1 The IncRNA APOLO interacts with the transcription factor WRKY42 to trigger root hair cell expansion in

2 response to cold

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26 SUMMARY

- 27 The IncRNA APOLO directly regulates the transcription of the root hair-master gene RHD6. In response to
- cold, APOLO is induced and it decoys the H3K27me3-binding protein LHP1 away from RHD6. In addition,
- 29 APOLO modulates the binding of the transcription factor WRKY42 to the RHD6 promoter at low tempera-
- 30 tures.
- 31

32 ABSTRACT

33

34 Plant long noncoding RNAs (IncRNAs) have emerged as important regulators of chromatin dynamics, 35 impacting on transcriptional programs leading to different developmental outputs. The IncRNA AUXIN 36 REGULATED PROMOTER LOOP (APOLO) directly recognizes multiple independent loci across the 37 Arabidopsis genome and modulates their three-dimensional chromatin conformation, leading to 38 transcriptional shifts. Here, we show that APOLO recognizes the locus encoding the root hair (RH) master 39 regulator ROOT HAIR DEFECTIVE 6 (RHD6) and controls RHD6 transcriptional activity leading to cold-40 enhanced RH elongation through the consequent activation of the transcription factor gene RHD6-like 41 RSL4. Furthermore, we demonstrate that APOLO interacts with the transcription factor WRKY42 and 42 modulates its binding to the RHD6 promoter. WRKY42 is required for the activation of RHD6 by low 43 temperatures and WRKY42 deregulation impairs cold-induced RH expansion. Collectively, our results 44 indicate that a novel ribonucleoprotein complex involving APOLO and WRKY42 forms a regulatory hub 45 which activates *RHD6* by shaping its epigenetic environment and integrates signals governing RH growth 46 and development.

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48 Words: 159

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50 Key words: root hairs; long noncoding RNAs; APOLO; RHD6; WRKY42; cold temperature

51 **INTRODUCTION**

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53 Root hairs (RHs) are single cell projections developed from specialized epidermal trichoblast cells able to increase their size several hundred times in a polar manner to reach and promote the uptake of 54 55 water-soluble nutrients, interact with soil microorganisms and support anchor to the plant. The specifi-56 cation of epidermal cells into RHs is a complex process whose underlying mechanisms are partially un-57 derstood. In Arabidopsis thaliana, RH cell fate is controlled by a developmental program involving a 58 complex of transcription factors (TFs) promoting the expression of the homeodomain protein GLABRA 2 59 (GL2) (Ryu et al., 2005; Song et al., 2011; Schiefelbein et al., 2014; Balcerowicz et al., 2015). GL2 blocks 60 RH development by inhibiting the transcription of the master regulator ROOT HAIR DEFECTIVE 6 (RHD6) 61 (Lin et al., 2015). In the cells that differentiate into RHs (known as trichoblasts), a second TF complex 62 suppresses GL2 expression (Schiefelbein et al., 2014), forcing the cells to enter the RH cell fate program 63 via the concomitant activation of RHD6 along with downstream TFs (Menand et al., 2007; Pires et al., 64 2013). Briefly, RHD6 together with its homolog RSL1 (ROOT HAIR DEFECTIVE 6 LIKE 1) induce the expres-65 sion of TFs from the bHLH family, including RSL2 (ROOT HAIR DEFECTIVE 6 LIKE 2) and RSL4 (ROOT HAIR DEFECTIVE 6 LIKE 4), ultimately triggering the differentiation of the RHs and their subsequent polarized 66 tip-growth (Karas et al., 2009; Yi et al., 2010; Bruex et al., 2012). In addition, it was proposed that RSL4 67 68 controls the expression of a small subset of nearly 125 genes (Won et al., 2009; Yi et al., 2010; Datta et 69 al., 2015; Vijayakumar et al., 2016), including several cell wall extensins (EXTs) (Ringli, 2010; Velasquez et 70 al., 2011) sufficient to promote RH growth (Hwang et al., 2017).

71 RH expansion is regulated both by cell-intrinsic factors (e.g. endogenous phytohormones such as 72 auxin) and external environmental signals (e.g. phosphate (Pi) availability in the soil) (Mangano et al., 73 2017; Bhosale et al., 2018). Pi starvation is one of the key environmental factors promoting rapid RH 74 growth (Yi et al., 2010; Datta et al., 2015; Vijayakumar et al., 2016). In Arabidopsis, it triggers RSL4 ex-75 pression via an enhanced auxin production, activating downstream effector genes mediating cell growth 76 (Yi et al., 2010; Datta et al., 2015; Mangano et al., 2017; Marzol et al., 2017; Bhosale et al., 2018). Ac-77 cordingly, several auxin-related TFs have been implicated in Pi-starvation signaling in roots, including 78 WRKY proteins that control the expression of the Pi transporter families Pi-permease PHO1 and PHOS-79 PHATE TRANSPORTER (PHT) (Devaiah et al., 2007; Chen et al., 2009; Wang et al., 2014; Su et al., 2015). 80 Under Pi-sufficient conditions, WRKY6 and WRKY42 bind to W-boxes of the PHO1 promoter and suppress 81 its expression. During Pi starvation, WRKY42 is degraded by the 26S proteasome pathway, resulting in 82 the activation of PHO1 transcription (Chen et al., 2009; Su et al., 2015). In addition, WRKY42 functions as

a positive regulator of *PHT1;1*, by binding to its promoter under Pi-sufficient condition (Su et al., 2015).
Overall, WRKY42 is part of the components activating root early-responses to Pi starvation, although its
role in controlling RH growth remains unexplored.

86 In recent years, plant long noncoding RNAs (IncRNAs) have emerged as important regulators of 87 gene expression, and several among them, have been functionally linked to Pi homeostasis. For instance, 88 the IncRNA INDUCED BY PHOSPHATE STARVATION 1 (IPS1) can sequester the Pi starvation-induced mi-89 croRNA miR-399, attenuating miR-399-mediated repression of PHO2, a gene encoding an E3 ligase af-90 fecting Pi uptake (Franco-Zorrilla et al., 2007). In addition, the cis-natural antisense (cis-NAT) transcript 91 PHO1;2, induced under Pi deficiency, was shown to promote the translation of the PHO1;2 mRNA in-92 volved in Pi loading into the xylem. The expression of this cis-NAT is associated with the transport of the 93 sense-antisense RNA pair toward the polysomes (Jabnoune et al., 2013). More recently, it was shown 94 that the IncRNA AUXIN REGULATED PROMOTER LOOP (APOLO) recognizes multiple spatially independent 95 genes by sequence complementarity and DNA-RNA duplex formation, known as R-loops. Upon recogni-96 tion, APOLO shapes the three-dimensional (3D) conformation of its target regions by decoying the Poly-97 comb Repressive Complex 1 (PRC1) component LIKE HETEROCHROMATIN PROTEIN 1 (LHP1), thereby 98 regulating their transcription (Ariel et al., 2014; Ariel et al., 2020).

99 Here, we show that the IncRNA *APOLO* directly regulates a subset of genes involved in RH devel-100 opment, including the master regulator of RH initiation *RHD6*. *APOLO* activates *RHD6* transcription by 101 modulating the formation of a local chromatin loop encompassing its promoter region, an epigenetic 102 regulatory mechanism likely involving PRC1 and PRC2 components. Furthermore, we found that *APOLO* 103 interacts with the TF WRKY42, forming a new hub that regulates *RHD6* to induce RH growth in response 104 to low temperatures. RHD6-mediated induction of RH expansion likely occurs through the transcriptional 105 activation of the TF-encoding gene *RSL4*, which emerged as a key factor in the response to cold.

106

107 **RESULTS**

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109 APOLO regulates root hair cell elongation in response to low temperatures

Based on Chromatin Isolation by RNA Purification (ChIRP-Seq) performed in wild-type (WT, Col-0) plants, it was previously reported that the IncRNA *APOLO* recognizes a subset of independent loci enriched in categories related to cell wall composition and organization (Ariel et al., 2020). A closer look at *APOLO bona fide* targets allowed us to identify seventeen genes involved in RH growth and expansion (**Supplementary Table 1**), a process dependent on cell wall remodeling molecules, including EXTs and EXT-related proteins (Ringli, 2010; Lamport et al., 2011; Velasquez et al., 2011; Velasquez et al., 2015; Marzol et al., 2018). Interestingly, according to single-cell RNA-seq datasets (Zhang et al., 2019) *APOLO* transcripts are enriched in RH cells (**Supplementary Figure 1A**). Notably, sixteen *APOLO* direct targets were upregulated upon *APOLO* over-expression (**Supplementary Table 2**). Furthermore, 52 additional RHrelated genes were upregulated in *35S:APOLO* seedlings (**Supplementary Table 2**) (Ariel et al., 2020). Among them, the RH central TFs *RHD6* (as a direct target), *RSL2* and *RSL4* (as indirectly regulated) were induced upon *APOLO* over-expression.

It was reported that the APOLO locus is targeted by the RNA-polymerase Pol V and silenced by 122 123 RNA-directed DNA Methylation (RdDM, (Ariel et al., 2014)). A search in a small RNA-Seq performed in WT 124 roots subjected to different temperature treatments (Gyula et al., 2018) revealed that RdDM-related 24nt 125 siRNA accumulation over the APOLO locus is less abundant at low temperatures (Supplementary Figure 126 **1B**), suggesting that APOLO transcription is regulated by cold. Accordingly, we found that APOLO tran-127 scriptional accumulation increases in roots after 24h at 10°C (Figure 1A). An analysis of the promoter 128 activity of the 5.2kb region upstream APOLO (Ariel et al., 2020) directing the expression of a GFP reporter 129 gene, additionally revealed a higher transcriptional activity at low temperatures in the RHs (Figure 1B). 130 Strikingly, we observed that two RNAi-APOLO (repression of approximately 90%; Ariel et al., 2014) and 131 35S:APOLO independent lines (over expression of 30-fold and 60-fold, respectively; Ariel et al., 2020) 132 exhibit a basal increase of RH length at 22°C, and uncovered a strong induction of RH elongation in WT 133 and RNAi-APOLO at 10°C, in contrast to 35S:APOLO lines (Figure 1C). Accordingly, RHD6 is induced in re-134 sponse to cold in WT roots, whereas RNAi-APOLO and 35S:APOLO roots display higher RHD6 basal levels 135 than the WT (Figure 1D). Notably, RHD6 transcript levels are not further induced by cold in 35S:APOLO 136 roots (Figure 1D), in agreement with the RH phenotype (Figure 1C). Collectively, our findings suggest that 137 APOLO participates in the induction of cold-mediated RH elongation and a deregulation of APOLO tran-138 script levels can impact RH growth.

139 Previous studies pointed out a key role of RHD6 (together with RSL1) in RH development, which 140 is mediated by RSL4 and RSL2 as downstream regulators of RH cell elongation (Menand et al., 2007; Pires 141 et al., 2013). Notably, RHD6, RSL2 and RSL4 transcript levels were upregulated in 35S:APOLO seedlings 142 (Supplementary Table 2; (Ariel et al., 2020)), although only RHD6 was identified as a direct target of 143 APOLO (Ariel et al, 2020). In agreement with RHD6 transcriptional behavior (Figure 1D), RSL2 and RSL4 144 basal transcript levels are higher in RNAi-APOLO and 35S:APOLO compared to WT. Interestingly, RSL2 145 and RSL4 are induced by cold in WT and RNAi-APOLO, but not in 35S:APOLO roots (Supplementary Fig-146 ure 2A), suggesting that low temperatures can activate these two genes still in the absence of APOLO

147 and bypassing RHD6 for RH expansion (Figure 1C). Thus, we assessed if these TFs were also controlling 148 the promotion of RH growth by low temperatures. To this end, we tested how rhd6/rsl1/rsl4 and rsl2, 149 rsl4 and double mutant plants rsl2/rsl4 respond to low temperatures in comparison with control condi-150 tions (Supplementary Figure 2A). The *rsl2* mutant was highly responsive to low temperatures in a similar 151 manner to WT while rsl4 was impaired in the response to cold. The double mutant rsl2/rsl4 and the triple 152 mutant rhd6/rsl1/rsl4 did not develop RHs in either of the two conditions. In addition, constitutive ex-153 pression of RSL4 (35S:RSL4) as well as its expression under the control of the RH specific EXPANSIN7 promoter (EXP7p:RSL4) boosted basal RH growth without further enhancement in response to cold 154 155 (Supplementary Figure 2B). These results demonstrate that RSL4 is a key factor mediating RHD6 activa-156 tion of RH growth at low temperature, and RSL2 might participate to a lower extent.

157 Nutrient unavailability is known to activate RH expansion through a transcriptional reprogram-158 ming governed by RHD6 and downstream TFs. The quantification of RH growth of WT plants in response 159 to increasing concentrations of nutrients (0.5X to 2.0X MS (Murashige and Skoog) medium) indicates that 160 high concentrations impair RH growth triggered by low temperatures (Supplementary Figure 3A). In a 161 similar way, an increase in agar concentration in the MS medium (from 0.8% to 2.5%), which likely restrains nutrient mobility (Singha, S., Townsend, E.C. and Oberly, 1985; Nonami and Boyer, 1989; 162 163 Ghashghaie et al., 1991; Buah et al., 1999) blocks cold-induced RH expansion (Supplementary Figure 3B). 164 Altogether, these observations suggest that low temperatures restrict nutrient mobility and availability in 165 the culture medium, leading to the promotion of polar RH growth.

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167 *APOLO* directly modulates the three-dimensional chromatin conformation of the root hair specific lo-168 **cus** *RHD6*

169 Among APOLO targets involved in RH development, we found the master regulator of RH initia-170 tion RHD6 (Menand et al., 2007; Pires et al., 2013). The epigenetic profile of the RHD6 locus corresponds 171 to typical APOLO targets (Figure 2A, (Ariel et al., 2020)), including H3K27me3 deposition (track 1), LHP1 172 recognition (track 2, chromatin immunoprecipitation (ChIP)-Seq, (Veluchamy et al., 2016)), and APOLO 173 binding regions (tracks 3 to 5, chromatin isolation by RNA purification (ChIRP)-Seq, (Ariel et al., 2020)). A 174 GAAGAA box, shown to be important for APOLO target recognition (Ariel et al., 2020) is located in the 175 RHD6 locus and coincides with APOLO binding site. In addition, a peak of DNA-RNA hybrid immunopre-176 cipitation (DRIP)-Seq from root samples indicates the presence of an R-loop coinciding with APOLO 177 recognition sites over RHD6 (tracks 6 to 8, (Xu et al., 2020)).

178 Remarkably, APOLO recognition and R-loop formation are also detectable over RHD6 neighbor 179 gene, located 3.2 kb upstream RHD6 transcription start site (Figure 2A). According to DpnII Hi-C datasets 180 from Arabidopsis seedlings (Liu et al., 2016), a chromatin loop encompassing the intergenic region up-181 stream RHD6 was detected (Figure 2B), and coincides with APOLO binding-sites (Figure 2A, ChIRP-Seq). 182 By performing a ChIRP-qPCR with two independent sets of biotinylated probes to purify APOLO (ODD and 183 EVEN; (Ariel et al., 2020)) and one additional set used as a negative control (LacZ), we confirmed that 184 APOLO RNA-RHD6 DNA interaction occurs in WT and is lost in APOLO knockdown (RNAi) seedlings ((Ariel 185 et al., 2014); Figure 2C). In addition, the quantification of relative RHD6 loop formation in RNAi-APOLO 186 and 35S:APOLO (Ariel et al., 2020) seedlings, revealed impaired loop formation in both lines (Figure 2D), 187 hinting at a stoichiometric requirement of APOLO for RHD6 chromatin loop formation. Chromatin loop 188 formation (Figure 2D) is in agreement with RHD6 basal levels in 35S:APOLO and RNAi-APOLO lines (Figure 189 **1D**), suggesting that the chromatin loop including *RHD6* promoter region precludes transcription. Alto-190 gether, our results indicate that APOLO IncRNA directly regulates RHD6 transcriptional activity by fine-191 tuning local chromatin 3D conformation.

192 It was previously reported that PRC2 actively participates in the regulation of RH growth (Ikeuchi 193 et al., 2015) and that the RHD6 locus exhibits H3K27me3 deposition and LHP1 recognition (Figure 2A; 194 (Veluchamy et al., 2016)). Considering that the IncRNA APOLO interacts with the PRC1 component LHP1 195 in vivo (Ariel et al., 2014; Ariel et al., 2020), we decided to explore the role of PRC1 and PRC2 in APOLO-196 mediated RHD6 activation at low temperatures. At 22°C, RHD6 suffers a reduction of H3K27me3 in the 197 PRC2 mutant curly leaf (clf), in contrast to the PRC1 mutant lhp1 (Supplementary Figure 4A; (Veluchamy 198 et al., 2016)). Interestingly, we observed that H3K27me3 deposition and LHP1 binding diminish in WT 199 roots treated for 24h at 10°C compared to 22°C (Supplementary Figure 4B), consistent with the induction 200 of RHD6 in response to cold (Figure 2E). Moreover, *lhp1* and *clf* mutants exhibit a basal decrease of RH 201 length together with a slight decrease of cold-induced RH elongation in *lhp1*, and a strong decrease of 202 cold-induced RH elongation in *clf* (Supplementary Figure 4D). Consistently, although the decrease in 203 H3K27me3 deposition results in higher basal transcript levels of RHD6 in the clf background, RHD6 tran-204 scriptional activation by cold is abolished in the *clf* and *lhp1* mutants (**Supplementary Figure 4C**), hinting 205 at an important role of chromatin rearrangement for *RHD6* activation in response to cold.

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207 APOLO interacts with the transcription factor WRKY42 to coordinate the activation of RHD6

208 In order to uncover novel actors involved in cold-induced transcriptional regulation of RH 209 growth, we aimed at identifying *APOLO* protein partners. To this end, we performed an exploratory

210 ChIRP on WT seedlings using two independent set of biotinylated probes to purify APOLO (ODD and 211 EVEN; (Ariel et al., 2014; Ariel et al., 2020)), and one additional set used as a negative control (against 212 LacZ), as recently described (Rigo et al., 2020). Co-purified proteins were precipitated and analyzed by 213 mass spectrometry. Among the potential APOLO partners (i.e. identified with at least two hits in ODD 214 and EVEN samples, but absent in LacZ-ChIRP), we found the WRKY42 protein, a TF involved in the re-215 sponse to Pi starvation (Su et al., 2015), an environmental condition that promotes RH cell expansion 216 (Bhosale, et al., 2018) in a similar manner to low temperatures. Thus, APOLO-WRKY42 interaction was 217 validated by RNA immuno-precipitation (RIP-qPCR) in tobacco leaves and in Arabidopsis plants transitory 218 or stably transformed with 35S:WRKY42:GFP, respectively (Figure 3A). Interestingly, according to the 219 Arabidopsis eFP Browser (Waese et al., 2017), WRKY42 is induced in roots when seedlings are subjected 220 to 4°C for 24h (Figure 3B). At 10°C, we observed that WRKY42 transcriptional accumulation augments 221 significantly in roots (Figure 3C). Notably, 13 out of the 17 APOLO targets contained between 1 and 4 222 canonical WRKY TF binding sites (W-box) in their promoters, including RHD6 (Supplementary Table 1). 223 By using a 35S:WRKY42:GFP line, we determined that WRKY42 can directly bind to the promoter region 224 of RHD6 (Figure 3D). Accordingly, the over-expression of WRKY42 (35S:WRKY42:GFP line) led to a basal 225 increase of RHD6 levels (Figure 3E) and RH elongation (Figure 3F) at ambient temperature, mimicking the 226 effect of cold. On the contrary, cold-mediated induction of RHD6 is abolished in the wrky42 mutant ((Su 227 et al., 2015); Figure 3E), which consistently exhibits shorter RHs at 22°C and almost no RH elongation at 228 low temperatures (Figure 3F). Taken together, these results suggest that the APOLO-interacting TF 229 WRKY42 is an important regulator of *RHD6*-mediated RH growth in response to cold.

230 We thus wondered to what extent WRKY42 regulates the epigenetic landscape of the RHD6 lo-231 cus. We first observed that H3K27me3 deposition over RHD6 is significantly augmented in the wrky42 232 mutant background, in contrast to AZG2, an APOLO target non-related to WRKY42 (Figure 4A), con-233 sistent with reduced RHD6 basal levels reported in wrky42 (Figure 3E). Therefore, we evaluated the mu-234 tual contribution of APOLO and WRKY42 to their respective recognition of the RHD6 locus. APOLO ChIRP-235 qPCR in the WT and wrky42 mutant (Figure 4B) revealed similar binding to RHD6, indicating that WRKY42 236 does not participate in APOLO-target recognition. Reciprocally, we assessed the control of APOLO over 237 WRKY42 recognition of the RHD6 locus. To this end, we transformed transiently A. thaliana leaves of WT, 238 35S:APOLO and RNAi-APOLO plants with the construct 35S:WRKY42:GFP. Observation using confocal 239 microscopy indicated that WRKY42:GFP is localized in the nucleus in the three genetic backgrounds 240 (Supplementary Figure 5). Remarkably, WRKY42:GFP ChIP-qPCR revealed that WRKY42 binding to RHD6 241 promoter is impaired both in 35S:APOLO and RNAi-APOLO plants compared to WT (Figure 4C), hinting at a stoichiometric role of *APOLO* in the modulation of TF-chromatin interaction. To further confirm our
observations, chromatin was extracted from *35S:WRKY42:GFP* seedlings and increasing amounts of *in vitro* transcribed *APOLO* were added before cross-link and regular WRKY42 ChIP over *RHD6*. Strikingly,
increasing concentrations of *APOLO* gradually decoy WRKY42 away from the *RHD6* locus (Figure 4D),
further supporting the hypothesis of a stoichiometric regulation of *APOLO* over the activity of its partner
TF.

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249 DISCUSSION

Our results indicate that the regulation of *RHD6* expression in response to cold depends on Polycomb-dependent H3K27me3 dynamic deposition. The WRKY42-*APOLO* complex modulates the epigenetic environment of *RHD6*, activating its transcription and promoting RH growth at low temperatures. *RHD6* activation further triggers the expression of *RSL2* and *RSL4* that control the transcriptional RH program inducing cell expansion in response to cold (**Figure 4E**).

255 Cell fate determination in the epidermis has been extensively studied (Ryu et al., 2005; Song et 256 al., 2011; Schiefelbein et al., 2014; Balcerowicz et al., 2015). Once trichoblast cells differentiate in the 257 root epidermis, RHs develop as fast polar growing protuberances in response to endogenous and envi-258 ronmental signals (Menand et al., 2007; Yi et al., 2010; Pires et al., 2013; Marzol et al., 2017). RHs are 259 one of the main entry points in the roots for water-soluble macronutrients, such as Pi and nitrates. Pi is 260 an essential element for plant growth and development, and the availability of this macronutrient is a 261 factor limiting plant productivity. In Arabidopsis roots, low Pi in the soil triggers auxin synthesis and 262 transport, enhancing RH elongation to promote Pi uptake (Bhosale et al., 2018). Thus, auxin mediates 263 low Pi-induced promotion of RH cell expansion. Under low soil Pi, auxin synthesis is enhanced specifically 264 in the root cap (Stepanova et al., 2008) and transported (mostly by AUX1, PIN2, and PGP4) from the apex 265 to the differentiation zone, specifically leading to an increase of auxin levels in trichoblasts (Jones et al., 266 2009; Bhosale et al., 2018; Wang et al., 2020). In response to the high-auxin microenvironment, RHs pro-267 trude from the root epidermis controlled by RHD6 and RSL1(Menand et al., 2007; Pires et al., 2013). High 268 levels of auxin in trichoblasts trigger a signaling cascade mediated by TIR1-ARF19 (and possibly also 269 ARF7) which directly induces the expression of RSL4 (and likely of RSL2) and promote RH elongation (Yi et 270 al., 2010; Mangano et al., 2017; Mangano et al., 2018b; Bhosale et al., 2018). ARF7 and ARF19 also acti-271 vate other RH genes independently of RSL4 (Schoenaers et al., 2018). Interestingly, our results indicate 272 that the IncRNA APOLO participates in the response to low temperatures. APOLO is directly activated by 273 ARF7 and regulates the transcriptional activity of its neighboring gene PINOID (PID) by shaping local 3D

274 chromatin conformation (Ariel et al., 2014; Ariel et al., 2020). PID encodes a kinase responsible for accu-275 rate auxin polar transport by localizing PIN2 in the root cell membrane (Friml et al., 2004). More recently, 276 it was shown that APOLO can recognize a subset of distant genes across the Arabidopsis genome, most 277 of them being related to auxin synthesis and signalling (Ariel et al., 2020). In this work, we demonstrate 278 that a group of RH related genes is directly regulated by APOLO in response to cold, including the RH 279 master regulator RHD6. Interestingly, RSL2 and RSL4 are still activated by cold in RNAi-APOLO roots in 280 contrast to RHD6 (Figure 1C and Supplementary Figure 2A), suggesting that additional yet uncovered factors, which might include ARF TFs, may trigger RH expansion in an APOLO/RHD6-independent man-281 282 ner, and hinting at RSL4 as a key regulator. Furthermore, we found that in addition to RHD6, APOLO di-283 rectly regulates 16 RH-related genes downstream RHD6 and RSLs (Supplementary Table 1), hinting at an 284 intricate regulatory network controlling RH expansion. Collectively, our results uncover a IncRNA-285 mediated epigenetic link between environmental signals and auxin homeostasis modulating RH growth. 286 Moreover, our observations suggest that low temperatures restrict nutrient mobility and availability in 287 the culture medium, leading to the promotion of polar RH growth. Further research will be needed to 288 determine what is the limiting nutrient mediating the effect of cold on RH growth.

289 Although substantial progress has been achieved in the identification of the molecular actors 290 controlling RH development, the impact of chromatin conformation in the transcriptional regulation of 291 central TFs remains poorly understood. In this study, we have revealed a new mechanism of gene regula-292 tion in RHs by which the IncRNA APOLO integrates chromatin-associated ribonucleoprotein complexes 293 together with the TF WYRK42, participating in the transcriptional activation of RHD6 and the down-294 stream RH gene network (Figure 4E). APOLO directly regulates the chromatin 3D conformation of the 295 genomic region encompassing the RHD6 locus and stoichiometrically recruits WYRK42, previously linked 296 to Pi-starvation (Chen et al., 2009; Su et al., 2015). Low levels of APOLO fail to recruit WRKY42 to RHD6 297 promoter region, whereas high levels of APOLO likely decoy WRKY42 from target chromatin, as it was 298 shown for APOLO regulation over LHP1 binding activity (Ariel et al., 2020). Notably, low and high levels of 299 APOLO result in higher RHD6 basal transcriptional accumulation (through LHP1 activity and change in 300 chromatin status), whereas both extremes impair RHD6 activation by cold (through WRKY42 binding 301 modulation; a model is shown in Supplementary Figure 6). Our results suggest that an WRKY42-APOLO 302 hub regulates RH cell elongation through the master regulator RHD6, although the APOLO-WYRKY42 hub 303 potentially targets several additional cell wall related genes (Supplementary Table 1) at the end of the 304 pathway controlled by RHD6 and the RHD6-downstream TFs RSL2/RSL4 (Mangano et al., 2017; Mangano 305 et al., 2018a).

306 Participation of epigenetic factors in root cell identity determination strongly suggests the de-307 fault pattern for epidermal cell fate that can be overridden by environmental stimuli (Guimil and 308 Dunand, 2006). Interestingly, it was reported that the expression of GLABRA2 (GL2), a gene encoding a 309 TF repressing *RHD6* in atrichoblasts, is tightly regulated at the epigenetic level. By using 3D fluorescence 310 in situ hybridization, it was shown that alternative states of chromatin organization around the GL2 locus 311 are required to control position-dependent cell-type specification in the root epidermis (Costa and Shaw, 312 2006). Furthermore, GL2 epigenetic regulation was proposed to be responsive to salt stress (Beyrne et 313 al., 2019). In addition, a comprehensive characterization of alternative mutant lines uncovered the role 314 of PRC2 in the regulation of RH development (Ikeuchi et al., 2015). Loss-of-function mutants in different 315 PRC2 subunits develop unicellular RHs but fail to retain the differentiated state, generating a disor-316 ganized cell mass from each single RH. It was shown that the resulting RHs are able to undergo a normal 317 endoreduplication program, increasing their nuclear ploidy, although they subsequently reinitiate mitotic 318 division and successive DNA replication. It was proposed that aberrant RH development in PRC2 related 319 mutants is due to the epigenetic deregulation of key regulatory genes such as WOUND INDUCED DEDIF-320 FERENTIATION 3 (WIND3) and LEAFY COTYLEDON 2 (LEC2) (Ikeuchi et al., 2015). Here, we showed that 321 the single mutants clf (PRC2) and lhp1 (PRC1) are affected in RH growth. In addition, we found that 322 H3K27me3 deposition and LHP1 binding to the RHD6 locus is modulated by cold. Moreover, we showed 323 that in the *clf* background, H3K27me3 deposition throughout the *RHD6* locus is partially impaired, and 324 that the *clf* mutant is affected in RH elongation promoted by cold. Notably, basal transcriptional levels of 325 *RHD6* are higher in the *clf* mutant (Supplementary Figure 4C). A phenotypic characterization revealed 326 that RH density is not altered in clf or lhp1 mutants, nor in APOLO and WRK42 deregulated lines (Sup-327 plementary Figure 7), suggesting that RHD6 is not ectopically expressed in epidermic cells. Thus, RHD6 328 over-accumulation may occur in inner cell layers of the root or in RHs, although their elongation in the *clf* 329 mutant may be blocked by additional perturbations of Polycomb-associated regulation of downstream 330 genes. Altogether, our results suggest that Polycomb proteins participate in the control of RH-related 331 genes transcriptional reprogramming at low temperatures.

Notably, CLF and LHP1 were shown to interact with a subset of IncRNAs in *Arabidopsis*, modulating the activity of PRC target genes (Lucero et al., 2020). Interestingly, several IncRNAs have been linked to the control of transcription in response to cold. *FLOWERING LOCUS C* is regulated by at least three IncRNAs. First, the alternative splicing of a set of antisense transcripts, collectively named as *COOLAIR*, depends on the prolonged exposure to cold, epigenetically repressing *FLC* (Marquardt et al., 2014). The use of the *COOLAIR* proximal poly(A) site results in down-regulation of *FLC* expression in a process involv338 ing FLOWERING LOCUS D (FLD), an H3K4me2 demethylase (Marquardt et al., 2014). A second IncRNA 339 called COLD ASSISTED INTRONIC NONCODING RNA (COLDAIR) is fully encoded in the sense strand of the 340 first intron of FLC. Similar to COOLAIR, its accumulation oscillates in response to low temperatures. It was 341 proposed that COLDAIR recruits the PRC2 component CLF to target FLC for H3K27me3 deposition (Heo 342 and Sung, 2011). More recently, a third IncRNA modulating FLC transcription was identified (Kim and 343 Sung, 2017). The cold-responsive lncRNA COLDWRAP is derived from the FLC proximal promoter and it also interacts with PRC2. It was suggested that COLDWRAP functions in cooperation with the IncRNA 344 COLDAIR to retain Polycomb at the FLC promoter through the formation of a repressive intragenic chro-345 346 matin loop (Kim and Sung, 2017). Another IncRNA named SVALKA was shown to mediate the response to 347 low temperatures (Kindgren et al., 2018). Interestingly, the activation of SVALKA by cold triggers the 348 transcription of a cryptic downstream lncRNA, which overlaps the antisense locus of the C-349 repeat/dehydration-responsive element Binding Factor 1 (CBF1), involved in the early response to cold in 350 Arabidopsis. Antisense transcription causes Pol II head-to-head collision modulating transcriptional ter-351 mination of CBF1 (Kindgren et al., 2018). Here, we show that the auxin-responsive lncRNA APOLO is also 352 transcriptionally modulated by cold. The differential accumulation of 24nt siRNAs across the APOLO lo-353 cus at low temperatures indicates that this activation is related to a decrease in RdDM. Moreover, we 354 showed here that the intergenic region between PID and APOLO acting as a divergent promoter is also 355 activated at low temperatures in RHs, as revealed by the reporter gene GFP. Thus, the IncRNA APOLO 356 integrates external signals into auxin-dependent developmental outputs in Arabidopsis.

357 In the last decade, IncRNAs have emerged as regulators of gene expression at different levels, 358 ranging from epigenetics to protein modifications and stability (Ariel et al., 2015). Notably, it has been 359 shown in animals that noncoding transcripts can be recognized by TFs. In humans, it was proposed that 360 the interaction with the IncRNA SMALL NUCLEOLAR RNA HOST GENE 15 (SNHG15) stabilizes the TF Slug 361 in colon cancer cells. It was shown that SNHG15 is recognized by the zinc finger domain of Slug prevent-362 ing its ubiquitination and degradation in living cells (Jiang et al., 2018). Also, the transcriptional activity of 363 the human gene DIHYDROFOLATE REDUCTASE (DHFR) is regulated by a lncRNA encoded in its proximal 364 promoter. It was proposed that the nascent noncoding transcript forms a hybrid with its parent DNA and 365 decoys the regulatory TF IIB away from the DHFR promoter, dissociating the transcriptional pre-initiation 366 complex in guiescent cells (Martianov et al., 2007). The IncRNA P21 ASSOCIATED ncRNA DNA DAMAGE 367 ACTIVATED (PANDA) was identified in human cancer and it was activated in response to DNA damage 368 (Hung et al., 2011). PANDA is transcribed from the promoter region of the CDKN1A gene and interacts 369 with the TF NF-YA to limit the expression of pro-apoptotic genes. The activity of PANDA has been linked 370 to the progression of different tumors (Kotake et al., 2016; Shi et al., 2019). Interestingly, it was shown 371 that in addition to NF-YA, PANDA interacts with the scaffold-attachment-factor A (SAFA) as well as PRC1 372 and PRC2 to modulate cell senescence. In proliferating cells, SAFA and PANDA recruit Polycomb compo-373 nents to repress the transcription of senescence-promoting genes. Conversely, the loss of SAFA-PANDA-374 PRC interactions allows expression of the senescence program (Puvvula et al., 2014). In this work, we 375 showed that the PRC1-interacting IncRNA APOLO can also be recognized by the TF WRKY42, hinting at 376 general IncRNA-mediated mechanisms linking Polycomb complexes with the transcriptional machinery 377 across kingdoms. Furthermore, our observations indicate that the deregulation of WRKY42 affects the 378 epigenetic environment of RHD6. It was previously shown that the addition of in vitro transcribed APOLO 379 to RNAi-APOLO chromatin extracts was able to partially restore R-loop formation over APOLO target 380 genes, and that high levels of APOLO may titer LHP1 away from chromatin (Ariel et al., 2020). Here we 381 show that the relative accumulation of the IncRNA APOLO can modulate the binding activity of its part-382 ner TF to common target genes. Collectively, our results strongly support that environmentally con-383 trolled cell fate in Arabidopsis relies on a transcriptional reprogramming governed by a network of epi-384 genetic regulatory complexes, IncRNAs, TFs and effector proteins.

385

386 MATERIALS AND METHODS

387

388 Plant Material and Growth Conditions. All the Arabidopsis thaliana lines used were in the Columbia-0 389 (Col-0) background. WRKY42 over expression transgenic plants were generated through Agrobacterium 390 tumefaciens (strain EHA105)-mediated transformation (Clough and Bent, 1998). 35S:WRKY42:GFP trans-391 formant lines were selected on MS/2 medium supplemented with kanamycin (40µg/mL) and WRKY42 392 expression levels were measured by RT-qPCR (primers used are listed in Supplementary Table 3). The 393 wrky42 mutant line belongs to the SALK collection (SALK 121674C), as the one previously characterized 394 (Su et al., 2015). The rhd6-3/rsl1-1/rsl4-1; rsl2; rsl4; rsl2rsl4, 35S:RSL4 and EXP7p:RSL4 transgenic lines 395 were previously described and characterized (Yi et al., 2010; Hwang et al., 2017) Homozygous plants 396 were obtained in our laboratory and genotyped using the oligonucleotides indicated in **Supplementary** 397 Table 3. Seeds were surface sterilized and stratified at 4°C for 2d before being grown under long day 398 conditions (16h light, 140µE.m⁻².sec⁻¹/8h dark), on ½-strength Murashige and Skoog media (1/2 MS) 399 (Duchefa, Netherlands) with 0.8% plant agar (Duchefa, Netherlands).

400

401 **Cloning procedure.** The coding region of *WRKY42* (AT4G04450) excluding the STOP codon was amplified 402 by PCR, cloned into the Gateway entry vector pENTR/D-TOPO (Invitrogen) and recombined by Gateway 403 technology (LR reaction) into the pK7FWG2,0 vector containing a *p35S-GFP* cassette 404 (http://www.psb.ugent.be/gateway/index.php).

405

406 Root hair phenotype characterization. For quantitative analyses of RH phenotypes, 10 fully elongated 407 RH from the root elongation zone of 15-20 roots, were measured on the same conditions for each par-408 ticular case and grown on vertical plates with $\frac{1}{2}$ -strength Murashige and Skoog media (1/2 MS) (Duchefa, 409 Netherlands) and 0.8% plant agar (Duchefa, Netherlands) for 5 days at 22° and 3 days at 10°C. Measure-410 ments were made after 8 days. Images were captured with an Olympus SZX7 Zoom Stereo Microscope 411 (Olympus, Japan) equipped with a Q-Colors digital camera and software Q Capture Pro 7(Olympus, Ja-412 pan). RH density was determined as the number of hairs in a representative area of the root elongation 413 zone using the same setup stated above and ImageJ software. Results were expressed as the mean ± 414 standard error (SE) and in the case of RH density, values were expressed per mm². All measurements 415 indicate the average of three independent experiments, each involving 15-20 seedlings.

416

417 Confocal microscopy analysis of root hairs. Confocal laser scanning microscopy was performed using 418 Zeiss LSM5 Pascal (Zeiss, Germany) and a 40x water-immersion objective, N/A=1.2. Fluorescence was 419 analyzed by using 488 nm laser for GFP excitation (Laser Intensity: 70%, Detector Gain:550, Amplifier 420 Offset:0.1, Amplifier Gain:1), and emitted fluorescence was recorded between 490 and 525nm for GFP 421 tag. Z stacks were done with an optical slice of 1µm, and fluorescence intensity was measured in 15µm 422 ROI (Region Of Interest) at the RH tip and summed for quantification of fluorescence using ImageJ. Five 423 replicates for each of ten roots and 15 hairs per root were observed. Col-0 wild type root hairs were used 424 as a negative control, to check autofluorescence signal occurrence and no signal were detected in the 425 wavelengths range stated above.

426

RNA extraction and RT-qPCR. Total RNA was extracted using TRIZol (Invitrogen) and 2 μg were subjected to DNase treatment according to the manufacturer's protocol (Thermo Scientific). One μg of DNase-free RNA was reverse-transcribed using Maxima H Minus Reverse Transcriptase (Thermo Scientific). RT-qPCR were performed using the LightCycler 480 SYBR Green I Master Kit on a LightCycler480 apparatus (Roche) using standard protocols (40 cycles, 60°C annealing). *PP2A* (AT1G13320; primers are listed in **Supplementary Table 3**) was used as reference.

433

RNA Immunoprecipitation. RNA immunoprecipitation (RIP) assays were performed on transiently transformed *N. benthamiana* leaves as described in (Sorenson and Bailey-Serres, 2015), or in 10-day-old *A. thaliana 35S:WRKY42:GFP* seedlings as described in (Bardou et al., 2014), using anti GFP (Abcam ab290) and anti-IgG (Abcam ab6702). RIP was performed using Invitrogen Protein A Dynabeads. Precipitated RNAs were prepared using TRI Reagent (Sigma-Aldrich), treated with DNase (Fermentas) and subjected to RT-qPCR (High Capacity cDNA Reverse Transcription Kit (Thermo); primers used are listed in Supplementary Table 3). Total RNAs were processed in parallel and considered the input sample.

441

442 Chromatin Immunoprecipitation. Chromatin immunoprecipitation (ChIP) assays were performed on 10day-old WT seedlings treated or not during 24h at 10 °C, using anti H3K27me3 (Diagenode pAb-195-050), 443 444 anti LHP1 (Covalab pab0923-P) and anti-IgG (Abcam ab6702), as described in (Ariel et al., 2020). Cross-445 linked chromatin was sonicated using a water bath Bioruptor Pico (Diagenode; 30sec ON/30sec OFF 446 pulses; 10 cycles; high intensity). ChIP was performed using Invitrogen Protein A Dynabeads. Precipitated DNA was recovered using Phenol:Chloroform:Isoamilic Acid (25:24:1; Sigma) and subjected to RT-qPCR 447 448 (primers used are listed in Supplementary Table 3). Untreated sonicated chromatin was processed in 449 parallel and considered the input sample. For in vitro competition assays, APOLO was transcribed using 450 the T7 transcription kit (Promega; (Ariel et al., 2020)). After regular chromatin isolation from 10-day-old 451 35S:WRKY42:GFP seedlings, the sample was split in 4 independent tubes and diluted to 1ml in Nuclei Lysis Buffer without SDS. 0 µg, 0.1 µg, 1 µg and 10 µg of APOLO were added to each sample respectively, 452 453 and incubated in rotation at 4 °C for 3h. Then, cross-linking was performed with 1% formaldehyde for 5 454 min at 4 °C, followed by 5 min with a final concentration of 50 mM glycine. SDS was added to a final con-455 centration of 0.1% prior to sonication and the subsequent steps of a regular ChIP protocol.

456 For ChIP in transiently transformed leaves, 3-week-old A. thaliana were transformed as previously de-457 scribed (Zhang et al., 2020) In brief, Agrobacterium tumefaciens strain GV3101 carrying 35S:WRKY42:GFP 458 construct were grown for 2 days in YEB-induced medium plates at 28 °C. Agrobacterium cells were 459 scraped and resuspended in washing solution (10 mM MgCl₂, 100 μ M acetosyringone). Infiltration solu-460 tion ($\frac{1}{M}$ MS [pH = 6.0], 1% sucrose, 100 μ M acetosyringone, 0.005% [v/v, 50 μ l/l] Silwet L-77) was pre-461 pared with the previous solution, adjusting the OD600=0.5. The infiltration was carried out in all leaves > 462 0.5cm in length of between 10 and 15 plants per genotype (WT, 35S:APOLO and RNAi-APOLO lines). After 463 infiltration, plants were kept in light for 1h and then in darkness for 24h. Finally they were transferred 464 back to light. Images and samples were obtained 3 days after infiltration. For image acquisition, infiltrat-

ed leaves were imaged with a Leica TCS SP8 confocal laser scanning microscope with excitation at 488
nm (Intensity=8%) and detection at 495-530 nm for GFP and 610-670 (gain 650) nm for chlorophyll fluorescence. Images were captured using 10X and 20X lenses, and processed using Fiji software (Schindelin
et al., 2012).

469

470 Chromatin Isolation by RNA Purification followed by gPCR or mass spectrometry. A method adapted 471 from the ChIRP protocol (Chu et al., 2012) was developed to allow the identification of plant DNA associated to specific IncRNAs, as described in (Ariel et al., 2014; Ariel et al., 2020). Briefly, plants were in vivo 472 473 crosslinked and cell nuclei were purified and extracted through sonication. The resulting supernatant 474 was hybridized against biotinylated complementary oligonucleotides that tile the IncRNA of interest and 475 IncRNA-associated chromatin was isolated using magnetic streptavidin beads. One hundred pmol of 476 probes against APOLO (ODD and EVEN set of probes (Ariel et al., 2014; Ariel et al., 2020)) and the corre-477 sponding negative set against LacZ were used for independent purification. Co-purified ribonucleopro-478 tein complexes were eluted and used to purify RNA or DNA, which were later subjected to downstream 479 assays for quantification as previously described (Ariel et al., 2020).

480 The exploratory identification of plant nuclear proteins bound to APOLO was performed as described in 481 (Rigo et al., 2020). Plant samples were prepared as for ChIRP-qPCR. For protein extraction, approximately 482 250 g of 7-day-old Col-0 plants grown on solid half-strength MS medium was irradiated three times with UV using a CROSSLINKERCL-508 (Uvitec) at 0.400 J/ cm². Protein purification samples for protein extrac-483 484 tion were DNase-treated according to the manufacturer (Thermo Scientific). After addition of 1.8ml of 485 TCA-acetone (5ml 6.1N TCA + 45ml acetone + 35μ l β -mercaptoethanol), samples were incubated over-486 night at -80°C. After centrifugation at 20000rpm for 20min at 4°C, the supernatant was discarded and 487 1.8ml of acetone wash buffer (120ml acetone, 84μ l β -mercaptoethanol) was added to the samples. Then, samples were incubated 1h at -20°C and centrifuged again at 20000 rpm for 20min at 4°C. The 488 489 supernatant was discarded, and the dry pellet was used for mass spectrometry analyses.

490

491 **Chromatin Conformation Capture.** Chromosome conformation capture (3C) was performed basically as 492 previously described in (Ariel et al., 2020) starting with 2g of seedlings. Digestions were performed over-493 night at 37°C with 400U *Dpn*II (NEB). DNA was ligated by incubation at 16°C for 5h in 4 ml volume using 494 100U of T4 DNA ligase (NEB). After reverse crosslinking and Proteinase K treatment (Invitrogen), DNA 495 was recovered by Phenol:Chloroform:Isoamilic Acid (25:24:1; Sigma) extraction and ethanol precipita-

- tion. Relative interaction frequency was calculated by qPCR (primers used are listed in Supplementary
 Table 3). A region free of *Dpn*II was used to normalize the amount of DNA.
- 498

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- 506

507 Author contributions

508 MM, JMP, LL, CFF, JRM, NM, AC, MB and FA performed the experiments. JB, MB, FI, MC, JE and FA ana-509 lyzed the data. JE and FA conceived the project. FA, JE and CFF wrote the manuscript with the contribu-510 tion of all authors.

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700

701 FIGURES AND LEGENDS

702

703 Figure 1. APOLO regulates root hair elongation in response to low temperatures

- A. APOLO transcript levels measured by RT-qPCR in roots of 10-day-old plants grown at 22 °C treated or
 not for 24h at 10 °C.
- 706 **B.** The APOLO promoter region (full intergenic region between PID and APOLO) is active in root hairs at 22

°C and 10 °C, using the *GFP* reporter gene. Images are representative of the lowest and highest GFP signal
detected in the observation of several roots. The quantification of the fluorescence is shown in the right

709 panel.

C. Quantification of RH length of 8-day-old Col-0, RNAi-APOLO and 35S:APOLO plants, respectively, at 22
°C and 10 °C. Each point is the mean ±error of the length of the 10 longest RHs identified in the root
elongation zone in a single root. Representative images of each genotype are shown below the graph.
The scale bas represent 500µm. Asterisks indicate significant differences between Col-0 and the corresponding genotype at the same temperature or significant differences between temperature treatments
on the same genotype at the same temperature (one way ANOVA followed by a Tukey-Kramer test;
"*"<0.05, "**"<0.01, "***"<0.001). NS stands for no statistically significant difference.</p>

717 D. *RHD6* transcript levels measured by RT-qPCR in roots of Col-0 plants vs. two independent RNAi-APOLO
 718 lines and two independent *35S:APOLO* lines grown at 22 °C treated or not for 24h at 10 °C.

In A and D, the error bars represent the SD of 3 biological replicates. The asterisks indicate that the dif ferences are significant (t test p<0.05). Values are normalized using the constant housekeeping transcript
 PP2A.

722

723 Figure 2. APOLO directly modulates chromatin three-dimensional conformation of the RHD6 locus

A. Epigenomic landscape of the *RHD6* locus. Lane 1: H3K27me3 deposition by ChIP-Seq (Veluchamy et al., 2016). Lane 2: LHP1 deposition by ChIP-Seq (Veluchamy et al., 2016). Lane 3 to 5: *APOLO* recognition by ChIRP-Seq (Lane 3 and 4, using ODD and EVEN sets of probes against *APOLO*, respectively; Lane 5, negative control using LacZ probes)(Ariel et al., 2020). Lane 6 to 8: R-loop formation by DRIP-Seq (R-loop Atlas, root samples,(Xu et al., 2020), on Watson strand (Lane 6) and Crick strand (Lane 7). DRIP negative control after RNAseH treatment is shown in Lane 8. Gene annotation is shown at the bottom.

B. Chromatin loops identified in the *RHD6* region by *Dpn*II HiC (Liu et al., 2016). Colors of the loops are
 related to the corresponding q-values indicated below. Black boxes in A and B indicate the same genomic
 locations, where the bases of the chromatin loop correlate with R-loop formation and *APOLO* recognition

733 (compared to gene annotations). In red the arrows indicate the position of the probes used for 3C-qPCR

shown in **D**.

735 C. APOLO association to DNA of the *RHD6* locus by ChIRP-qPCR in Col-0 and RNAi-APOLO plants. The
 736 background level was determined using a set of probes against LacZ RNA.

737 **D.** Relative chromatin loop formation by 3C-qPCR deduced from **B**, in Col-0 plants vs. 35S:APOLO and

738 RNAi lines for the region upstream *RHD6*. The probes used for 3C-qPCR are indicated in red in panel **B**.

739 In C and D, the error bars represent the SD of 3 biological replicates. The asterisks indicate that the dif-

- 740 ferences are significant (t test "*"<0.05, "***"<0.001).
- 741

742 Figure 3. The transcription factor WRKY42 interacts with APOLO, directly regulates RHD6 and partici-

743 pates in the response to cold

744 **A.** APOLO-WRKY42 in vivo interaction demonstrated by RNA Immunoprecipitation (RIP)-qPCR in transi-

ently transformed tobacco leaves and stably transformed Arabidopsis plants. In tobacco, WRKY42:GFP
 and *APOLO* were co-transformed under the control of the 35S constitutive promoter. In Arabidopsis,

747 35S:WRKY42:GFP transformed plants were used to detect the interaction with endogenous APOLO. Anti-

748 IgG antibodies were used as a negative control. The RNA *PP2A* was considered as a RIP negative control

749 in Arabidopsis plants.

B. Arabidopsis eFP Browser (Waese et al., 2017) plot representing the increase of *WRKY42* transcript
levels in roots of seedlings treated for 24h at 4°C.

752 C. WRKY42 transcript levels measured by RT-qPCR in roots of plants grown at 22°C treated or not for 24h
753 at 10°C.

D. Chromatin Immunoprecipitation (ChIP)-qPCR assay revealing regulation by WRKY42 of *RHD6* by direct
 recognition of its promoter region. Probes amplifying *PP2A* were used as a negative control of the exper-

iment. Anti-IgG antibodies were used as a negative control for each pair of probes.

757 E. *RHD6* transcript levels measured by RT-qPCR in roots of Col-0 plants vs. *wrky42* mutants and
 758 35S:WRKY42:GFP lines grown at 22°C treated or not for 24h at 10°C.

F. Quantification of RH length of Col-0, *wrky42* and *35S:WRKY42:GFP* plants at 22°C and 10°C. Each point is the mean of the length of the 10 longest RHs identified in a single root. Representative images of each genotype are shown on the right. The scale bas represent 500μm. Asterisks indicate significant differences between Col-0 and the corresponding genotype at the same temperature (one way ANOVA followed by a Tukey-Kramer test; "*"<0.05, "***"<0.001).

- In A, C, D and E, the error bars represent the SD of 3 biological replicates. The asterisks indicate that the
- 765 differences are significant (t test p<0.05). NS stands for no statistically significant difference. In C and E,
- values are normalized using the constant housekeeping transcript *PP2A*.
- 767

768 Figure 4. The WRKY42-APOLO hub activates RHD6 transcription in response to low temperatures, pro-

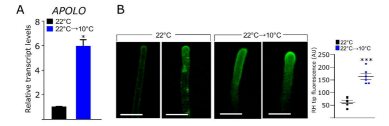
- 769 moting root hair expansion
- 770 A. H3K27me3 deposition in Col-0 vs. wrky42 mutant 10-day-old seedlings grown at 23°C over RHD6 and a
- 771 WRKY42-independent APOLO target (AZG2; (Ariel et al., 2020)). An immunoprecipitation (ChIP-qPCR) was
- performed using an anti-IgG antibody as a negative control.

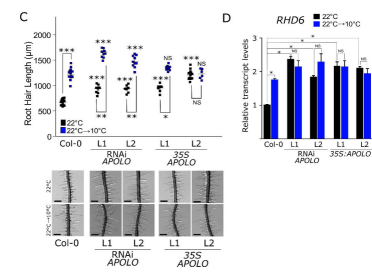
B. APOLO association to DNA of the *RHD6* locus by ChIRP-qPCR in Col-0 and *wrky42* mutant plants. The
background level was determined using a set of probes against LacZ RNA.

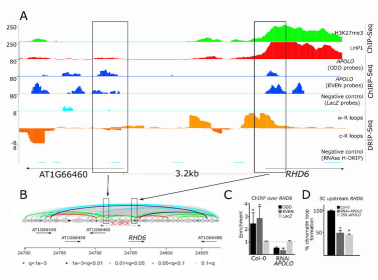
C. WRKY42 binding to *RHD6* promoter in transiently transformed leaves of Col-0, RNAi-*APOLO* and
 355:APOLO plants. WRKY42:GFP was immunoprecipitated using the anti-GFP antibody (ChIP-qPCR). An
 immunoprecipitation was performed using an anti-IgG antibody as a negative control.

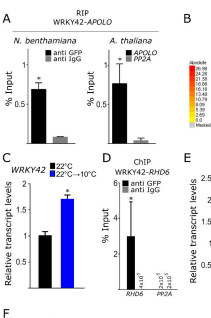
- 778 **D.** WRKY42 immunoprecipitation over *RHD6* with the addition of increasing concentrations of *in vitro* 779 transcribed *APOLO* prior to cross-linking.
- 780 E. Proposed model for the action of APOLO in RH growth under low temperatures. At 22 °C, RHD6 mod-781 erate transcription is regulated by APOLO, H3K27me3 and LHP1, maintaining a repressive chromatin loop 782 encompassing the RHD6 promoter region. In response to cold, APOLO levels increase, still recognizing the 783 RHD6 locus by sequence complementarity (R-loops). Then, LHP1 (brown balls) is decoyed away from 784 chromatin and PRC2-dependent H3K27me3 (green balls) deposition decreases in a process involving 785 WRKY42 and leading to the opening of the RHD6 promoter region. Additionally, APOLO and WRKY42 786 jointly activate RHD6 transcription. RHD6 activates the transcription of downstream TF genes RSL2 and 4 787 promoting RH cell expansion. Yet unknown factors activate RSL2 and 4 independently from APOLO and 788 RHD6 in response to cold. Furthermore, APOLO directly regulates a subset of downstream genes, likely 789 including a subset of them controlled also by WRKY42.
- 790 In A to D the error bar represents the SD of 3 biological replicates. The asterisk indicates that the differ-
- 791 ence is significant (t test "*"<0.05, "**"<0.01).

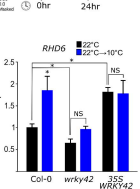
792









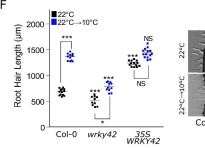


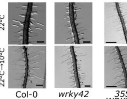
WRKY42

shoot

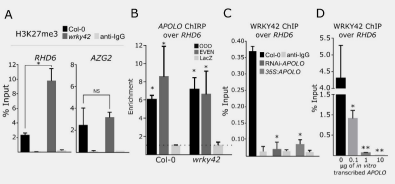
root

4°C



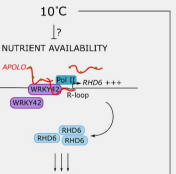


35S WRKY42



Е

22°C



?

RSL2/4

RSL2/4

RSL2/4

