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1 Autoregulation of cluster root and nodule development by white lupin CCR1 receptor-2 like kinase

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30 ABSTRACT

31 Root development is controlled by local and systemic regulatory mechanisms that optimize 32 mineral nutrient uptake and carbon allocation. The Autoregulation of Nodulation (AoN) pathway 33 defines a negative regulation of nodule development in Legumes as a way to regulate the costly 34 production of nitrogen-fixing organs. This pathway is defined as a response to symbiotic 35 interaction and has been shown to also control root formation to some extent. However, it remains unclear if root and nodule development are under coordinated genetic regulation. Here, 36 37 we identified mutants with altered root development in white lupin, constitutively producing 38 specialized lateral roots called cluster roots. We showed that the CCR1 receptor-kinase 39 negatively regulates cluster root and nodule development and targets common molecular modules 40 such as NIN/LBD16-NFYA, defining a novel pathway that we named Autoregulation of 41 Development (AoDev). AoDev defines a negative systemic pathway controlling several types of 42 root organ development, independently of symbiotic partners and nutrient availability.

43 INTRODUCTION

44 Plasticity of plant root development enables the exploration and exploitation of soil resources 45 to meet the demand for nutrients. As plant development is essentially post-embryonic, systemic 46 integration pathways play a central role in coordinating plant growth and nutrient acquisition at 47 the whole-organism level. Specifically, the demand for nitrogen (N) triggers a cascade of signaling events that orchestrate systemic responses to optimize nutrient uptake and allocation^{1,2}. 48 49 Recent studies have highlighted the involvement of root-secreted small peptides and their cognate leucine-rich repeat receptor-like kinases (LRR-RLK) as pivotal regulators of local cell-to-cell 50 communication and systemic root-to-shoot-to-root signaling pathways^{3,4}. These pathways are of 51 52 critical importance in both governing plant-wide developmental processes and maintaining 53 nutrient homeostasis, ultimately contributing to optimal plant fitness.

54 In legume plants, Autoregulation of Nodulation (AoN) has long been recognized as a pivotal negative systemic regulator of nodulation induced by rhizobial colonization⁵⁻⁷. Its 55 56 characterization has been achieved through a combination of EMS-based forward genetics, 57 natural variation approaches and grafting experiments in model species such as *Lotus japonicus*, Glycine max, and Medicago truncatula⁸⁻¹¹. Under conditions of nitrate deficiency, legume roots 58 are predisposed to colonization by N-fixing rhizobacteria, resulting in the formation of symbiotic 59 nodules. The AoN pathway encompasses two distinct mechanisms that finely tune the regulation 60 of root nodulation to balance shoot and root development, thereby preventing excessive 61 proliferation of energy-consuming nodules^{7,12}. The first mechanism involves the production of 62 CLAVATA3/EMBRYOSURROUNDING REGION-RELATED (CLE) peptides in the roots, 63 64 followed by their translocation via the xylem to the shoot. Upon reaching the shoot, CLE peptides 65 interact with LiHAR1/GmNARK/MtSUNN LRR-RLK receptors, leading to the inhibition of 66 nodulation. The second mechanism, elucidated in *M. truncatula*, and also responsive to nitrate 67 supply, involves the action of root secreted C-TERMINALLY ENCODED PEPTIDE (CEP) 68 peptides and the MtCRA2 LRR-RLK receptor in shoot tissues. This mechanism not only governs lateral root development but also, unlike the CLE-dependent route, activates nodulation¹³. Recent 69 70 investigations in M. truncatula, have identified the microRNA miR2111, and TOO MUCH LOVE (TML) Kelch-repeat F-box proteins, as components of the downward signaling cascade in 71 both routes^{14,15}. 72

White lupin (WL, *Lupinus albus*) is a hardy legume crop capable of thriving in nitrogen- and
phosphate-deficient soils, owing to the remarkable developmental plasticity of its root system^{16,17}.
In response to N deficiency, white lupin roots establish a symbiotic relationship with *Bradyrhizobium lupini*, resulting in the formation of nitrogen-fixing nodules. Additionally,

phosphate starvation triggers the development of specialized lateral roots known as "cluster 77 roots" (CRs)¹⁸. These CRs are characterized by densely clustered, short third-order lateral roots 78 termed rootlets, which serve as highly efficient organs for nutrient acquisition^{19,20}. Through the 79 secretion of protons, organic acids, and phosphatases, CRs facilitate the solubilization of 80 phosphate pools that are otherwise inaccessible to conventional root systems. In contrast, narrow-81 leaved lupin (NLL, *L. angustifolius*) is another resilient crop species that does not produce CRs²¹. 82 However, they exhibit the capacity to mobilize phosphate pools through enhanced phosphatase 83 activity²². Despite the fact that white lupin has been employed as a model for root exsudative 84 85 activities, it remains unclear how cluster roots are induced and how their development is 86 controlled at the whole plant level.

87 Here, we employed a forward genetic approach based on screening of a mutagenized white 88 lupin population for enhanced CR production in P-rich conditions, and identified the Lupinus 89 albus CONSTITUTIVE CLUSTER ROOT 1 (LalbCCR1) gene. This gene encodes an LRR-RLK, 90 exhibiting synteny with the LRR-RLKs involved in AoN, namely LiHAR1, GmNARK and 91 *MtSUNN*. CCR1 functions in a systemic pathway that restricts CR numbers, reminiscent of AoN, 92 even in the absence of rhizobial symbiosis and in the presence of nitrogen. We demonstrate that 93 by controlling both CR and nodule numbers, CCR1 is involved in a global root developmental 94 pathway, which prevents the overproduction of root organs that excessively deplete carbon 95 resources. We named this regulatory mechanism Autoregulation of Development (AoDev).

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97 RESULTS

98 Genetic identification of *LalbCCR1* as a LRR-RLK inhibiting CR development

99 In phosphate-deficient medium (-P), white lupin exhibits numerous CRs in the upper part of the 100 root system, whereas P-rich medium (+P) inhibits their development (Fig. 1a). We screened 800 101 M2 batches of an ethyl-methanesulfonate (EMS)-mutagenized WL AMIGA population for plants 102 exhibiting numerous CRs in the P-rich suppressive condition. We identified four independent 103 lines of recessive mutants (ems5, ems96, ems120, ems413). These mutants consistently develop 104 two to five times more CRs compared to wild-type plants independently of the Pi supply (Fig. 1a-105 b; Extended data Fig. 1a). Additionally, they have shorter lateral and primary roots resulting in a 106 marginally lower dry weight compared to wild-type (Fig. 1b; Extended data Fig. 1b). CRs of the 107 mutants display a higher rootlet density, as if all the potential sites of third LR initiation have 108 been unlocked (Fig. 1c-d). Physiologically, the CRs of the mutants are fully active, excreting 109 large amounts of protons, and displaying elevated phosphatase and reductase activities (Extended

110 data Fig. 1c). The pronounced overproduction of CRs observed in the mutants supports the 111 involvement of the mutated gene in an inhibitory pathway of CR development in wild-type 112 plants.

113 The mutations are recessive, segregate with the mutant phenotype, and the four independent 114 mutants fall into the same complementation class that we named *ccr1* (*constitutive cluster root 1*) 115 (Extended data Fig. 2a-b). Bulk-Segregant Analysis combined with mapping-by-sequencing 116 (BSA-seq) was performed in parallel on the four allelic *ccr1* mutant lines. This analysis unveiled 117 a linkage of causal mutations to the beginning of chromosome 3 in all four ccr1 lines (Extended 118 data Fig. 2c). Further analysis of this genomic region across the four *ccr1* alleles pinpointed the 119 presence of causal SNPs within the CONSTITUTIVE CLUSTER ROOT 1 gene (LalbCCR1, 120 Lalb Chr03g0025491, https://www.whitelupin.fr). LalbCCR1 encodes a putative protein like 121 kinase of the LRR-RLK family XI-1 (Fig. 2a) with a canonical 3D-structure calculated by 122 homology modeling using ColabFold v1.5.5: AlphaFold2 with MMseqs2 (Fig. 2b). Specifically, 123 the ccr1-2 allele carries a G1370A SNP resulting in a premature stop-codon within the LRR 124 domain; both ccr1-1 and ccr1-3 alleles contain a G2099A SNP causing a G700E substitution in 125 the kinase domain; and ccr1-4 exhibits a C2437T SNP resulting in an H813Y change within the 126 kinase domain (Fig. 2a). Notably, the G>E substitution found in *ccr1-1* and *ccr1-3* has also been identified in a hypernodulating pea line (P91, Pssym29-7)^{23,24}. This mutation is also present in a 127 LalbCCR1 paralog, that we named LalbCCR1-like (Fig. 2c). The pangenome analysis of WL 128 129 confirms the presence of this SNP in LalbCCR1-like across all WL accessions²⁵. Therefore, it is 130 reasonable to infer that LalbCCR1-like is non-functional in WL, resulting in the absence of 131 functional redundancy between LalbCCR1 and LalbCCR1-like, thereby explaining the efficiency 132 of EMS mutagenesis in isolating mutants with a highly constitutive CR phenotype. In order to 133 describe the relationship between LalbCCR1, LalbCCR1-like and other ortholog genes, a 134 phylogenetic tree was constructed, including ten of the nearest BLAST-sequence homologs of 135 LalbCCR1 in Arabidopsis thaliana, and LRR-RLKs from model legumes known to be involved 136 in AoN: LiHAR1, GmNARK, MtSUNN, and MtCRA2 (Fig. 2d). LalbCCR1 clustered with 137 GmNARK, LjHAR1, MtSUNN, as the nearest LRR-RLK/XI homolog of AtCVL1. Further 138 analysis of genomic loci revealed that the three legume LRR-RLK genes MtSUNN, LiHAR1, 139 GmNARK are syntenic to LalbCCR1 (Fig. 2e).

140

141 LalbCCR1 controls CR and nodule development through a systemic shoot-to-root142 signaling pathway.

143 MtSUNN, LiHAR1 and GmNARK are well known to be involved in AoN with corresponding mutants displaying remarkable supernodulation phenotypes^{5,11,26,27}. Based on the synteny 144 observed between LalbCCR1 and these three LRR-RLKs, we assessed the nodulation phenotype 145 146 of the independent ccr1-1 and ccr1-2 mutants after inoculation with Bradyrhizobium lupini. Both 147 mutant lines exhibited hypernodulating phenotypes (Extended data Fig. 3a-f). The efficiency of 148 nitrogen fixation was highlighted by the healthier appearance and greener foliage of nodulated 149 plants compared to the non-nodulated ones, as well as by the reddish coloration of the nodules. 150 These hypernodulation phenotypes in *ccr1* mutants provide additional validation for the accurate 151 identification of the causal gene in a species where stable transformation and subsequent 152 complementation assays are not feasible. We then used grafting experiments to test the systemic 153 nature of hypernodulation phenotypes observed in ccr1 mutants. Our results demonstrated that 154 *ccr1* shoots promote hypernodulation, characterized by clusters of nodules intertwined with short 155 rootlets, when grafted onto wild-type roots (Fig. 3a-c; Extended data Fig. 3g-j). Therefore, 156 LalbCCR1 governs nodule development from shoots through a systemic shoot-to-root pathway. 157 Analysis of LalbCCR1 expression in various WL organs revealed predominant expression in 158 petioles and hypocotyls irrespective of P conditions (Extended data Fig. 4). Thus, LalbCCR1 159 appears to be the ortholog of *MtSUNN*, *LiHAR1*, *GmNARK* for AoN.

160 Subsequently, we conducted grafting experiments to elucidate whether LalbCCR1 controls CR 161 development via a systemic mechanism akin to AoN. All grafting experiments were conducted 162 under P-rich conditions, which are favorable for the grafting process and known to inhibit CR 163 development. In this CR-suppressing condition, significant CR development occurred when ccr1 164 shoots were grafted onto wild-type rootstocks, while conversely, grafting wild-type shoots 165 significantly inhibited CR formation on ccr1 rootstocks (Fig. 3 d,e and, for ccr1-2, Extended data 166 Fig. 5a,b). These grafting outcomes unequivocally demonstrate that LalbCCR1 governs CR 167 development via a systemic shoot-to-root pathway, reminiscent of AoN. They highlight that 168 LalbCCR1 drives a common systemic shoot-to-root pathway that regulates both nodule and CR 169 development in WL.

170

171 Identification of a common negative systemic signal across *Lupinus* species

We took advantage of the inability of NLL to form cluster roots to test whether the CCR1dependent inhibitory signal is conserved across *Lupinus* species. Since WL and NLL possess differing chromosomal numbers precluding interspecific crosses, we performed interspecific grafting experiments. Grafting *ccr1* mutant shoots onto NLL rootstocks induced a remarkable 176 transformation in NLL root architecture, leading to the development of clusters of short tertiary 177 roots, that could be described as CR-like (Fig. 4 a,b and, for ccr1-2, Extended data Fig.5 c). This 178 striking modification occurred in P-rich medium, beneficial for grafting, indicating that the P-179 starvation signal is not involved in this phenotype. Grafting of *ccr1* shoots was thus sufficient to 180 initiate the formation of CR-like structures in a non-CR producing lupin species, suggesting the existence of a potent inhibitory systemic pathway in NLL. We also observed that grafting wild-181 182 type WL shoots onto NLL rootstocks, triggered the development of some tertiary short roots, 183 albeit to a lesser extent than when Lalbccr1 was used as scion. A dose-effect relationship 184 appeared in the induction of CR development of NLL rootstocks across the three grafting 185 scenarios: NLL/NLL, WL/NLL and Lalbccr1/NLL, with an increasing proportion of short 186 tertiary roots respectively (Fig. 4a,b). An inverse developmental gradient was observed when 187 Lalbccr1 was used as rootstock. Grafting wild-type WL shoots onto Lalbccr1 rootstocks, 188 inhibited CR development, albeit to a lesser extent than when NLL was used as scion (Fig. 4c,d 189 and, for ccr1-2, Extended data Fig. 5b,d). Indeed, NLL shoots profoundly and strikingly inhibited 190 the development of CRs on ccr1 roots. It should be noted that neither LangCCR1 nor LangCCR1-191 like carry the ccr1-1 SNP while LalbCCR1-like was found to harbor it (Fig. 2d). The strong 192 inhibitory effect of NLL shoots could be explained by the fact that NLL lupin likely possesses 193 two functional LRR-RLK CCR1s whereas WL has only one. These allelic variations in CCR1 194 and CCR1-like LRR-RLKs between WL and NLL may explain the observed phenotypic gradient 195 in our grafting experiments and should be explored further. This finding underscores the major 196 role played by these LRR-RLKs in regulating root architecture development in Lupinus spp through a conserved inhibitory systemic shoot-to-root pathway. 197

198

199 Lateral root developmental genes are involved in early CR development

200 White lupin CR serves as an excellent biological model for investigating tertiary roots 201 development, as it undergoes successive emergence of numerous rootlets along one lateral root 202 (LR), establishing a continuous spatial and temporal gradient of developmental stages. Taking advantage of this model, we conducted sampling of rootlet formation by collecting 1 cm-long 203 root segments every 12h, up to 132h as described before²⁸ to generate a comprehensive temporal 204 transcriptomics dataset of cluster root development. Additionally, pieces of LRs were utilized as 205 206 control (Fig. 5a; Supplementary Table 1). Principal component analysis of sample distribution 207 delineates an elliptical trajectory progressing from state T000, proximal to LR, to timepoints 208 T024-036-048 (hours), which exhibit the greatest deviation from LR samples along axis 1. These

209 three T024-036-048 samples are not distinctly differentiated, unlike the more consistent states 210 observed at T120 and T132, when rootlets had completed their growth (Fig. 5b). In order to 211 identify genes specifically expressed during the early developmental stages of rootlets, we 212 retrieved 2349 differentially regulated genes between LR and timepoints T024-036-048 213 (Absolute Log2(fold change) > 2; FDR < 0.05) (Supplementary Table 2). Subsequent Gene 214 Ontology (GO) enrichment analysis highlighted terms associated with "cell division", "cell 215 cycle", "response to auxin" and "root development", among the up-regulated genes with the most 216 prevalent counts indicating a clear alignment with our objective to target the early developmental 217 stages of rootlet formation (Fig. 5c; Supplementary Table 3). Conversely, GO enrichment 218 analysis of down-regulated genes resulted in a less interpretable outcome, with numerous GO 219 terms spanning diverse pathways (Extended data Fig. 6; Supplementary Table 3). Following 220 filtration for up-regulated transcription factors, we obtained 144 genes (Supplementary Table 4), 221 and we compiled a list of transcription factor genes up-regulated from 8 families known to be 222 involved in root development: AP2/EREB, LOB domain, ARF, GRAS, Homeodomain, NAM, 223 PLATZ and STY-LRP1 (Supplementary Table 5). We retrieved 55 WL genes, with the 224 AP2/EREB family representing nearly half with a total of 27 genes. We employed the 225 "Orthologous gene search" tool available on our WL website (https://www.whitelupin.fr), 226 combining the Orthologous Matrix Algorithm (OMA) and NCBI BLAST searches to retrieve 227 genes homologous to Arabidopsis thaliana. Within the AP2/EREB family, we discovered genes 228 associated with ethylene and cytokinin signaling, such as ERF and CRF, alongside wellcharacterized genes implicated in early LR patterning, including 4 PUCHI, 2 AIL6/7, and 3 229 230 PLT1/2 genes. Additionally, within other families, we identified genes crucial for LR 231 development, including LBD16, LBD29, WOX5, LRP1, SMB, SCR (Fig. 5d). The temporal 232 expression patterns of these genes align closely with well-established transcriptional dynamics observed during A. thaliana LR development²⁹, providing compelling evidence that rootlets 233 234 undergo analogous developmental processes as LR (Fig. 5e) and pinpointing sets of early and 235 dynamic gene expression patterns as potential targets of the AoDev pathway.

236

Both developmental and *NIN* genes are up-regulated in *ccr1* plants independently
of rhizobial infection

To elucidate the regulatory role of *LalbCCR1* in CR development and identify potential targets of AoDev, we performed a RNAseq transcriptomic analysis focusing on LR transitioning into CR, at timepoints close to T024-036-048 of prior RNAseq data, in wild-type and *ccr1-1* mutant 242 plants grown in P-deficient medium (Fig. 6a; Supplementary Table 6). This growing condition 243 was chosen to induce CR formation in wild-type plants and reveal early dynamic changes rather 244 than on/off responses. Differential gene expression analysis between wild-type and ccr1-1 245 (Absolute Log2(fold change) > 1; FDR < 0.05), revealed 1845 deregulated genes, with 798 up-246 regulated and 1047 down-regulated genes (Supplementary Table 7). GO enrichment analysis 247 revealed the activation of oxidative and phosphate stress response pathways in the *ccr1* mutant, 248 highlighting its ability to perceive and respond to phosphate deprivation (Extended data Fig. 7a; Supplementary Table 8). Nevertheless, it is noteworthy that the PHR1 genes are not up-regulated 249 250 in *ccr1* plants compared to wild-type, suggesting that the phosphate starvation response is not 251 differentially engaged in the mutant line compared to wild-type plants. Interestingly, a significant 252 up-regulation of several CLE and CEP peptides was observed, indicating a widespread 253 deregulation of the AoN signaling within the mutant line, consistent with previous results. 254 Analysis of down-regulated genes did not retrieve relevant outcomes (Extended data Fig. 7b). 255 Further filtering for up-regulated genes identified 28 transcription factors that were shared with 256 the up-regulated transcription factors in early stages of CR development (Fig. 6b, Supplementary 257 Tables 5 "Common gene in the Venn diagram"). Notably, transcription factors such as LBD16, 258 LBD29, WOX5, PUCHI, AIL6/7, and PLT1/2 were found. These results were confirmed in 259 another *ccr1* allele by measuring gene expression levels on LRs transitioning into CRs from *ccr1*-260 2 plants by RT-qPCR (Extended data Fig. 7c). Altogether, using two complementary 261 transcriptomics approaches, we revealed that genes expressed early during CR development and 262 known to be important regulators in other models for LR development (such as LBD16) are up-263 regulated in the ccr1 mutants.

264 Analyzing the deregulated genes between *ccr1-1* mutant and wild-type plants, we observed that 265 CLE and TML genes exhibited the expected regulatory patterns corresponding to their role in 266 AoN : up-regulation for CLE and down-regulation for TML (Fig. 6c). However, unexpectedly, 267 we found that two NIN genes and two NF-YA genes, specific to the symbiotic pathway, were 268 slightly but significantly up-regulated in *ccr1-1* plants compared to wild-type, while other genes 269 involved in the nodulation pathway, including SYMRK, CCamK, or NSP1, did not show 270 significant up-regulation in *ccr1-1* background (Fig. 6c; Supplementary Table 9). Additionally, 271 *PHR1* genes did not exhibit up-regulation in *ccr1-1* mutants compared to wild-type. RT-qPCR 272 expression analysis of a second *ccr1* allele (*ccr1-2*) validated these regulations for NIN, NF-YA, 273 NSP1 and PHR1 (Extended Fig. 7d). Given the expression of LalbCCR1 in both roots and shoots, 274 we investigated whether the observed regulation of NIN and NF-YA was due to local or systemic 275 events. Grafting experiments were conducted, and LRs transitioning into CRs were sampled from

276 recovering roots. Expression analysis revealed that LBD16, NIN and NF-YA genes were up-277 regulated in wild-type rootstocks when ccr1 mutants were used as scions, while TML, NSP1 or 278 PHR1 were not (Fig. 6d). These findings suggest that in wild-type plants, a shoot-to-root 279 systemic pathway involving LalbCCR1 leads to the direct or indirect inhibition of NIN gene 280 expression, which may control both LBD16 and NF-YA expression independently of nodulation 281 and PHR1 pathways. Therefore LalbCCR1 regulates genes involved in both cluster root and 282 nodule development in a shoot-to-root control, defining the Autoregulation of Development 283 pathway.

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286 DISCUSSION

287 The root system of White Lupin exhibits a robust developmental plasticity, producing nodules 288 under conditions of nitrogen deficiency in the presence of compatible rhizobia, and CRs in the 289 event of phosphate deprivation. Specifically, CR development has been poorly studied and lacks 290 genetic and molecular description, due to the absence of established model species among the 291 numerous species producing these structures (mainly trees and shrubs, mostly perennials). We took advantage of the recently published genome and pangenome in white lupin^{25,30} to conduct a 292 293 forward genetic screen and identified four allelic ccr1 mutants consistently producing CRs, 294 irrespective of phosphate availability. Independent mapping of the 4 mutants pinpointed LalbCCR1 as the causative gene responsible for inhibiting CR development in wild-type plants. 295 296 Strikingly, LalbCCR1 encodes an LRR-RLK sharing synteny with the CLV1-like LRR-RLKs well-established in legume AoN^{8,9,31}. We demonstrated that LalbCCR1 governs both AoN and 297 298 CR development through a systemic shoot-to-root pathway, indicating the involvement of the 299 same systemic pathway in both CR formation and nodule organogenesis. This observation 300 prompted us to define an Autoregulation of Development (AoDev) pathway that encompasses the 301 regulation of both structures (Fig. 6e).

302 Recent evidence indicates the incorporation of part of the LR developmental program into root nodule development in leguminous plants^{32,33}. The Nodule Inception (NIN) transcription factor 303 304 plays a pivotal role in this integration, serving as a key regulator that bridges the two programs by modulating the expression of both LBD16 and NF-YA genes^{34,35}. In our study, transcriptomic 305 306 analysis of CR development highlighted the critical role of LBD16 as a transcriptional factor essential for CR initiation, akin to its importance in LR patterning³⁶. Subsequently, we conducted 307 308 another transcriptomic analysis comparing young wild-type and ccr1-1 LR, grown in a 309 phosphate-deficient medium. Our analysis revealed the deregulation of several genes associated

310 with AoN, such as CLE, CEP and TML, as expected. Meanwhile intriguingly, we also observed 311 up-regulation of NIN and NF-YA genes in ccr1 mutants compared to wild-type, despite the 312 absence of rhizobial infection and nodule organogenesis. Canonical genes involved in rhizobia 313 infection, such as SYMRK, CCamK, or NSP1/2 did not show any up-regulation, consistent with the absence of rhizobia. On the other hand, PHR1, a transcription factor known to be induced 314 upon phosphate deficiency and to up-regulate NIN expression³⁷, did not exhibit an up-regulation 315 in *ccr1-1* compared to wild-type samples. Therefore, this P-starvation pathway cannot be 316 317 responsible for the local increase in NIN expression in the ccr1-1 roots. These two observations, 318 namely up-regulation of the NIN/LBD16-NFYA module and the absence of regulation by 319 rhizobial and nutrient (phosphate) signals in the *ccr1* mutant, advocate for a developmental level 320 of regulation. The AoDev pathway hereby targets shared molecular components to refrain from 321 organ formation.

322 Grafting experiment, backed up by qPCR analysis performed on LR rootstocks, definitively 323 established the systemic nature of the up-regulation of NIN/LBD16-NFYA module, confirming 324 the differential expression patterns observed in the RNAseq dataset. This systemic regulation is 325 significant, as leguminous plants exhibit intricate local and systemic regulatory networks involving AoN LRR-RLKs to modulate responses to nutrient availability, particularly for N and 326 327 P supply³⁸. For instance, in soybean, rhizobial infection triggers a systemic root-to-shoot-to-root inhibition of nodulation via the RIC1/2-NARK module, while the same NARK receptor can 328 329 locally suppress nodulation through the production of NIC1 peptides induced by a nitrate excess³⁹. Importantly, the systemic regulation intrinsic to AoN remained unaffected by nitrate 330 331 availability, as did the systemic control of CR development mediated by LalbCCR1 with respect 332 to phosphate availability, once again suggesting that the AoDev pathway acts at a strict 333 developmental level.

334 Both nodules and CRs serve as substantial carbon-sinks, primarily due to carbon demand of 335 rhizobia within nodules and the carbon losses resulting from organic acid secretion by CRs. This 336 developmental control is intricately regulated by shoot, in order to maintain optimal nodule or CR 337 numbers. Common regulation by the AoDev pathway suggests a mutualization of carbon sink 338 control and a potential recycling of genetic pathways. This raises several questions such as 339 whether CR systemic regulation was duplicated from AoN in white lupin, and whether non-340 legume species harbor a systemic regulation of LR development independent of nutrient 341 availability.

342 Indeed, the scenario is rendered more complex by recent findings in Medicago, elucidating the 343 involvement of an MtCLE35 peptide perceived by the SUNN shoot LRR-RLK in the nitrate-

mediated inhibition of nodulation, while MtCLE12/13-SUNN LRR-RLK orchestrates nitrate-344 independent systemic regulation of nodulation⁴⁰. The interplay between N and P in governing 345 both nodule and cluster root development is complex^{18,38}, with numerous local and systemic 346 control pathways, involving different peptide-LRR-RLK modules. However, the systemic AoDev 347 pathway appears to be the dominant regulatory force³⁹. Further studies will be needed to fully 348 understand how systemic control of root system plasticity by the AoDev pathway is orchestrated 349 350 and interacts with local components in legume and also in non-legume species. 351 352 353 REFERENCES 354 355 1. Ruffel, S. et al. Nitrogen economics of root foraging: Transitive closure of the nitrate-356 cytokinin relay and distinct systemic signaling for N supply vs. demand. Proc Natl Acad Sci 357 USA 108, 18524–18529 (2011). 358 2. Jia, Z. & von Wirén, N. Signaling pathways underlying nitrogen-dependent changes in root 359 system architecture: from model to crop species. J Exp Bot 71, 4393–4404 (2020). 360 3. Ohkubo, Y., Tanaka, M., Tabata, R., Ogawa-Ohnishi, M. & Matsubayashi, Y. Shoot-to-root 361 mobile polypeptides involved in systemic regulation of nitrogen acquisition. Nature Plants 3, 362 1-6 (2017). 363 4. Jeon, B. W. et al. Recent advances in peptide signaling during Arabidopsis root development. 364 Journal of experimental botany 72, 2889–2902 (2021). 365 5. Caetano-Anollés, G. & Gresshoff, P. M. PLANT GENETIC CONTROL OF 366 NODULATION. Annual Review of Microbiology 45, 345–382 (1991). 367 6. Oka-Kira, E. & Kawaguchi, M. Long-distance signaling to control root nodule number. 368 Current Opinion in Plant Biology 9, 496–502 (2006). 7. Li, Y. et al. Progress in the Self-Regulation System in Legume Nodule Development-AON 369 370 (Autoregulation of Nodulation). International Journal of Molecular Sciences 23, 6676 371 (2022). 372 8. Nishimura, R. et al. HAR1 mediates systemic regulation of symbiotic organ development. 373 Nature 420, 426–429 (2002). 374 9. Krusell, L. et al. Shoot control of root development and nodulation is mediated by a receptor-375 like kinase. Nature 420, 422-426 (2002). 376 10. Searle, I. R. et al. Long-Distance Signaling in Nodulation Directed by a CLAVATA1-Like 377 Receptor Kinase. Science 299, 109-112 (2003).

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- 486
- 487 METHODS
- 488

489 Plant material and cultivation

490 White lupin (Lupinus albus) cv. AMIGA (Florimond Desprez, France) and narrow-leaved lupin 491 (L. angustifolius) cv. TANJIL (CSIRO, Australia) were used in this study. Seeds were germinated 492 on a vermiculite substrate for 4 days, after which they were cultivated in 200 L hydroponic tanks 493 containing the following well-aerated nutritive solution: 400 µM Ca(NO₃), 54 µM MgSO₄, 0.24 494 μM MnSO₄, 0.1 μM ZnSO₄, 0.018 μM CuSO₄, 2.4 μM H₃BO₃, 0.03 μM Na₂MoO₄, 10 μM Fe-495 EDTA and either 200 µM K₂SO₄ for phosphate-deficient (-P) or 400 µM of KH₂PO₄ for 496 phosphate-rich (+P) experiments. Growth chambers are set to a photoperiod of 16h light / 8h 497 dark, 25 C day / 20 C night, 65% relative humidity and photon flux density of 200 μ mol m⁻² s⁻¹. 498 Grafting experiments were conducted at 7 days post-germination. The scion was prepared by 499 trimming to a V-shape and inserted into a vertical slit created in the rootstock. Post-grafting, roots 500 recovery rates ranged from 20 to 60% over the subsequent 3 to 7 days, contingent to the specific 501 scion-rootstock pairing. Phenotyping analyses were conducted on 20-day-old plants. The number 502 of LRs exhibiting clusters of rootlets, referred to as cluster roots (CR) in this study, were counted. 503 Measurements of primary and LR lengths, as well as root dry weight, were also measured. Data 504 were plotted and statistically analyzed using GraphPad Prism software 10.2. In order to visualize 505 physiological activities of *ccr1-1* root systems, roots were spread on 0.8% agar plates containing 506 either, 0.005% (m/v) bromocresol purple buffered in Tris-HCl pH 6 for testing proton excretion, 507 or 0.013% (m/v) 5-Bromo-4-chloro-3-indolyl phosphate buffered in sodium acetate pH 5 for 508 testing phosphatase activity, or 330µM bathophenanthroline disulfonic acide disodium salt, 100 509 µM FeNaEDTA for testing ferric reductase activity.

510

511 EMS population and genetic screen

512 A large-scale mutagenesis was conducted on AMIGA seeds using 0.4% ethyl methanesulfonate

513 (EMS) for 3 hours and deactivation with sodium thiosulfate 2.5% for 5 minutes. M1 seedlings

514 were subsequently cultivated in the Cerience experimental fields (Poitiers, France) and the pods 515 from each individual M1 plant were harvested, resulting in an EMS-mutagenized population of 5000 M2 batches. Finally, 36 seeds from each of 800 M2 batches were screened in P-rich 516 517 medium to identify plants exhibiting constitutive cluster roots. Plants with the same constitutive 518 cluster root phenotype were found in 4 independent batches and amplified. They were crossed for 519 allelic test and back-crossed with AMIGA for the mapping by sequencing strategy. Pools of 50 to 520 90 F2 plants with a mutant phenotype (homozygous) and similarly sized pools of plants with a 521 wild-type phenotype (WT batch) were harvested for DNA extraction and sequencing by Illumina 522 HiSeq at Get-PlaGe core facility (INRAe, Toulouse, France). A mean coverage of 50X to 100X was obtained across samples. Cutadapt v1.15⁴¹ has been used to remove IlluminaTruseq adapter 523 524 from the sequencing data and to remove bases with a quality score lower than 20, in both 5' and 525 3' end of the reads. Pairs of reads containing one read with a length lower than 35 have been discarded. We used BWA-MEM v0.7.17⁴² to map reads to the white lupin reference genome. 526 Picard MarkDuplicates v2.20.1 (https://github.com/broadinstitute/picard) has been used to detect 527 and remove PCR and Optical duplicates. We then used GATK HaplotypeCaller v4.1.4.1⁴³ tool to 528 call variants and snpEff 4.3t⁴⁴ to annotate them. The duplicate free mapped reads have been used 529 as input for the mutmap pipeline $v2.1.2^{45}$. 530

531

532 Microscopy

533 Root segments transitioning to CR were fixed with 4% paraformaldehyde for 120 min at room 534 temperature under vacuum treatment and then washed twice for 2 min in 1X PBS before being 535 embedded in 3% (w/v) agarose resin in PBS. Longitudinal root sections of 100 µm were cut with 536 a vibrating microtome (Microcut H1200, BioRad). The sections were stained with 2 µg/mL 4,6 537 diamidino-2-phenylindole (DAPI). Fluorescence was observed using a ZEISS Axio Observer 538 microscope, with a plan-apochromatic 20X/0.8 objective and the following filters: BP 325-390 nm for excitation and BP 445/50 for emission. Mosaic pictures were taken using the Apotome 539 540 module. Images were captured with OrcaFlash (Hamamastu) controlled with the ZEISS Zen blue 541 Software. In nodulation experiments, nodules were observed in dark field with the OLYMPUS 542 SZX16 stereo microscope and images were taken with a DP72 camera.

543

544 Structural and Phylogenetic analysis

AlphaFold structure prediction was performed using ColabFold v1.5.5: AlphaFold2 with MMseqs2^{46,47}, providing the amino acid sequence of LalbCCR1 (LalbChr03g0025491). PyMol v2.5.4 was used to visualize and modify the protein structure to make the transmembrane domain 548 apparent, and to indicate the positions of amino acid substitutions within the structure. Closest 549 LalbCCR1 homologs from white lupin, narrow-leaved lupin, Glycine max, Lotus japonicus, Medicago truncatula and Arabidopsis thaliana were retrieved using NCBI blastp tool. Alignment 550 of the kinase domain portion was conducted using MUSCLE⁴⁸ alignment software on 551 NGPhylogeny website with default settings, and the output was refined with JalView 2.11.3.3⁴⁹ 552 Phylogenetic analysis was performed using the PhyML/OneClick workflow with default settings 553 on NGPhylogeny⁵⁰. The resultant phylogenetic tree was generated using iTOl v6⁵¹. Synteny 554 analysis utilized genome assemblies from *M. truncatula* A17 r5.0⁵², *G. max* Williams 82 v4.0⁵³, 555 L. japonicus MG20 v 3.0^{54} and L. albus v 1.0^{30} . It was carried out using Easyfig 2.2.5⁵⁵ with blastn 556 557 and a minimum identity value for the blast at 0.7. The output of Easyfig was subsequently edited 558 with Inkscape 1.2.1.

559

560 Nodulation assays

561 For nodulation experiments plants were grown in Magenta GA-7 pot filled with leached and 562 sterilized zeolite substrate (Siliz 14, Somez, France) supplied with a nutrient solution 563 corresponding to the previously described P-rich medium but without nitrogen. Seeds were 564 sterilized with calcium hypochlorite, germinated in Petri dishes and then transferred into Magenta 565 pots. The *Bradyrhizobium lupini* MIAE428 strain (previously named LL13)⁵⁶ was used. Inoculum was produced by cultivating the strain in modified yeast mannitol (YM) medium 566 567 (mannitol 10 g/L, yeast extract 1 g/L, K₂HPO₄ 0.5 g/L, NaCl 50 mg/L magnesium sulfate 7H₂O 100 mg/L, calcium chloride 40 mg/L, glutamic acid 0.43 g/L, FeCl₃ 4 mg/L) supplemented with 568 569 nalidixic acid 20 µg/L, in the dark for 4 days at 28 °C. One mL inoculum was applied one week 570 after the seedlings were transferred to the pots or one week after grafting. Nodule numbers per 571 plant were assessed and the leaf chlorophyll content was indirectly estimated using a Chlorophyll 572 meter SPAD (Konica-Minolta) on the third youngest leaf. Data were plotted and statistically 573 analyzed using GraphPad Prism software 10.2.

574

575 Gene expression analysis

576 Developmental temporal transcriptome. The sampling began (T0) on eight-day-old plants 577 grown in P-deficient conditions. A total of eight 1 cm-long transitioning CR segments from four 578 independently grown plants was sampled, at a distance of 1 cm from the primary root, in the 579 upper part of the root system where LRs are transitioning to CR, every 12 h for 5 days, covering 580 the entire rootlet developmental process (T0 to T132). As a control, 1 cm-long lateral root 581 segments not transitioning to CR were collected. Four biological replications were produced for 582 each experiment. Total RNA was extracted from all frozen samples using the Direct-zol RNA 583 MiniPrep kit (Zymo Research) according to the manufacturer's recommendations. A total of 52 584 independent root RNA-seq libraries were constructed and sequenced at Get-PlaGe core facility 585 (INRAe, Toulouse, France). The Illumina TruSeq Stranded mRNA Sample Preparation Kit 586 (Illumina Inc.) was used according to the manufacturer's protocol. Paired-end sequencing was 587 performed, generating 2 x 150 bp reads using TruSeq SBS kit v3 sequencing chemistry on an Illumina NovaSeq instrument. Raw reads were cleaned using Cutadapt v1.15⁴¹, by removing 588 589 bases with a quality score lower than 30, in both 5' and 3' end of the reads, as well as TruSeq 590 Illumina adapters. Pairs of reads containing one read with a length lower than 35 have been 591 discarded. The quality-checked RNA-seq reads were mapped on the white lupin genome 592 reference using Hisat2 v2.1.0, with the following parameters "--rna-strandness RF --dta". 593 Transcripts were assembled and quantified using Stringtie v1.3.4d with the options "--rf -e -B -u -594 M 1".

ccr1-1 mutant transcriptome. All LRs of four ten-day-old plants grown in P-deficient conditions were harvested from the upper part of the root system, corresponding to the zone where LRs are transitioning to CR. Eight independent RNA-seq libraries were constructed and processed as described for developmental temporal transcriptome.

599 Normalization, differential expression and gene ontology enrichment analysis were performed using the DIANEbeta R package⁵⁷ (https://shinyapps.southgreen.fr/app/dianebeta) v1.1.0.1. The 600 TCC R package with the "tmm" normalization method was used, with prior removal of 601 602 differentially expressed genes. For each analysis, Log2(fold change) and False Discovery Rate adjusted p-value (FDR) were provided in the text and figure legends. SRPlot online⁵⁸ was used 603 604 for generating the PCA, heatmap, and the GO plots, and GraphPad Prism software 10.2 for the 605 statistical analysis and kinetic expression data plotting. The GO terms used for enrichment are 606 available for download at https://www.whitelupin.fr/download.html.

- 607
- 608 RT-qPCR experiments

For *LalbCCR1* expression in different organs, 11-day-old plants grown under either P-deficient or P-rich conditions were sampled. Samples included for lateral roots (LR), root apical meristem (RAM), shoot apical meristem (SAM), leaf, petiole and hypocotyl. Cluster root (CR) samples were collected exclusively from plants grown on P-deficient conditions. CRs were collected from the upper part of the root system, while LRs were collected below. The apices of LR and CR were removed. Petioles and leaves were collected from the second leaf. Each sample contained tissues from 3 individual plants and 3 biological replicates were collected for each plant part. For 616 experiments confirming transcriptomic data in another allele besides ccr1-1, samples were 617 collected from the root system of ccr1-2 mutant plants following the same protocol as for the 618 transcriptomic study. For the grafted plants, this protocol was applied to roots recovering after the 619 grafting operation. At least five biological replicates were collected and analyzed in each 620 experiment. Total RNA was extracted using the Direct-zol RNA MiniPrep kit (Zymo Research) according to the manufacturer's recommendations. RNA concentration was measured on a 621 622 NanoDrop (ND1000) spectrophotometer. Poly(dT) cDNAs were synthetized from 2 µg total 623 RNA using the RevertAid First Strand cDNA Synthesis (ThermoFisher). Gene expression was 624 measured by quantitative Real Time-Polymerase Chain Reaction (gRT-PCR) (LightCycler 480, 625 Roche Diagnostics) using the SYBR Premix Ex Taq (Tli RNaseH, Takara, Clontech). Expression 626 levels were normalized to a putative initiation factor LalbEIF-4 (Lalb Chr07g0195211) or to a 627 LalbPolvubiquitin (Lalb Chr06g0164891). Two technical replicates were performed for all gRT-628 PCR experiments. Specific primer pairs are described on the Supplementary Table 10. Relative 629 gene expression levels were calculated according to the $\Delta\Delta$ Ct method, using LR (for organ 630 expression), or WT and WT/WT, samples for ccr1-2 and grafted plants respectively. 631

632 Data availability

633 FASTQ raw sequence files are available at NCBI under the Bioproject number PRJNA1124865

- 634 for the temporal RNAseq (Sequence Read Archive accession numbers SAMN41865670-82) and
- 635 number PRJNA1125199 for the ccr1-1 RNAseq (Sequence Read Archive accession numbers
- 636 SRR29446565-79).
- 637 *Reviewer links for temporal dataset:*
- 638 <u>https://dataview.ncbi.nlm.nih.gov/object/PRJNA1124865?reviewer=851s8ptss62fl7h178k2j5f8bu</u>
- 639 and for ccr1-1 dataset:
- 640 *https://dataview.ncbi.nlm.nih.gov/object/PRJNA1125199?reviewer=m11hh43j385of3s7hgrnsd7uc*
- 641

p

642

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652

653 Author contributions

FD and FG performed the initial genetic screen and subsequent genetic analysis. FD, AB, LM, VF, CMC, IV performed phenotypic analysis. LM, CMC, FD and AB performed grafting experiments and expression analysis. BH generated the temporal RNAseq dataset and FD the *ccr1* dataset. EIA performed the *CCR1* expression analysis. AS, HP and LM performed bioinformatic analyses. LM analyzed the data. LM and FD generated the figures. LM and BP conceived the project, obtained funding and wrote the article.

660

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667

668 **Competing interests**

669 The authors declare they have no competing interests.

670 FIGURE LEGENDS

671

672 Figure 1. Phenotypes of the four allelic *ccr1* mutants

- a) Representative images of the upper part of the root system from 20-day-old wild-type (WT)
- and the four *ccr1* mutant plants grown on either phosphate-rich medium (+P) or phosphate-
- 675 deficient medium (-P). Scale bar = 1 cm.

b) Quantitative analysis of various root traits in wild-type (WT) and *ccr1* mutant plants, including

- 677 CR abundance within the upper 10 cm of the root systems, maximum lateral root length, root
- 678 system dry weight. Statistical analysis was performed using two-way ANOVA with Tuckey 679 correction, p < 0.05.
- 680 c) Apotome imaging of a CR section from the *ems5* mutant compared to wild-type (WT) plants.
- 681 DAPI staining revealed rootlet primordia, identifiable by their small nuclei. Scale bar = 100 682 μ m.
- d) Density distribution of rootlet along 1 cm of cluster root in the *ems5* mutant. Statistical
- analysis was performed using two-way ANOVA with Tuckey correction, n = 8, p < 0.05
- 685

Figure 2. LalbCCR1 is a LRR-RLK syntenic with LRR-RLKs involved in Autoregulation of Nodulation.

a) *LalbCCR1* gene (*LalbChr03g0025491*) and LalbCCR1 protein structures. The *LalbCCR1* gene
lacks introns. EMS-induced mutations are indicated by a triangle with the corresponding
mutant names. The predicted LalbCCR1 protein contains a LRR domain, a transmembrane
domain and a kinase domain. The specific amino-acid substitutions resulting from EMSinduced SNPs are given.

b) The 3D-structural model of LalbCCR1 generated by AlphaFold2, highlighting the mutated
amino-acid positions within the kinase domain of *ccr1-1* and *ccr1-4* mutants.

c) Phylogenetic tree of LalbCCR1-related proteins across narrow-leaved lupin (*Lupinus angustifolius*), *Lotus japonicus*, *Glycine max*, *Medicago truncatula* and *Arabidopsis thaliana*.

- d) Allelic variations in the kinase domain segment containing the *ccr1-1/3* G>A mutation across
- 698 closest-homologs of LalbCCR1 in L. albus, L. angustifolius, Lotus japonicus, Glycine max,
- 699 *Medicago truncatula* and *Arabidopsis thaliana*. Notably, the *LalbCCR1* paralog, *LalbCCR1*-
- 700 *like* exhibits the same punctual mutation observed in *ccr1-1/3* mutants, leading to a non-
- functional protein in *L. albus*.
- e) Syntenic relationships among *LalbCCR1*, *GlmNARK*, *LjHAR1*, *MtSUNN* loci, indicating
- 703 genomic organization across species at this locus.

705	Figure 3. <i>LalbCCR1</i> controls CR nodulation and development via a systemic signaling
706	pathway.
707	The four graft combinations involving wild-type (WT) and Lalbccr1-1 mutant plants were
708	cultivated on either P-rich or N-deprivated medium and inoculation with Bradyrhizobia lupini
709	was performed in Magenta boxes, or in tanks containing N- and P-rich conditions for phenotypic
710	evaluation of the root systems.
711	a) Representative images of nodulated rootstocks of the four 3-week-old graft combinations
712	inoculated with Bradyrhizobia lupini. The legends of the images indicate scion/rootstock.
713	Scale bar = 0.7 cm.
714	b) Quantification of nodule numbers per plant on the rootstocks of the four 3-week-old graft
715	combinations. The legends of the images indicate scion/rootstock. Error bars represent mean \pm
716	SD, with statistical analysis performed using Kruskal-Wallis test, $n = 8$, *adjusted p-value =
717	0.0305, ***adjusted p-value = 0.0003 .
718	c) Magnification of the nodules formed on the rootstocks of grafted plants, revealing compact
719	clusters of nodules on ccr1-1/WT and ccr1-1/ ccr1-1 plants. The legends of the images
720	indicate scion/rootstock. Scale bar = 0.2 cm.
721	d) Root systems of the four 3-week-old graft combinations. The legends of the images indicate
722	scion/rootstock. Heterografted plants, ccr1-1/WT and WT/ccr1-1, demonstrate that the
723	constitutive CR phenotype is primarily influenced by the ccr1-1 mutation in the scion rather
724	than in the rootstock. Scale bar = 1 cm .
725	e) Quantitative assessment of CR abundance within the upper 10 cm of the rootstocks. Error bars
726	represent mean \pm SD, and statistical analysis was performed using Kruskal-Wallis test, n = 10,
727	***adjusted p-value = 0.0006 , ****adjusted p-value< 0.0001 .
728	
729	Figure 4. Phenotypes of interspecific grafts between L. albus, L. angustifolius, and the ccr1-1
730	mutant.
731	a) Representative images illustrating rootstocks phenotypes of grafted plants using NLL (L.
732	angustifolius) as rootstock and NLL, WL (L. albus) or the mutant Lalbccr1-1 as scions. The
733	Lalbccr1-1 scion triggered the emergence of numerous tertiary roots on NLL rootstock.
734	Legends of the images indicate the scion/rootstock combination. Scale bar = 1 cm.
735	b) Quantitative evaluation of the abundance of LR with more than 15 tertiary rootlets on NLL
736	rootstocks, with statistical analysis performed using Kruskal-Wallis test; n = 11 to 24; adjusted
737	p-values are indicated: *p = 0.0110, ****p<0.0001.

738 c) Representative images illustrating rootstocks phenotypes of grafted plants using the Lalbccr1-1 739 mutant as rootstock and NLL (L. angustifolius), WL (L. albus) or Lalbccr1-1 as scions. The 740 NLL scion markedly suppressed the formation of CRs on Lalbccr1-1 rootstock, whereas WL 741 did so to a lesser extent. Notably, *LalbCCR1-like* gene carried the same mutation as in the 742 Lalbccr1-1 mutant, whereas LangCCR1-like does not. Legends of the images indicate the 743 scion/rootstock combination. Scale bar = 1 cm. 744 d) Ouantitative evaluation of the abundance of LR with more than 15 tertiary rootlets on 745 *Lalbccr1-1* rootstocks, with statistical analysis performed using Kruskal-Wallis test, n = 10 to 746 11, adjusted p-values are indicated : *p = 0.013, ****p < 0.0001. 747 748 Figure 5. Transcriptional expression analysis during CR formation in white lupin. 749 a) Illustrations depicting the developmental stages of 1-cm long segments of LR sampled for 750 developmental temporal RNAseq transcriptomic analysis. Samples were collected every 12h 751 over a period of 132h. Segments of LR without developing rootlets served as control. Four 752 biological replicates were performed, each containing eight 1-cm-long LR segments coming 753 from four different plants. Scale bar = 2 mm. 754 b) Scatter plot presenting the distribution of the RNAseq dataset with the four replicates for each 755 timepoint and LR control samples in the two principal components, explaining 46.79% of the 756 total variance. 757 c) GO enrichment analysis of early up-regulated genes between LR and timepoints T024-036-048 758 (Absolute Log2(fold change) > 2, FDR < 0.05). GO terms presenting counts > 10 are 759 displayed. They are associated with cell division, response to hormones and root development. 760 Normalization, differential expression and GO term analysis were performed using DIANE R package⁵⁷. 761 762 d) Clustered heatmap displaying the normalized Z-scores of gene expression for up-regulated 763 transcription factors from 8 families known to be involved in root development (AP2/EREB, 764 LOB domain, ARF, GRAS, Homeodomain, NAM, PLATZ and STY-LRP1), comparing LR 765 with timepoints T024-036-048 (Absolute Log2(fold change) > 2, adjust p-value (FDR) < 766 0.05). Transcription factors known to participate in A. thaliana LR patterning are highlighted. 767 e) Kinetic profiles illustrating changes over the 132h early developmental stages of six major LR 768 patterning transcription factors involved in CR development. 769 770 Figure 6. Transcriptional expression analysis of ccr1-1 mutant during CR development and 771 the proposed model for the AoDev pathway.

- a) Illustrations depicting the developmental stages of roots used for sampling the LRs from WT
 and *ccr1-1* plants for the RNAseq. LRs longer than 3 cm from three plants were pooled, and
 four biological replicates were collected. Scale bar = 1 cm.
- b) Venn diagram illustrating the overlap of transcription factors-encoding genes up-regulated at
 timepoints T024-036-048 (hours) versus LR samples in the temporal developmental RNAseq
- dataset (minimal gene count sum across conditions = 1200, Absolute Log2(fold change) > 2,
- FDR < 0.05) compared to those up-regulated in the *ccr1-1* CR versus wild-type samples
- (minimal gene count sum across conditions = 80, Absolute Log2(fold change) > 1, FDR <
- 780 0.05). Normalization and differential expression analysis were performed using DIANE R
- package⁵⁷. Among the 28 shared genes, 13 emblematic transcription factors involved in
- 782 LR/CR initiation are listed.
- c) Log2(fold change) of selected key genes in the *ccr1-1* mutants relative to wild-type plants,
- focusing on the AoN, the nodulation pathway, and the *LBD16* and *PHR1* genes. Expected
- regulations were observed for AoN genes with up-regulation of *CLE* genes and down-
- regulation of *TML* genes. Nodulation genes were not significantly deregulated in *ccr1-1*
- 787 mutant compared to wild-type, except for NIN and NF-YA. PHR1 genes were not significantly
- deregulated in *ccr1-1* mutant compared to wild-type. Fold changes and adjusted p-values
- (FDR) were calculated using DIANE R package⁵⁷. In the graph: ns p-adjust>0.05, *p-
- 790 adjust<0.05, **p-adjust<0.01, ***p-adjust<0.001, ****p-adjust<0.0001.
- d) RT-qPCR analysis of the expression levels of selected genes in wild-type rootstocks of grafted
 plants using either wild-type (WT/WT) or *ccr1-1* mutants (WT/*ccr1-1*) as scions. The grafting
- experiment revealed that *LBD16*, *TML*, *NIN*, *NF-YA* genes are systemically regulated by
- *LalbCCR1* via a shoot-to-root systemic pathway. Additionally, it confirmed that the *NSP1*
- nodulation gene or *PHR1* genes are not significantly deregulated when comparing WT and
- 796 ccr1-1 in the growing conditions used. Error bars represent mean \pm SE, and statistical analysis
- 797 was performed using Two-tailed Mann Whitney tests: ns p-value>0/05, *p-value <0.05, **p-
- 798 value <0.0, ***p-value <0.00, ****p-value<0.0001.
- e) Proposed model for the AoDev pathway, depicting a systemic root-to-shoot-to root signaling
- 800 mechanism involving the LRR-RLK CCR1, which represses the *NIN/LBD16/NF-YA* module,
- 801 leading to the inhibition of both CR and nodule development.





M. truncatula Chr04

34.924 - 34.720 Mb

Figure 3







Figure 6





AoN

p<0.0001

p=0.927

0.0

WINT

cert-flowr

Lalb_Chr13g0303751 (AP2/ERF domain)	PUCHI
Lalb_Chr07g0177601 (AP2/ERF domain)	PUCHI
Lalb_Chr18g0055601 (AP2/ERF domain)	PUCHI
Lalb_Chr01g0003331 (AP2/ERF domain)	PLT1/2
Lalb_Chr19g0138171 (AP2/ERF domain)	PLT1/2
Lalb_Chr16g0381611 (AP2/ERF domain)	AIL6/7
Lalb_Chr12g0199451 (AP2/ERF domain)	AIL5
Lalb_Chr06g0162491 (LOB domain)	LBD16
Lalb_Chr01g0019701 (LOB domain)	LBD29
Lalb_Chr18g0052701 (HOMEOBOX)	WOX5
Lalb_Chr09g0322101 (SHI RELATED seq.)	LRP-related
Lalb_Chr16g0377691 (SHI RELATED seq.)	LRP-related
Lalb_Chr14g0369671 (SHI RELATED seq.)	LRP-related

С

Lalb_Chr01g0006721 (CLE)-Lalb_Chr01g0022391 (CLE) Lalb_Chr03g0031061 (CLE) Lalb_Chr19g0133921 (CLE)-Lalb_Chr01g0006711 (CLE)-Lalb_Chr22g0360431 (TML1) Lalb_Chr21g0317471 (TML2a) Lalb_Chr09g0325781 (LaSYMRK) Lalb_Chr19g0132601 (LaCCamK) Lalb_Chr25g0281441 (NSP1) Lalb_Chr23g0270931 (NSP2a) Lalb_Chr23g0270961 (NSP2b) Lalb_Chr10g0107661 (NINa) Lalb_Chr04g0246951 (NINb) Lalb_Chr25g0287931 (NF-YAb) Lalb_Chr01g0017821 (NF-YAa) Lalb_Chr06g0162491 (LaLBD16a) Lalb_Chr02g0152691 (PHR1a) Lalb_Chr06g0170661 (PHR1b) Lalb_Chr20g0113691 (PHR1c)



0.0 WINT cert-1 INT Relative expression

2

WINT

cert-1 INT



