total of 134 TF-promoter interactions between 42 TFs and genomic regions of 7 *EXPA*s (Figures 2C–2E, 3D, and S2–S4).

Extensive redundancy in the binding of germination-associated *EXPA* promoters was identified as a result of this Y1h screening (Figure 2E; Data S2). A total of 23 of the 42 TFs bound to multiple promoter fragments (55%), whereas each *EXPA* promoter had multiple TFs binding their promoters. Biases in the TF families binding to *EXPA* promoter sequences were also observed, and the TCP and AP2 TF families were enriched (Figure 2C).

EXPA Promoter-Binding TFs Also Interact with DELLA Proteins

Previous work investigating gene expression in seeds has demonstrated the control of GA-promoted germination, and *EXPA* gene expression is regulated by DELLA proteins [18, 33]. To exert their function, DELLA proteins physically interact with TFs such that they sequester their capacity to bind target DNA sequences [33, 34]. We sought to establish whether the TFs identified as *EXPA* promoter-binding proteins also interact with DELLA proteins.

Yeast-2-hybrid (Y2h) screening using truncated (M5) versions of the DELLA proteins REPRESSOR-OF-*ga1-3* (RGA) and GIB-BERELLIC ACID INSENSITIVE (GAI) [19, 35] identified multiple TF-DELLA interactions including that of CYCLING DOF FACTOR 2 (CDF2) and CDF3, DECREASE WAX BIOSYNTHESIS (DEWAX), PHYTOCHROME A SIGNAL TRANSDUCTION 1

Article



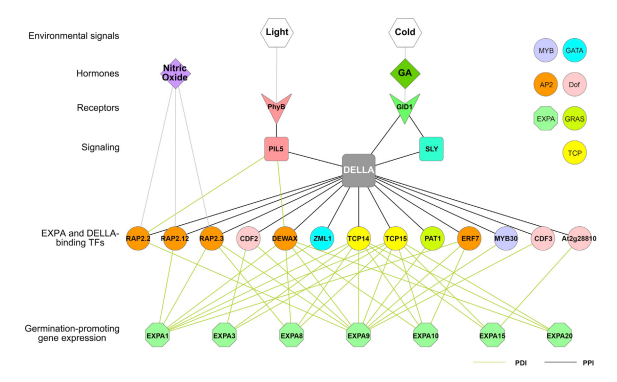


Figure 4. Signal Integration and Growth Regulatory Network in Arabidopsis Seeds

Molecular regulatory network linking the perception of environmental and hormonal cues to downstream gene expression driving cellular biomechanical changes and embryo growth. TFs are shown as circles, *EXPA* nodes as green octagons, receptors as arrowheads, hormones as squares, and environmental inputs as hexagons. Green edges indicate protein-DNA interactions, black edges indicate protein-protein interactions, and gray edges indicate the regulatory relationships between nodes. Node color indicates TF family after the legend on the right.

(PAT1), Dof-type zinc finger (Dof, At2g28810), ZIM-LIKE 1 (ZML1/GATA24), MYB DOMAIN PROTEIN 30 (MYB30), ETHYLENE RESPONSE FACTOR 7 (ERF7), RELATED TO AP2 3 (RAP2.3), and TEOSINTE BRANCHED1/CYCLOIDEA/PCF 15 (TCP15) (Figures 3A and 3B). Several other *EXPA* promoterbinding TFs had been previously reported to interact with DELLA proteins including TCP14 [36] and RAP2.2/RAP2.12 [37]. Of the 42 TFs identified that bind *EXPA* promoters, 13 also physically interact with DELLA proteins (Figure 3C).

A Molecular Interaction Network Underpinning Seed Germination

Results of the Y1h screen identified TFs that bind to germination-associated *EXPA* genes. These TFs were compared with the Y2h screen identifying DELLA-interacting TFs. This resulted in a network consisting of 134 protein-DNA interactions involving 42 TFs, 7 *EXPA*s, and 13 DELLA-TF protein-protein interactions (Data S2). The subset of this molecular interaction network providing direct links between the perception of GA and the binding of *EXPA* promoter fragments is presented in Figure 4.

The ERFVII family of TFs including RAP2.2, RAP2.3, and RAP2.12 acts as sensors of nitric oxide (NO) in plants [38]. The Y1h screen identifies them as also binding the promoters of *EXPA* genes, providing a potential link between this signaling molecule and the biomechanical modulation of plant growth (Figure 4).

The TF PHYTOCHROME INTERACTING FACTOR3 LIKE5 (PIL5) mediates light-regulated control of seed germination, and interacts with the red light receptor PHYTOCHROME [39].

PIL5 also physically interacts with DELLA proteins [40] and binds to their promoter sequences [41]. Despite the central role of this TF in the control of germination, *EXPA* genes upregulated during germination are not direct targets of PIL5 [42]. The TFs ERF4, CRF1, DEWAX, and RAP2.2 are, however, direct transcriptional targets of PIL5 and bind *EXPA* promoter sequences in the Y1h assay. Both DEWAX and RAP2.2 also interact with DELLA, providing a link between the perception of light and the binding of *EXPA* promoter sequences.

The organization of this molecular interaction network reveals a hierarchical structure of signal integration underpinning the control of seed germination (Figure 4) [4]. The interactions between these molecular agents identify the paths of environmental and hormonal signal perception, their integration, and ultimately the promotion of embryo cell growth through the induction of *EXPA* gene expression.

EXPA Promoter-Binding Proteins Impact GA-Mediated Germination

Whether the TFs identified through molecular interaction screening play a role in GA-mediated *Arabidopsis* seed germination through their control of *EXPA* gene expression was investigated. A total of 13 TFs was screened that both interact with DELLA proteins and bind to the *EXPA* promoter sequences (Figure S5).

Mutant seeds were initially screened on the GA-synthesis inhibitor PAC, identifying potential GA-mediated germination phenotypes for *TCP14/15*, *ZML1*, and *RAP2.2/2.3/2.12* on 10 µM



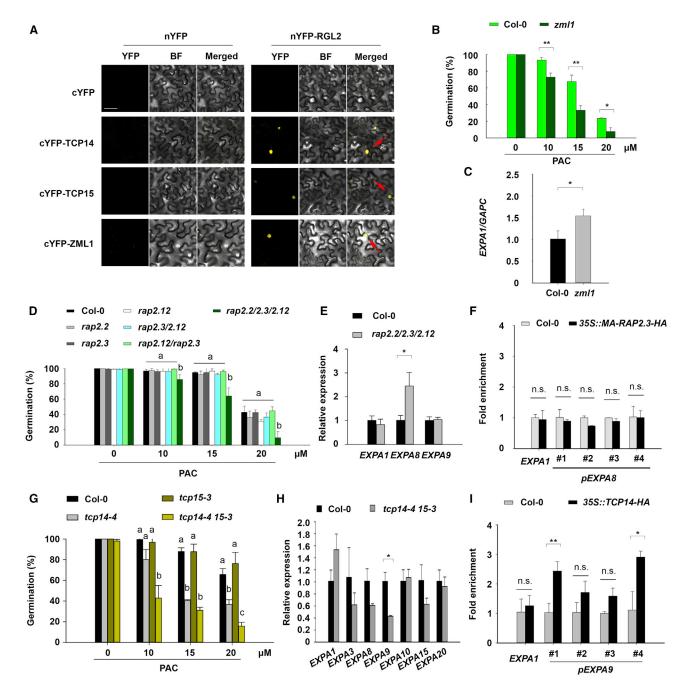


Figure 5. Functional Validation of the Signal Integration Network

(A) BiFC assay investigating the interaction between DELLA and TFs in plant cells. Fusion proteins were co-expressed in tobacco leaves by using *Agrobacterium* infiltration and images represent co-transfected cells with visible fluorescence. YFP, fluorescence of yellow fluorescent protein; BF, bright field; merged, merger of the YFP and bright-field images. Red arrows indicate the position of YFP signal. The scale bar indicates 50 μm.

- (B) Germination of *zml1* imbibed in different concentrations of PAC.
- (C) Relative expression of the ZML1 target EXPA1 in the zml1 background determined by using qPCR.
- (D) Same as (B) using combinations of rap2.2 rap2.3 and rap2.12 mutant seeds.
- (E) Relative expression of the ERFVII TF targets EXPA1, EXPA8, and EXPA9 in the rap2.2 rap2.3 rap2.12 mutant background determined by qPCR.
- (F) ChIP using HA-tagged RAP2.3 on the EXPA8 promoter. The EXPA1-coding region was used as a negative control and no specific signals were observed in the negative-control region.
- (G) Same as (B) using tcp14 and tcp15 mutant seeds.
- (H) qPCR analysis of $\it EXPA$ genes targeted by TCP14 and TCP15.

(legend continued on next page)

Article



PAC (Figures S5E–S5M). Before proceeding with further functional analysis of these genes, the interactions between these TFs and DELLA were investigated *in planta*. Although each TCP14/15 and ERFVII TF has been previously reported to interact with DELLAs [36, 37, 40], this has not been established for ZML1. Using bimolecular fluorescence complementation (BiFC) in tobacco leaves, we confirmed the interaction between ZML1 and RGL2 within plant cells (Figure 5A).

Phenotyping mutant seeds on a broader range of PAC concentrations confirmed GA-mediated germination phenotypes for *zml1* (Figure 5B). The regulatory relationship between ZML1 and its putative target *EXPA1* was examined by looking at *EXPA* transcript abundance in the *zml1*-null mutant background. This showed *EXPA1* transcript abundance to be significantly higher than in the wild type (Figure 5C). The inability to recover a transgenic line harboring a detectable epitope-tagged version of this protein prevented the further study of this interaction.

Compared with the wild type, the *rap2.2/2.3/2.12* triplemutant background showed reduced germination in the presence of PAC, suggesting these ERFVII proteins promote embryo growth in response to GA (Figure 5D). Of the *EXPA* targets of these TFs, the transcript abundance of *EXPA8* was significantly higher in the triple mutant than in the wild type (Figure 5E). Chromatin immunoprecipitation (ChIP) analysis using *35S::MA-RAP2.3-HA* (hemagglutinin), however, did not identify the enrichment of this protein on its putative *EXPA8* promoter fragment targets (Figure 5F). The expression level of *EXPA8* might therefore be indirectly regulated by these *ERFVII* genes.

The *tcp14-4* single and *tcp14-4* 15-3 double mutants showed reduced germination in the presence of PAC (Figure 5G). A role for *TCP14* and *TCP15* in GA-stimulated seed germination has been reported previously [43], and was proposed to occur through the regulation of the cell cycle by TCPs within the cells of the radicle. Although the overall contribution of cell division in the radicle to overall embryo growth and germination remains unclear [5, 6], the putative control of *EXPA*s by these TCP TFs was examined as a putative explanation for this reduced germination phenotype.

In the *tcp14-4 15-3* double mutant, the transcript abundance of *EXPA9* was significantly decreased, whereas the expression of the other 6 putative *EXPA* targets was not altered (Figure 5H). ChIP using a *35S::TCP14-HA* construct identified an enrichment of this protein on the promoter of *EXPA9* (Figure 5I). These results collectively suggest TCP14 to be a direct positive regulator of *EXPA9* expression.

By taking together (1) *EXPA* expression is capable of stimulating GA-mediated seed germination (Figure 1C), (2) the *tcp14* mutant shows a reduced capacity to germinate under GA-limited conditions (Figure 5G), and (3) *EXPA9* gene expression in the absence of *TCP14* and *TCP15* (Figure 5H), we sought to determine whether the expression of *EXPA9* is sufficient to account for the reduced germination phenotype observed in the *tcp14-4 tcp15-3* mutant.

To test this, the estrogen-inducible XVE::EXPA9 construct was introduced into the tcp14-4 tcp15-3 loss-of-function mutant.

Although no significant difference in germination was observed under standard germination conditions, induced expression of *EXPA9* in *tcp14-4 tcp15-3* led to an enhanced germination response in the presence of PAC (Figures 6 and S6). The ectopic expression of *EXPA9* is therefore capable of partially rescuing GA-mediated germination defects in *tcp14-4 tcp15-3*, while partly explaining the mechanistic basis for this phenotype.

A Resource for Plant Growth and Signal Integration

This study identified 42 TFs capable of binding *EXPA* promoter sequences, 13 of which also interact with DELLA proteins. These TFs represent putative regulators of *EXPA* gene expression, and therefore cell wall biomechanics [10]. The identification of these TFs as direct modulators of plant cell growth provides a focused resource awaiting further characterization. This resource might be used to identify further mechanistic interactions underpinning the modulation of plant growth by linking genetic pathways to the biomechanical changes driving cell expansion. In addition to the provided network file (Data S2), these interactions from this publication have been submitted to the IMEx Consortium (http://www.imexconsortium.org) through IntAct [44] and assigned the identifier IM-27910.

DISCUSSION

The regulation of plant development is highly plastic in response to the environment [1]. In this study, we reveal the genetic complexity that underpins signal integration in the control of embryo growth during seed germination. These insights reveal the extent and nature of the redundancy that underlies the control of the downstream gene expression promoting GA-mediated embryo growth. This is observed in terms of the number of *EXPA* genes that are induced during germination, and the large number of TFs that bind to the promoters of these genes (Figure 2E). This network further reveals the paths of information flow from the perception of environmental signals to the downstream gene expression that drives the seed-to-seedling developmental transition (Figure 4).

Although a variety of environmental, hormonal, and genetic factors have been described to regulate seed germination [3], how these signals are integrated and transduced into embryo growth remains unclear. This study addressed this gap by performing targeted molecular interaction mapping to establish a network underpinning signal integration and seed-germination-driving gene expression in *Arabidopsis* (Figure 4). The network linked the perception of environmental (light quality) and hormonal (GA and NO) signals to the gene expression responsible for the biomechanical changes driving the seed-to-seedling transition (*EXPA*).

Although *EXPA* gene expression was shown to promote GAmediated embryo growth (Figure 1C), the regulation of these downstream targets is highly redundant, and multiple TFs bound to their associated upstream promoter fragments (Figure 2E). This redundancy in the integration of signals into *EXPA* gene



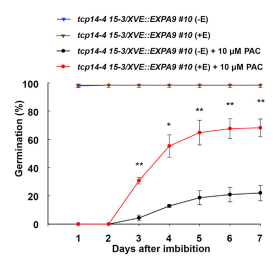


Figure 6. Functional Complementation of TCP Germination by EXPA Expression

Impact of ectopic expression of *EXPA9* on seed germination upon 10 μ M PAC treatment in the *tcp14-4 tcp15-3* mutant background. Data were statistically analyzed by using Student's *t* test (*p < 0.05, **p < 0.01). See also Figure S6.

expression likely provides robustness to the germination process, whereby a single TF-EXPA interaction rarely impacts the seed-to-seedling transition (Figure S5). This redundancy could explain the relatively small number of GA-related germination phenotypes identified in seeds carrying mutations in single genes or restricted to single gene families. Despite this redundancy, phenotypes in null mutants of the ERFVII (RAP/2.2/2.3/2.12), TCP14/15, and ZML1 TFs were identified (Figures 5B, 5D, and 5G).

The *tcp14 tcp15* mutant showed reduced germination in GA-limited conditions (Figure 5G), and TCP14 directly bound and promoted *EXPA9* expression (Figures 5H and 5l). A GA-mediated germination phenotype for TCP14 and TCP15 has been proposed previously to act through the control of the cell cycle in the radicle [43]. This work extends the role of these genes to the promotion of cell expansion through the control of *EXPA9* gene expression, a proposal consistent with the description of this TF as a promoter of embryo growth potential [45]. This finding further suggests the mechanistic basis of reduced germination in the *tcp14 tcp15* mutant background to be at least partially due to a reduction in *EXPA* expression, as supported by the partial rescue of this phenotype under GA-limiting conditions (Figure 6).

The uncovering of this network provides insight into a fundamental gap in our understanding of a complex biological process: how multiple environmental inputs are integrated to create a single developmental output, in this instance, how environmental signals are used to regulate the gene expression altering cellular biomechanics and embryo growth in seeds. It further highlights the importance of understanding the complexity of transcriptional regulation as a whole system, while providing a resource for further exploitation by understanding how the activity of the TFs that bind to *EXPAs* modulates plant growth.

The identification of downstream targets of developmental processes enables the mechanistic basis of phenotypes to be established and the comprehensive mapping of the molecular interactions underpinning developmental phase transitions. Similar approaches and the resources generated by these studies represent powerful ways to understand plant development.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
 - Lead Contact
 - Materials Availability
 - Data and Code Availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
 - Plant Material and growth conditions
- METHOD DETAILS
 - Plasmid construction and plant transformation
 - Phenotypic analyses
 - Gene expression analyses
 - Yeast one hybrid screening of Arabidopsis cDNA libraries
 - Protein-protein interaction assays
 - O BiFC protein-protein interaction in planta
 - Chromatin Immunoprecipitation
 - Bioinformatics analyses
- QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.cub.2020.07.012.

ACKNOWLEDGMENTS

We thank George Coupland (Max Planck Institute for Plant Breeding Research), Brendan Davies (University of Leeds), Yuda Fang (Shanghai Institutes for Biological Sciences), Michael Holdsworth (University of Nottingham), Åsa Strand (Umea University), and Mi Chung Suh (Chonnam National University) for sharing research materials. This work was supported by the Biotechnology and Biological Sciences Research Council and Syngenta B.V. (BB/N009754/1) to G.W.B., BBSRC/EPSRC-funded Systems Approaches to Biological Research (grant BB/F005806/1) to K.D., and Sonderforschungsbereich (924) to C.S. We also thank Abigail Cooper and Guillaume Chomicki for technical support.

AUTHOR CONTRIBUTIONS

H.X. performed all phenotypic and molecular analyses; K.D. supervised the protein-DNA interaction screening; O.L. and C.S. provided the protein-protein interaction screening; G.W.B. conceived and supervised the project; G.W.B., T.B., R.B., and F.L. designed the experiments; and H.X. and G.W.B. wrote the manuscript with input from all co-authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: January 8, 2020 Revised: June 22, 2020 Accepted: July 2, 2020 Published: August 6, 2020

Article



REFERENCES

- Domagalska, M.A., and Leyser, O. (2011). Signal integration in the control of shoot branching. Nat. Rev. Mol. Cell Biol. 12, 211–221.
- Koornneef, M., Bentsink, L., and Hilhorst, H. (2002). Seed dormancy and germination. Curr. Opin. Plant Biol. 5, 33–36.
- Holdsworth, M.J., Bentsink, L., and Soppe, W.J. (2008). Molecular networks regulating *Arabidopsis* seed maturation, after-ripening, dormancy and germination. New Phytol. 179, 33–54.
- Bassel, G.W. (2016). To grow or not to grow? Trends Plant Sci. 21, 498–505.
- Sliwinska, E., Bassel, G.W., and Bewley, J.D. (2009). Germination of Arabidopsis thaliana seeds is not completed as a result of elongation of the radicle but of the adjacent transition zone and lower hypocotyl. J. Exp. Bot. 60, 3587–3594.
- Bassel, G.W., Stamm, P., Mosca, G., Barbier de Reuille, P., Gibbs, D.J., Winter, R., Janka, A., Holdsworth, M.J., and Smith, R.S. (2014). Mechanical constraints imposed by 3D cellular geometry and arrangement modulate growth patterns in the *Arabidopsis* embryo. Proc. Natl. Acad. Sci. USA 111, 8685–8690.
- Nakabayashi, K., Okamoto, M., Koshiba, T., Kamiya, Y., and Nambara, E. (2005). Genome-wide profiling of stored mRNA in *Arabidopsis thaliana* seed germination: epigenetic and genetic regulation of transcription in seed. Plant J. 41, 697–709.
- Ogawa, M., Hanada, A., Yamauchi, Y., Kuwahara, A., Kamiya, Y., and Yamaguchi, S. (2003). Gibberellin biosynthesis and response during Arabidopsis seed germination. Plant Cell 15, 1591–1604.
- Narsai, R., Gouil, Q., Secco, D., Srivastava, A., Karpievitch, Y.V., Liew, L.C., Lister, R., Lewsey, M.G., and Whelan, J. (2017). Extensive transcriptomic and epigenomic remodelling occurs during *Arabidopsis thaliana* germination. Genome Biol. 18, 172.
- Cosgrove, D.J. (2000). Loosening of plant cell walls by expansins. Nature 407, 321–326.
- Chen, F., Nonogaki, H., and Bradford, K.J. (2002). A gibberellin-regulated xyloglucan endotransglycosylase gene is expressed in the endosperm cap during tomato seed germination. J. Exp. Bot. 53, 215–223.
- Ren, C., and Kermode, A.R. (2000). An increase in pectin methyl esterase activity accompanies dormancy breakage and germination of yellow cedar seeds. Plant Physiol. 124, 231–242.
- Sitrit, Y., Hadfield, K.A., Bennett, A.B., Bradford, K.J., and Downie, A.B. (1999). Expression of a polygalacturonase associated with tomato seed germination. Plant Physiol. 121, 419–428.
- Sánchez-Montesino, R., Bouza-Morcillo, L., Marquez, J., Ghita, M., Duran-Nebreda, S., Gómez, L., Holdsworth, M.J., Bassel, G., and Oñate-Sánchez, L. (2019). A regulatory module controlling GA-mediated endosperm cell expansion is critical for seed germination in *Arabidopsis*. Mol. Plant 12, 71–85.
- Debeaujon, I., and Koornneef, M. (2000). Gibberellin requirement for Arabidopsis seed germination is determined both by testa characteristics and embryonic abscisic acid. Plant Physiol. 122, 415–424.
- Bassel, G.W., Mullen, R.T., and Bewley, J.D. (2008). Procera is a putative DELLA mutant in tomato (Solanum lycopersicum): effects on the seed and vegetative plant. J. Exp. Bot. 59, 585–593.
- 17. Lee, K.P., Piskurewicz, U., Turecková, V., Strnad, M., and Lopez-Molina, L. (2010). A seed coat bedding assay shows that RGL2-dependent release of abscisic acid by the endosperm controls embryo growth in *Arabidopsis* dormant seeds. Proc. Natl. Acad. Sci. USA 107, 19108–19113.
- Lee, S., Cheng, H., King, K.E., Wang, W., He, Y., Hussain, A., Lo, J., Harberd, N.P., and Peng, J. (2002). Gibberellin regulates *Arabidopsis* seed germination via RGL2, a GAI/RGA-like gene whose expression is up-regulated following imbibition. Genes Dev. 16, 646–658.
- de Lucas, M., Davière, J.-M., Rodríguez-Falcón, M., Pontin, M., Iglesias-Pedraz, J.M., Lorrain, S., Fankhauser, C., Blázquez, M.A., Titarenko, E.,

- and Prat, S. (2008). A molecular framework for light and gibberellin control of cell elongation. Nature *451*, 480–484.
- Achard, P., Cheng, H., De Grauwe, L., Decat, J., Schoutteten, H., Moritz, T., Van Der Straeten, D., Peng, J., and Harberd, N.P. (2006). Integration of plant responses to environmentally activated phytohormonal signals. Science 311, 91–94.
- Mukhtar, M.S., Carvunis, A.-R., Dreze, M., Epple, P., Steinbrenner, J., Moore, J., Tasan, M., Galli, M., Hao, T., Nishimura, M.T., et al. (2011). Independently evolved virulence effectors converge onto hubs in a plant immune system network. Science 333, 596–601.
- Taylor-Teeples, M., Lin, L., de Lucas, M., Turco, G., Toal, T.W., Gaudinier, A., Young, N.F., Trabucco, G.M., Veling, M.T., Lamothe, R., et al. (2015). An *Arabidopsis* gene regulatory network for secondary cell wall synthesis. Nature 517, 571–575.
- Gaudinier, A., Rodriguez-Medina, J., Zhang, L., Olson, A., Liseron-Monfils, C., Bågman, A.-M., Foret, J., Abbitt, S., Tang, M., Li, B., et al. (2018). Transcriptional regulation of nitrogen-associated metabolism and growth. Nature 563, 259–264.
- O'Malley, R.C., Huang, S.-C., Song, L., Lewsey, M.G., Bartlett, A., Nery, J.R., Galli, M., Gallavotti, A., and Ecker, J.R. (2016). Cistrome and epicistrome features shape the regulatory DNA landscape. Cell 165, 1280–1292.
- Dekkers, B.J.W., Pearce, S., van Bolderen-Veldkamp, R.P., Marshall, A., Widera, P., Gilbert, J., Drost, H.-G., Bassel, G.W., Müller, K., King, J.R., et al. (2013). Transcriptional dynamics of two seed compartments with opposing roles in *Arabidopsis* seed germination. Plant Physiol. *163*, 205–215.
- 26. Linkies, A., Müller, K., Morris, K., Turecková, V., Wenk, M., Cadman, C.S., Corbineau, F., Strnad, M., Lynn, J.R., Finch-Savage, W.E., and Leubner-Metzger, G. (2009). Ethylene interacts with abscisic acid to regulate endosperm rupture during germination: a comparative approach using Lepidium sativum and Arabidopsis thaliana. Plant Cell 21, 3803–3822.
- Stamm, P., Topham, A.T., Mukhtar, N.K., Jackson, M.D., Tomé, D.F., Beynon, J.L., and Bassel, G.W. (2017). The transcription factor ATHB5 affects GA-mediated plasticity in hypocotyl cell growth during seed germination. Plant Physiol. 173, 907–917.
- Kende, H., Bradford, K., Brummell, D., Cho, H.-T., Cosgrove, D., Fleming, A., Gehring, C., Lee, Y., McQueen-Mason, S., Rose, J., and Voesenek, L.A. (2004). Nomenclature for members of the expansin superfamily of genes and proteins. Plant Mol. Biol. 55, 311–314.
- Graeber, K., Linkies, A., Steinbrecher, T., Mummenhoff, K., Tarkowská, D., Turečková, V., Ignatz, M., Sperber, K., Voegele, A., de Jong, H., et al. (2014). DELAY OF GERMINATION 1 mediates a conserved coatdormancy mechanism for the temperature- and gibberellin-dependent control of seed germination. Proc. Natl. Acad. Sci. USA 111, E3571– E3580.
- Papdi, C., Abrahám, E., Joseph, M.P., Popescu, C., Koncz, C., and Szabados, L. (2008). Functional identification of *Arabidopsis* stress regulatory genes using the controlled cDNA overexpression system. Plant Physiol. 147, 528–542.
- Piskurewicz, U., Jikumaru, Y., Kinoshita, N., Nambara, E., Kamiya, Y., and Lopez-Molina, L. (2008). The gibberellic acid signaling repressor RGL2 inhibits Arabidopsis seed germination by stimulating abscisic acid synthesis and ABI5 activity. Plant Cell 20, 2729–2745.
- 32. Castrillo, G., Turck, F., Leveugle, M., Lecharny, A., Carbonero, P., Coupland, G., Paz-Ares, J., and Oñate-Sánchez, L. (2011). Speeding cis-trans regulation discovery by phylogenomic analyses coupled with screenings of an arrayed library of Arabidopsis transcription factors. PLoS One 6, e21524.
- Cao, D., Cheng, H., Wu, W., Soo, H.M., and Peng, J. (2006). Gibberellin mobilizes distinct DELLA-dependent transcriptomes to regulate seed germination and floral development in *Arabidopsis*. Plant Physiol. 142, 509–525.
- Davière, J.-M., and Achard, P. (2013). Gibberellin signaling in plants. Development 140, 1147–1151.



- 35. Lantzouni, O., Alkofer, A., Falter-Braun, P., and Schwechheimer, C. (2020). GROWTH-REGULATING FACTORS interact with DELLAs and regulate growth in cold stress. Plant Cell 32, 1018-1034.
- 36. Davière, J.-M., Wild, M., Regnault, T., Baumberger, N., Eisler, H., Genschik, P., and Achard, P. (2014). Class I TCP-DELLA interactions in inflorescence shoot apex determine plant height. Curr. Biol. 24, 1923-
- 37. Marín-de la Rosa, N., Sotillo, B., Miskolczi, P., Gibbs, D.J., Vicente, J., Carbonero, P., Oñate-Sánchez, L., Holdsworth, M.J., Bhalerao, R., Alabadí, D., and Blázquez, M.A. (2014). Large-scale identification of gibberellin-related transcription factors defines group VII ETHYLENE RESPONSE FACTORS as functional DELLA partners. Plant Physiol. 166,
- 38. Gibbs. D.J., Md Isa, N., Movahedi, M., Lozano-Juste, J., Mendiondo, G.M., Berckhan, S., Marín-de la Rosa, N., Vicente Conde, J., Sousa Correia, C., Pearce, S.P., et al. (2014). Nitric oxide sensing in plants is mediated by proteolytic control of group VII ERF transcription factors. Mol. Cell 53, 369-379.
- 39. Oh, E., Kim, J., Park, E., Kim, J.-I., Kang, C., and Choi, G. (2004). PIL5, a phytochrome-interacting basic helix-loop-helix protein, is a key negative regulator of seed germination in Arabidopsis thaliana. Plant Cell 16, 3045-3058
- 40. Gallego-Bartolomé, J., Minguet, E.G., Marín, J.A., Prat, S., Blázquez, M.A., and Alabadí, D. (2010). Transcriptional diversification and functional conservation between DELLA proteins in Arabidopsis. Mol. Biol. Evol. 27, 1247-1256
- 41. Oh, E., Yamaguchi, S., Hu, J., Yusuke, J., Jung, B., Paik, I., Lee, H.-S., Sun, T.P., Kamiya, Y., and Choi, G. (2007). PIL5, a phytochrome-interacting bHLH protein, regulates gibberellin responsiveness by binding directly to the GAI and RGA promoters in Arabidopsis seeds. Plant Cell 19, 1192-1208.
- 42. Oh, E., Kang, H., Yamaguchi, S., Park, J., Lee, D., Kamiya, Y., and Choi, G. (2009). Genome-wide analysis of genes targeted by PHYTOCHROME INTERACTING FACTOR 3-LIKE5 during seed germination in Arabidopsis. Plant Cell 21, 403-419.
- 43. Resentini, F., Felipo-Benavent, A., Colombo, L., Blázquez, M.A., Alabadí, D., and Masiero, S. (2015). TCP14 and TCP15 mediate the promotion of seed germination by gibberellins in Arabidopsis thaliana. Mol. Plant 8, 482-485
- 44. Orchard, S., Ammari, M., Aranda, B., Breuza, L., Briganti, L., Broackes-Carter, F., Campbell, N.H., Chavali, G., Chen, C., del-Toro, N., et al. (2014). The MIntAct project-IntAct as a common curation platform for 11 molecular interaction databases. Nucleic Acids Res. 42, D358-D363.
- 45. Tatematsu, K., Nakabayashi, K., Kamiya, Y., and Nambara, E. (2008). Transcription factor AtTCP14 regulates embryonic growth potential during seed germination in Arabidopsis thaliana. Plant J. 53, 42-52.
- 46. Fornara, F., Panigrahi, K.C.S., Gissot, L., Sauerbrunn, N., Rühl, M., Jarillo, J.A., and Coupland, G. (2009). Arabidopsis DOF transcription factors act redundantly to reduce CONSTANS expression and are essential for a photoperiodic flowering response. Dev. Cell 17, 75-86.

- 47. Sun, Z., Guo, T., Liu, Y., Liu, Q., and Fang, Y. (2015). The roles of Arabidopsis CDF2 in transcriptional and posttranscriptional regulation of primary microRNAs. PLoS Genet. 11, e1005598.
- 48. Go, Y.S., Kim, H., Kim, H.J., and Suh, M.C. (2014). Arabidopsis cuticular wax biosynthesis is negatively regulated by the DEWAX gene encoding an AP2/ERF-type transcription factor. Plant Cell 26, 1666-1680.
- 49. Shaikhali, J., de Dios Barajas-Lopéz, J., Ötvös, K., Kremnev, D., Garcia, A.S., Srivastava, V., Wingsle, G., Bako, L., and Strand, A. (2012). The CRYPTOCHROME1-dependent response to excess light is mediated through the transcriptional activators ZINC FINGER PROTFIN EXPRESSED IN INFLORESCENCE MERISTEM LIKE1 and ZML2 in Arabidopsis. Plant Cell 24, 3009-3025.
- 50. Abbas, M., Berckhan, S., Rooney, D.J., Gibbs, D.J., Vicente Conde, J., Sousa Correia, C., Bassel, G.W., Marín-de la Rosa, N., León, J., Alabadí, D., et al. (2015). Oxygen sensing coordinates photomorphogenesis to facilitate seedling survival. Curr. Biol. 25, 1483-1488.
- 51. Kieffer, M., Master, V., Waites, R., and Davies, B. (2011). TCP14 and TCP15 affect internode length and leaf shape in Arabidopsis. Plant J. 68,
- 52. Walter, M., Chaban, C., Schütze, K., Batistic, O., Weckermann, K., Näke, C., Blazevic, D., Grefen, C., Schumacher, K., Oecking, C., et al. (2004). Visualization of protein interactions in living plant cells using bimolecular fluorescence complementation. Plant J. 40, 428-438.
- 53. Shannon, P., Markiel, A., Ozier, O., Baliga, N.S., Wang, J.T., Ramage, D., Amin, N., Schwikowski, B., and Ideker, T. (2003). Cytoscape: a software environment for integrated models of biomolecular interaction networks. Genome Res. 13, 2498-2504.
- 54. Pavlidis, P., and Noble, W.S. (2003). Matrix2png: a utility for visualizing matrix data. Bioinformatics 19, 295-296.
- 55. Zhang, X., Henriques, R., Lin, S.-S., Niu, Q.-W., and Chua, N.-H. (2006). Agrobacterium-mediated transformation of Arabidopsis thaliana using the floral dip method. Nat Protoc 1, 641-646.
- 56. Dill, A., Thomas, S.G., Hu, J., Steber, C.M., and Sun, T.P. (2004). The Arabidopsis F-box protein SLEEPY1 targets gibberellin signaling repressors for gibberellin-induced degradation. Plant Cell 16, 1392-1405.
- 57. Hickman, R., Hill, C., Penfold, C.A., Breeze, E., Bowden, L., Moore, J.D., Zhang, P., Jackson, A., Cooke, E., Bewicke-Copley, F., et al. (2013). A local regulatory network around three NAC transcription factors in stress responses and senescence in Arabidopsis leaves. Plant J. 75, 26-39.
- 58. Pruneda-Paz, J.L., Breton, G., Nagel, D.H., Kang, S.E., Bonaldi, K., Doherty, C.J., Ravelo, S., Galli, M., Ecker, J.R., and Kay, S.A. (2014). A genome-scale resource for the functional characterization of Arabidopsis transcription factors. Cell Rep. 8, 622-632.
- 59. Altmann, M., Altmann, S., Falter, C., and Falter-Braun, P. (2018). Highquality yeast-2-hybrid interaction network mapping. Curr. Protoc. Plant Biol. 3, e20067.
- 60. Gendrel, A.-V., Lippman, Z., Martienssen, R., and Colot, V. (2005). Profiling histone modification patterns in plants using genomic tiling microarrays. Nat. Methods 2, 213-218.

Current Biology Article



STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		····
Anti-HA tag antibody - ChIP Grade	Abcam	Cat#ab9110
Goat Anti-Mouse IgG-Peroxidase antibody	Sigma-Aldrich	Cat#A4416
Bacterial and Virus Strains	Cigina radion	
Escherichia coli DH5α	[27]	
Agrobacterium tumefaciens (strain GV3101)	[27]	
Chemicals, Peptides, and Recombinant Proteins		
Phusion DNA Polymerase	NEB	M0530
Murashige & Skoog Medium	Duchefa Biochemie	Cat#M0221
β-estradiol	Sigma-Aldrich	Cat#E2758
3-Amino-1,2,4-triazole	Sigma-Aldrich	Cat#A8056
Taq DNA Polymerase	NEB	Cat#M0273
Critical Commercial Assays		
RNeasy PowerPlant Kit	QIAGEN	Cat#13500-50
RNeasy Plant Mini Kit	QIAGEN	Cat#74903
Plant DNeasy Kit	QIAGEN	Cat#69104
Gateway BP Clonase II Enzyme mix	Invitrogen	Cat#11789020
Gateway LR Clonase II Enzyme mix	Invitrogen	Cat#11791020
SuperScript II Reverse Transcriptase	Invitrogen	Cat#18064022
Brilliant II QPCR Master Mix with ROX	Agilent	Cat#600806
Pierce Protein A/G Magnetic Beads	Invitrogen	Cat#88802
Experimental Models: Cell Lines	minagen	Cati 60002
Yeast strain AH109 (MATa)	Clontech	N/A
Yeast strain Y187 (MATa)	Clontech	Cat#630457
Yeast strain Y8800 (MATa)	Clontech	N/A
Yeast strain Y8930 (MATa)	Clontech	N/A
Experimental Models: Organisms/Strains	Cionicon	1471
Arabidopsis thaliana Col-0	[27]	
Arabidopsis thaliana At2g28810	Arabidopsis Biological Resource Center	Cat#SALK_056801C
Arabidopsis thaliana erf7	Arabidopsis Biological Resource Center	Cat#SALK_032229
Arabidopsis thaliana myb30	Arabidopsis Biological Resource Center	Cat#SALK_027644C
Arabidopsis thaliana pat1	Arabidopsis Biological Resource Center	Cat#SALK_064220C
Arabidopsis thaliana TCP14 TPTD	Arabidopsis Biological Resource Center	Cat#TPT_3.47620.1D
Arabidopsis thaliana TCP15 TPTC	Arabidopsis Biological Resource Center	Cat#TPT_1.69690.1C
Arabidopsis thaliana RAP2.3 TPTD	Arabidopsis Biological Resource Center	Cat#TFT_3.16770.1D
Arabidopsis thaliana Col-0 XVE::EXPA9	This study	N/A
Arabidopsis thaliana top14-4 15-3	This study This study	N/A
XVE:EXPA9	iiis study	IVA
Arabidopsis thaliana tcp14-4 35S::TCP14- HA	This study	N/A
Arabidopsis thaliana cdf2-1	[46]	N/A
Arabidopsis thaliana cdf3-1	[46]	N/A
Arabidopsis thaliana cdf2-13-1	[46]	N/A
Arabidopsis thaliana CDF2-OX	[47]	N/A
,		(Continued on next page)

(Continued on next page)





Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Arabidopsis thaliana dewax	[48]	N/A
Arabidopsis thaliana iDEWAX	[48]	N/A
Arabidopsis thaliana zml1	[49]	N/A
Arabidopsis thaliana rap2.2	[50]	N/A
Arabidopsis thaliana ebp	[50]	N/A
Arabidopsis thaliana rap2.12	[50]	N/A
Arabidopsis thaliana rap2.2 2.12	[50]	N/A
Arabidopsis thaliana rap2.12 ebp	[50]	N/A
Arabidopsis thaliana rap2.2 2.3 2.12	[50]	N/A
Arabidopsis thaliana tcp14-4	[51]	N/A
Arabidopsis thaliana tcp15-3	[51]	N/A
Arabidopsis thaliana tcp14-4 15-3	[51]	N/A
Oligonucleotides		
Primers	Table S1	N/A
Recombinant DNA		
pHISLEU2GW	[32]	N/A
pHISLEU2GW-pEXPA1	This study	N/A
pHISLEU2GW-pEXPA3	This study	N/A
pHISLEU2GW-pEXPA8	This study	N/A
pHISLEU2GW-pEXPA9	This study	N/A
pHISLEU2GW-pEXPA10	This study	N/A
pHISLEU2GW-pEXPA15	This study	N/A
pHISLEU2GW-pEXPA20	This study	N/A
oDEST-AD	Invitrogen	N/A
oDEST-DB	Invitrogen	N/A
oDEST-DB-RGA∆N	This study	N/A
oDEST-DB-GAIAN	This study	N/A
oSPYNE	[52]	N/A
oSPYNE-RGL2	This study	N/A
oSPYCE	[52]	N/A
pSPYCE-TCP14	This study	N/A
oSPYCE-TCP15	This study	N/A
oSPYCE-ZML1	This study	N/A
oGWB14	RIKEN BRC	N/A
oGWB14-TCP14	This study	N/A
pER8-GW	[30]	N/A
oER8-GW-EXPA9	This study	N/A
Software and Algorithms		
CytoScape	[53]	https://cytoscape.org/
matrix2png	[54]	https://matrix2png.msl.ubc.ca/index.htm

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources should be directed to and will be fulfilled by the Lead Contact, George W. Bassel (george.bassel@warwick.ac.uk).

Materials Availability

This study did not generate new unique reagents.

Article



Data and Code Availability

All datasets generated or analyzed during this study are included in the manuscript. The interactions from this publication have been submitted to the IMEx (http://www.imexconsortium.org) consortium through IntAct [44] and assigned the identifier IM-27910.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Plant Material and growth conditions

Seeds were grown on half-strength Murashige and Skoog (MS) medium with 0.8% Agar under 16h/8h light/dark cycles. 2-week-old seedlings were transferred to soil and grown in a greenhouse. Freshly harvested seeds were stored at room temperature for 1-2 months. *Arabidopsis* T-DNA insertion lines were obtained from *Arabidopsis* Biological Resource Center (ABRC): *At2g28810* mutant (SALK_056801C), *erf7* (SALK_032229), *myb30* (SALK_027644C), *pat1* (SALK_064220C). Following lines were kindly provided by authors and have been previously described: *cdf2-1*, *cdf3-1*, *cdf2-13-1*, and *CDF2-OX* [46, 47]; *dewax* and *iDEWAX* [48]; *zml1* [49]; *rap2.2*, *ebp*, *rap2.12*, *rap2.22.12*, *rap2.12* ebp, and *rap2.2 2.3 2.12* [50]; *tcp14-4*, *tcp15-3*, and *tcp14-4 15-3* [51].

METHOD DETAILS

Plasmid construction and plant transformation

To generate the estrogen-inducible XVE::EXPA9 construct, full-length cDNA coding sequence of EXPA9 was amplified using primers described in Table S1 with Phusion DNA Polymerase (NEB, M0530), and recombined in pDONRzeo using BP clonase II (Invitrogen). The fragment was subsequently transferred from the entry vector into the estrogen inducible pER8GW [30] by LR clonase reaction (Invitrogen).

For the 35S::TCP14-HA construct, the coding sequence without the stop codon of TCP14 was amplified and inserted into pDONR-zeo by BP reaction, then recombined with pGWB14 by LR reaction (Invitrogen).

The resulting plasmids were transformed into *Agrobacterium tumefaciens* strain GV3101 and introduced into *Arabidopsis thaliana* by the floral dipping method [55]. Transgenic progeny seeds were selected on half-strength MS medium with 20 mg/L hygromycin [55].

Phenotypic analyses

Germination assays were performed by surface sterilizing seeds and pipetting them onto 1/2 MS medium supplemented with PAC as required to achieve specific concentrations. For estrogen inducible lines, seeds were either treated with 30 μ M β -estradiol or ethanol (mock). After sowing, seeds were kept at 4°C under dark conditions for 3 days, and then transferred to a controlled growth chamber under 16h/8h light-dark cycles at 22°C.

Gene expression analyses

Total RNA was isolated either from *Arabidopsis* seeds using the RNeasy PowerPlant Kit or from seedlings using RNeasy Plant Mini Kit (QIAGEN) according to the manufacturer's instructions. DNA in RNA samples was removed with DNase I (Thermo Fisher Scientific) and RNA was reverse-transcribed into cDNA using SuperScript II Reverse Transcriptase (Invitrogen). Quantitative PCR was performed in 96-well blocks with Brilliant II QPCR Master Mix with ROX (Agilent, #600806) on the AriaMx Real-Time PCR system. Gene expression was normalized using internal control *GLYCERALDEHYDE-3-PHOSPHATE DEHYDROGENASE C SUBUNIT* (*GAPC*) (At3g04120) [56]. RT-PCR was performed with Taq DNA Polymerase (NEB, #M0273) on a thermal cycler. Analysis of *EXPA9* were subjected to amplication for 26 and 30 cycles, and analysis of *GAPC* was followed by 26 cycles.

Yeast one hybrid screening of Arabidopsis cDNA libraries

TFs included in the Y1h screening library were determined by examining those expressed in the germinating embryo at 7 HAI and 12 HAI [25], prior to or concurrent with the induction of *EXPA* gene expression. 168 embryo expressed TFs were available in the REGIA + REGULATORS RR Library [32], in which TF cDNAs fused to an N-terminal GAL4-activation domain in *pDEST22* (Invitrogen). Prey clones were transformed into yeast strain AH109 (MATa, Clontech) according to manufacturer's instructions.

Y1h screening was performed as described previously [57]. Promoter sequences for EXPA1 (At1g69530), EXPA3 (At2g37640), EXPA8 (At2g40610), EXPA9 (At5g02260), EXPA10 (At1g26770), EXPA15 (At2g03090), and EXPA20 (At4g38210) consisting of 300 ~400 bp fragments with 100 bp overlaps were cloned into pDONRZeo, and then into the destination vector pHISLEU2GW. Bait vectors were then transformed into yeast strain Y187 (MATa, Clontech) and tested for autoactivation. Several fragments required the addition of 3-amino-1,2,4-triazol (3-AT) to enable selection of interacting proteins. Mating was performed by mixing 3 μ L of each bait and prey onto YPDA plates, with subsequent replica-plating (after 24 h) onto SD-Leu-Trp and SD-Leu-Trp-His (+ 3-AT where required) and grown for two days to confirm mating and provide selection for interaction. All positive interactions in yeast were re-tested and sequenced.

Protein-protein interaction assays

Protein interaction screening in yeast was performed using N-terminal truncations of RGA and GAI. These were cloned into *pDEST-DB* and screened by yeast mating against *EXPA* promoter-binding TFs fused to a GAL4 activation domain (AD) expressed via *pDEST-*





AD [58] following previously described protocols [59]. To detect AD auto-activators, the AD-fused transcription factor collection in MATa Y8800 yeast strains was mated with the empty pDEST-DB expressing MATa Y8930, then selected using 1 mM 3-amino-1,2,4-triazole (3-AT). Protein interaction screens were repeated twice with 1 mM 3-AT and a second time with 2 and 3 mM 3-AT for RGA and GAI, respectively.

BiFC protein-protein interaction in planta

Full-length TF coding sequences and RGL2 were cloned into pSPYCE or pSPYNE vectors containing either C- or N-terminal portions of yellow fluorescence protein [52] and transformed into Agrobacterium strain GV3101. Bacteria were grown overnight at 28°C in LB medium and resuspended in infiltration buffer (10 mM MES, pH5.7, 10 mM MgCl₂, and 150 mM acetosyringone). To visualize the fluorescence signal, different combinations of bacteria cells were co-infiltrated into the leaves of 4-week-old Nicotiana benthamiana. YFP signals were assayed 3 days after inoculation and excited at 488 nm using a Zeiss LSM510 microscope.

Chromatin Immunoprecipitation

ChIP was performed as described previously with minor modifications [60]. Seedlings (6 days after imbibition) grown on half-strength MS medium supplemented with 10 μM GA₃ were used for CHIP-qPCR analyses. Chromatin was isolated from 2 g of seedlings and sheared by sonication to fragments of 400 bp. Following isolation, sonicated chromatin was precleared with Dynabeads Protein A/G (Invitrogen) and then immunoprecipitated overnight using Anti-HA (Abcam, ab9110) polyclonal antibodies. qPCR analyses were performed using Brilliant II QPCR Master Mix with ROX (Agilent, #600806). Fold enrichment was calculated by normalizing the amount of target fragment first to the internal control gene (ACTIN2) and then to the corresponding amount in the wild-type (Col-0). The primers used for ChIP-gPCR are listed in Table S1.

Bioinformatics analyses

The network graph was visualized using CytoScape (version 3.7) [53]. Expression profile of EXPAs was generated in the Matrix2png program (https://matrix2png.msl.ubc.ca/index.html) [54].

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analysis was implemented using either EXCEL or SPSS software. Data shown in the figures are representing an average of biological replicates. All seed germination test had at least 50 seeds in each biological replication. Two-tailed Student's t test was performed with the t.test() function in EXCEL. One-way ANOVAs with post hoc Turkey test was carried out by SPSS (version 16.0) at a significance level of p < 0.05. Details of statistical tests are provided in figure legends.