1 miR167-ARF8, an auxin-responsive module involved in the formation of

2 root-knot nematode-induced galls in tomato

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15

16 Abstract

- Root-knot nematodes (RKN) from genus *Meloidogyne* induce the dedifferentiation of
 root vascular cells into giant multinucleate feeding cells. These feeding cells result
 from an extensive reprogramming of gene expression in targeted root cells, as shown
 by transcriptomic analyses of galls or giant cells from various plant species.
- Small non-coding RNAs, and messenger RNAs from tomato (*Solanum lycopersicum*)
 galls and uninfected roots were sequenced. *De novo* microRNA prediction in the
 tomato genome identified microRNAs expressed in galls and uninfected roots.
 Statistical analyses identified 174 miRNA genes differentially expressed in galls at 7
 and/or 14 days post infection (dpi).
- Integrative analyses combining small non-coding RNA and transcriptome datasets
 with the specific sequencing of cleaved transcripts identified miRNA targets in tomato
 galls. Functional analyses of promoter-GUS fusions and CRISPR-Cas9 mutants
 highlighted the role of the miR167-regulated transcription factor AUXIN RESPONSE
 FACTOR 8 (ARF8) in giant cell formation.
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32 Key words: root-knot nematodes, galls, microRNAs, tomato, auxin, ARF8

33 Introduction

Root-knot nematodes (RKNs) are major crop pests causing massive yield losses, estimated at 34 35 millions of Euros annually, worldwide (Blok et al., 2008; Abad & Williamson, 2010). These 36 microscopic worms of genus *Meloidogyne* have a wide host spectrum encompassing more 37 than 5,000 plant species, and a wide geographic distribution. After infecting the root, these 38 obligatory parasites induce the *de novo* formation of a specialized feeding site that is crucial 39 for nematode survival. The second-stage RKN juveniles (J2) penetrate the roots and migrate 40 within them; they then inject a cocktail of molecules into five to seven root parenchyma cells 41 (Favery et al., 2016). In response to RKN signals, targeted root parenchyma cells 42 dedifferentiate into giant multinucleate hypermetabolic feeding cells. These "giant cells" form 43 the feeding site supplying the nematode with the nutrients it requires for its development 44 (Favery et al., 2020). Dedifferentiation into giant cells involves an initial phase of successive 45 mitoses without cytokinesis, followed a second phase of endoreduplication (de Almeida 46 Engler & Gheysen, 2013). During feeding cell formation, the neighboring cells begin to 47 divide. This whole process results in a swelling of the root to form a gall, the characteristic 48 symptom of RKN infection. Feeding site formation involves several biological processes, 49 including the cell cycle (de Almeida-Engler et al., 2011), metabolic reprogramming (Marella 50 et al., 2013), cytoskeleton organization (Caillaud et al., 2008), and auxin signaling (Gheysen 51 & Mitchum, 2019). Auxin (indole-3 acetic acid, IAA), is a major plant hormone that plays a 52 key role in root development by regulating cell division and the establishment/maintenance of 53 root primordia (De Smet et al., 2007; Weijers & Wagner, 2016). The formation of RKN-54 induced feeding sites has been shown to involve a peak in auxin levels (Karczmarek et al., 55 2004; Absmanner et al., 2013) and gall transcriptome analyses have shown that auxin biosynthesis and auxin-responsive genes are upregulated in A. thaliana early galls, whereas 56 57 the genes encoding repressors of auxin response genes are repressed (Barcala et al., 2010). 58 Multiple transcriptome analyses have been performed on RKN-infected roots, galls or

specifically on giant feeding cells, from various plant species, including tomato, initially by microarrays, and more recently by RNA sequencing (Bar-Or *et al.*, 2005; Portillo *et al.*, 2013; Shukla *et al.*, 2018). Four time points in feeding site formation have frequently been investigated in transcriptome analyses: the early phase of feeding site formation at 3 days post infection (dpi), 7 dpi, a time point corresponding to multiple mitoses without cytokinesis, 14 dpi, corresponding to the endoreduplication phase and cell expansion, and, finally, 21 dpi when the feeding cells are mature and fully functional. All these analyses revealed a massive bioRxiv preprint doi: https://doi.org/10.1101/2022.07.29.501986; this version posted August 1, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

reprogramming of plant gene expression in response to nematode infection, with about 10%
of protein-coding genes displaying changes in expression levels in response to RKN infection
(Cabrera et al., 2014). However, it remains unclear how this reprogramming occurs and how
these genes are regulated.

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71 Small non-coding microRNAs may act as the master regulators of this reprogramming of 72 gene expression (Jaubert-Possamai et al., 2019). MicroRNAs are major repressors of gene 73 expression in eukaryotes. Within the plant genome, microRNAs are encoded by MIR genes, 74 often organized into multigene families and transcribed as a single-stranded RNA precursor 75 that folds into a typical hairpin structure. This hairpin precursor is processed to generate a 76 microRNA duplex consisting of two complementary strands of 20-22 nucleotides. One of the 77 two strands is then loaded into the ARGONAUTE-1 protein and guides the RNA silencing 78 complex (RISC) to target messenger RNAs through miRNA/mRNA sequence 79 complementarity. The targeting of an mRNA by a microRNA induces its degradation or the 80 inhibition of its translation, depending on the mRNA/miRNA sequence complementarity. 81 Several recent studies have identified microRNAs expressed in galls (Jaubert-Possamai et al., 82 2019) induced by RKN in Arabidopsis thaliana (Cabrera et al., 2016; Medina et al., 2017), in 83 tomato (Solanum lycopersicum; Zhao et al., 2015; Kaur et al., 2017), in cotton (Gossypium 84 hirsutum; Pan et al., 2019; Cai et al., 2021) or in rice (Oryza sativa; Verstraeten et al., 2021). 85 However, the roles of only four microRNAs have been validated by functional analyses: the 86 miR390/tasiRNA/ARF3 module (Cabrera et al., 2016), the miR159/MYB33 pair (Medina et 87 al., 2017), the miR172/TOE1/FT module (Díaz-Manzano et al., 2018) in Arabidopsis and the 88 miR319/TCP4 pair in tomato (Zhao et al., 2015).

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We investigated the gene regulation network involved in the plant response to RKN through a combination of transcriptome, microRNome and degradome sequencing in uninfected roots of tomato and tomato galls induced by the RKN *M. incognita* at two key time points in gall development: 7 and 14 dpi. We identified 12 miRNA/targeted transcript pairs as robust candidates for the regulation of gall formation. A key role in tomato gall formation was demonstrated for the auxin-responsive miR167/*ARF8* transcript pair in functional analyses.

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99 Materials and methods

100 Biological materials, growth conditions and nematode infection

101 For in vitro experiments, seeds of Solanum lycopersicum cv St Pierre or Micro-Tom (wildtype, WT or transgenic *pARF8A::GUS* and *pARF8B::GUS* lines (Bouzroud *et al.*, 2018)) 102 103 were surface-sterilized with chlorine solution (44% active chlorine) and washed three times 104 with water. Ten to 15 sterile seeds were sown on a Gamborg B5 (Duchefa Biochemie) agar 105 plates (1x Gamborg B5; pH = 6.4; 1% Sucrose; 0,7% Agar), placed at 24°C for 48 hours for 106 germination, and finally transferred in a growth chamber (8h light; 16h dark, 20°C). M. 107 incognita (Morelos strain) J2s were sterilized with HgCl2 (0.01%) and streptomycin (0.7%)as described before (Caillaud & Favery, 2016). One to two weeks after germination, roots 108 109 were inoculated with 1,000 sterile J2s per petri dishes. 110 For in soil infection assay, S. lycopersicum (cv Micro-Tom) WT plants and CRISPR lines

111 (arf8a, arf8b and arf8ab) were sown and individually transferred in pots filled with a mixture 112 of sand and soil (1:1, vol:vol), kept at 4°C for 48 hours, then transferred in a growth chamber 113 (16h light and 8h dark, at 24°C). Two weeks after germination, each plant was inoculated 114 with 200 J2s. Infection rate was evaluated six weeks after inoculation. The root system of 115 each plant was collected, rinsed with tap water, weighted and stained for 30 s. in eosin 116 (Sigma) solution (0.5%). Galls and egg masses were counted for each root under the 117 binocular magnifier MZFLIII (Leica). Mann–Whitney $U\Box$ tests ($\alpha = 2.5\%$) were performed 118 to determine the significance of the differences in the numbers of egg masses and galls per 119 root observed between mutants and WT.

120

121 BABB clearing

For giant cell area measurements, galls were collected 21 days post-infection (dpi), cleared in
benzyl alcohol/benzyl benzoate (BABB) as previously described (Cabrera *et al.*, 2018; Mejias *et al.*, 2021) and examined under an inverted confocal microscope (model LSM 880; Zeiss).
The mean areas of giant cells in each gall, for WT and CRISPR lines, for two biological
replicates, were measured with Zeiss ZEN software. The impact of the mutation on the giant
cell surface was analysed using a Mann & Whitney Test.

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129 **RNA extraction**

130Total RNAs, including small RNAs (< 200 nt), were isolated from *in vitro* tomato (St Pierre)

galls or uninfected root fragments at 7 and 14 dpi. Approximately 40 galls or uninfected roots

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132 devoid meristems were independently frozen into powder by using a tissue lyser (Retsch;

133 MM301) at 30 Hertz frequency for 30 seconds with 4 mm tungsten balls (Retsch; MM301).

134 Total RNAs were extracted from these samples with the miRNeasy Mini Kit (Qiagen),

- according to the manufacturer's instructions, with three additional washes in RPE buffer.
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137 RNA sequencing

Small RNA libraries were generated by ligation, reverse transcription and amplification (11 cycles) from total RNAs (1 µg), with the reagents of the NEB Next Small RNA Library Prep Set for Illumina. Libraries were then quantified with the Bioanalyzer High Sensitivity DNA Kit (Agilent) and sequenced at the Nice-Sophia Antipolis functional genomics platform (France Géenomique, IPMC, Sophia Antipolis, France). The full raw sequencing data were submitted to the GEO database (http://www.ncbi.nlm.nih.gov/geo/), accession number PRJNA799360.

PolyA-RNA libraries from St Pierre tomato were generated from 500 ng of total RNA using
Truseq Stranded mRNA kit (Illumina). Libraries were sequenced on a NextSeq 500 platform
(Illumina) with 2□×□75bp paired-end chemistry as described in (Mejias *et al.*, 2022). RNAseq data are available at SRA database accession number #PRJNA799360. PolyA sequencing
of galls from the *arf*8 mutants and microtom wild type was performed by the Beiging
Genomics Institute (BGI) by using the DNBSeq technology. RNA-seq data are available at
SRA database accession number #XXX

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153 miRNAome and transcriptome analysis

154 For each small RNA library, adapters were trimmed and reads matching ribosomal RNA, 155 mitochondrial RNA and repeat sequences were removed by performing Blast analyses with 156 the sequences listed in the Rfam database (Nawrocki et al., 2015). The STAR 2.5 aligner (: --157 twopassMode Basic --alignEndsType EndToEnd) was then used to align the trimmed reads (Dobin et al., 2013) on a virtual concatenated genome generated from the S. lycopersicum 158 159 genome (V3.01, annotation V3.2) and the *M. incognita* genome (Blanc-Mathieu *et al.*, 2017). 160 Each read was attributed to the S. lycopersicum and/or M. incognita genome on the basis of 161 the best alignment obtained. Low-quality mapped reads were removed. The htseq-count 162 package version 0.9.1 (Anders *et al.*, 2015) was used to count reads mapping perfectly onto 163 the S. lycopersicum genome. The counts for protein coding genes from each replicate were 164 used for differential expression analysis with the R package EdgeR version 3.4.1 (Robinson et 165 al., 2009) and DSeq2 (Anders & Huber, 2010). Differentially expressed miRNAs, identified 166

with a false discovery rate of 5% (adjusted pvalue<0.05; Benjamini-Hochberg adjustment).

167 *De novo* microRNA encoding genes were predicted in tomato genome V3.0 by using three

168 algorithms MirCat (Paicu et al., 2017), Shortstack (Axtell, 2013) and MirDeep plant (Yang &

169 Li, 2011) with default parameters. The sequence homology between newly predicted miRNA 170 mature sequences and mature miRNA sequences listed in miRBase 22.1 was analysed by 171 using SSearch algorithm (Kozomara et al., 2019). The HTSEQCOUNT package (Anders et 172 al., 2015) was used to count reads mapping perfectly onto the predicted S. lycopersicum 173 mature microRNA 5P or 3P sequence. Reads mapping to multiple loci were counted for each 174 of the loci concerned. The counts for mature miRNAs (5P and 3P) from each replicate were 175 used for differential expression analysis by using DSeq2 statistical analysis (Anders and 176 Huber, 2010). Mature miRNAs with an adjusted p value below 0.05 were considered as

177 differentially expressed.

178 GO analyses of genes differentially expressed in galls were performed by using over-179 representation test from PANTHER analysis tools (Mi et al. 2019) with a Fisher's exact test,

180 a FDR threshold of 0.05 and by selecting "Biological Process » as GO category.

181

182 **Degradome analysis**

183 Degradome libraries were constructed from total RNAs extracted from galls at 7 and 14 dpi 184 by Vertis Biotechnologie (Freising, Germany) using the parallel analysis of RNA ends 185 (PARE) protocol described by German et al. (2009). The PARE libraries were sequenced on 186 an Illumina High Sequencing 2000 platform. The full raw sequencing data were submitted to 187 the GEO database (http://www.ncbi.nlm.nih.gov/geo/), accession XXX. To identify miRNA 188 targets, degradome reads were analysed and classified by using the CleaveLand 4.0 (Addo-189 Quaye et al., 2009) algorithm with default parameters. All hits are classified into five 190 categories based on the abundance of the diagnostic cleavage tag relative to the overall profile 191 of degradome tags matching the targets.

192

193 **GUS staining analysis**

194 We localized the promoter activity in tomatoes transgenic lines expressing a reporter gene 195 GUS fused to the promoter of the two tomato genes ARF8A (Solyc02g037530) and ARF8B 196 (Solyc03g031970) (pARF8A:GUS and pARF8B:GUS lines) (Bouzroud et al., 2018). We 197 inoculated 21-day-old seedlings in vitro, as described above. GUS staining was performed 7 198 and 14 dpi as previously described (Noureddine et al. 2022), and the roots were observed under a Zeiss Axioplan 2 microscope. Stained galls were dissected, fixed by incubation in 1%

- 200 glutaraldehyde and 4% formaldehyde in 50 mM sodium phosphate buffer pH 7.2, dehydrated,
- and embedded in Technovit 7100 (Heraeus Kulzer, Wehrheim, Germany), according to the
- 202 manufacturer's instructions. Sections were cut and mounted in DPX (VWR International Ltd,
- 203 Poole, UK), and observed under a Zeiss Axioplan 2 microscope (Zeiss, Jena, Germany).
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205 **Quantitative RT-PCR**

206 Total RNA was extracted from galls and uninfected roots produced in soil with the miRNeasy 207 kit (QIAGEN) according to the manufacturer's instruction. 500ng of total RNA were 208 subjected to reverse transcription with the Superscript IV reverse transcriptase (Invitrogen). 209 qPCR analyses were performed as described by Nguyen *et al.* (2018). We performed qPCR on triplicate samples of each cDNA from three independent biological replicates. SIPSKR1 210 211 (Solyc01g008140) and a gene coding for a Sucrose Synthase (SuSy3, Solyc07g042550) were 212 used for the normalization of qRT-PCR data. Quantifications and statistical analyses were 213 performed with SATqPCR (Rancurel et al., 2019), and the results are expressed as 214 normalized relative quantities. Primers used to amplify the premature miRNAs and the 215 transcripts are listed in Table S1.

216

217 Generation of ARF8 mutants by CRISPR/Cas9

For CRISPR/Cas9 construct, the sgRNA sequence (AAGCTTTCAACATCAGGAA) commune to SIARF8a and SIARF8b was designed by using the CRISPR-P website tool (http://cbi.hzau.edu.cn/crispr/). The sgRNA was cloned into pAGM4723 final vector by golden gate ligation method. Construct was confirmed by sequencing before introduction into the C58 *Agrobacterium tumefaciens* strain. Tomato seedlings were used for the next step plant transformation according to Hao et al. (2015).

224

225 **Results**

226 Gall formation results from a massive reprogramming of gene expression in root cells

Two statistical methods, DSeq2 and EdgeR, were used to compare transcript levels between tomato galls and uninfected roots. These two methods were applied to 19,918 genes, and those found to be differentially expressed by both methods, with an adjusted *p*-value below 0.05, were identified as differentially expressed genes (DEG). We found 1,958 DEGs at 7 dpi (**Table S2**) and 3,468 DEGs at 14 dpi (**Table S3 and Figure 1a**). In total, 1,239 genes were 232 identified as DEGs at both 7 and 14 dpi, including 625 genes downregulated and 600 genes 233 upregulated at both time points and 14 DEGs with opposite patterns of change in expression 234 levels at 7 and 14 dpi. The 719 genes displaying differential expression in galls specifically at 235 7 dpi comprised 327 upregulated and 392 downregulated genes. The 2,229 genes displaying 236 differential expression in galls specifically at 14 dpi comprised 1,006 upregulated and 1,223 237 downregulated genes. The change in gene expression in galls detected by sequencing was 238 confirmed by RT-qPCR for six of these genes at 7 dpi and six at 14 dpi (Figure S1 and 239 Table S4). Gene ontology (GO) analysis DEGs in galls at 7 and/or 14 dpi revealed an 240 overrepresentation of genes associated with biological processes previously reported to be 241 involved in the formation of giant cells (**Table S5**), including i) "cell division", with multiple 242 categories linked to cytokinesis and cell wall biogenesis, ii) "response to auxin", iii) 243 "response to endogenous stimulus" (including response to hormone and to cytokinin) and iv) 244 "response to abiotic stress". This analysis confirms that a massive reprogramming of gene 245 expression in root cells underlies the formation of galls and feeding cells.

246

247 microRNAs regulate gene expression in galls

248 For the identification of regulators of gene expression in galls, we constructed libraries of 249 small non-coding RNAs from tomato galls and uninfected roots, from three independent 250 replicates at two points of gall development: 7 and 14 dpi. These libraries were sequenced, 251 generating a total of 333,949,327 raw reads (**Table S6**). The reads were cleaned and mapped 252 to a virtual genome constructed from the S. lycopersicum genome (genome V3.0; ITAG3.3) 253 concatenated with the genome of *M. incognita* (genome V2.0; Blanc-Mathieu et al., 2017) to 254 reflect the dual composition of root galls. A de novo prediction of microRNAs was then 255 performed (**Table S7**), based on an integration of the results of three prediction algorithms: 256 MirCat, Shortstack and MirDeep plant.

257

258 Levels of microRNA expression were compared between galls and uninfected roots in DSeq2 259 statistical analyses (Anders & Huber, 2010). We identified 174 mature microRNAs (5P 260 and/or 3P) corresponding to 148 MIR genes as differentially expressed (DE) between 261 uninfected roots and galls at 7 and/or 14 dpi (Table S8). We identified 129 of the 174 mature 262 microRNAs DE in galls as specifically DE at 7 dpi, 11 as specifically DE at 14 dpi and 34 263 mature microRNAs were found to be DE in galls at both 7 and 14 dpi (Figure 1b). These 148 264 MIR genes DE in galls comprised 65 known MIR genes listed in miRbase (Kozomara et al., 265 2019) and 73 previously unknown MIR genes. The 65 known MIR genes DE in galls are

organized into 20 miRNA families, 14 of which are conserved between tomato and other

267 plants, the other six being specific to tomato.

268

Integration of data from the transcriptome, small RNAs and degradome sequencing to construct a gene-microRNA regulation network in galls

271 Once the miRNAs expressed in galls had been identified, the transcripts cleaved by the 272 microRNAs in galls were identified by degradome sequencing (German et al., 2009) on 273 mRNA extracted from galls at 7 (G7) and 14 dpi (G14). The CleaveLand pipeline (Addo-274 Quaye et al., 2009) was used to analyze degradome sequencing data and to predict the 275 mRNAs cleaved by miRNAs in galls. We restricted our analysis to the highest confidence 276 targets by selecting CleaveLand categories 0 and 1 with a degradome p-value below 0.05. In 277 total, 153 transcripts targeted by microRNAs in galls were identified (**Table S9**), including 58 278 targets common to both the G7 and G14 libraries, whereas 45 targets were identified 279 specifically in the G7 library and 50 were found only in the G14 library. We identified 111 280 targets of 135 known miRNAs from 39 known miRNA families. The 298 newly identified 281 miRNAs in galls included 46 that targeted 47 transcripts in galls at 7 and/or 14 dpi.

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283 We integrated transcriptome, microRNA and degradome sequencing data to construct a a 284 gene-miRNA regulation network putatively involved in the gall formation. Transcriptome 285 analysis showed that 32 of the 153 transcripts identified as targeted by microRNAs expressed 286 in galls were DE in galls. Nineteen of the targeted genes were DE in galls at both 7 and 14 287 dpi; 11 of these genes were upregulated and eight were downregulated. Five targeted genes 288 were specifically DE at 7 dpi, including three transcripts that were upregulated and two that 289 were downregulated in galls. At 14 dpi, only eight transcripts identified as targets were DE, 290 three of which were upregulated, the other five being downregulated. Most plant miRNAs 291 silence gene expression by cleaving the targeted transcripts. An inverse correlation of 292 expression profiles between the microRNA and its target gene is, therefore, usually expected. 293 We identified 12 miRNA/mRNA pairs for which such an inverse correlation of expression 294 levels was observed (**Table 1**). These miRNA/mRNA pairs are the most robust candidates for 295 involvement in gall formation.

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ARF8 auxin-related transcription factors are expressed in nematode-induced feeding sites

Among the 12 microRNA/mRNA pairs with opposite patterns of gene expression, two of the strongest genes candidates for regulation by a microRNA are the *AUXIN RESPONSE FACTORS 8A* (*Solyc02g037530*) and *8B* (*Solyc03g031970*), both of which are cleaved miR167. These two genes are ARF transcription factors, which relay auxin signaling at the transcriptional level by regulating the expression of auxin-responsive genes (Guilfoyle & Hagen, 2007). Transcriptomic analyses of galls showed that *ARF8B* was overexpressed in tomato galls at 7 and 14 dpi and *ARF8A* was overexpressed at 14 dpi (**Table 1**).

308

309 Five *MIR167* genes were identified in the tomato genome, including the four described by Liu 310 et al. (2014), all of which have the same mature sequence and are downregulated in galls, 311 whereas both the ARF8 genes were found to be upregulated. The fifth MIR167 gene was 312 annotated as SLYMIR167B in miRBase (Kozomara et al., 2019) and was not DE in galls. 313 However, it was expressed at a much lower level than the other four MIR167 genes (Figure 314 S2). ARF8B transcripts have been shown to be cleaved by miR167 in tomato (Liu et al., 315 2014) and this regulation in conserved in A. thaliana (Wu et al., 2006). The downregulation 316 of miR167 and the upregulation of ARF8A and ARF8B observed in galls suggest that, by 317 repressing MIR167 expression, the RKN prevents the ARF8 silencing by miR167 that occurs 318 in uninfected roots.

319

320 We investigated the spatiotemporal expression of ARF8A and ARF8B in RKN-infected roots 321 in vivo further, by analyzing the activity of both the ARF8A and ARF8B promoters in 322 transgenic tomato lines expressing promoter-GUS fusions (Bouzroud et al., 2018). Strong 323 blue staining indicating GUS activity was observed 7 and 14 dpi in galls from two 324 independent pARF8B::GUS and pARF8A::GUS lines and in root tips from uninfected roots 325 (Figure 2a-f). Histological sections of the galls showed strong GUS staining within the giant 326 feeding cells and in neighboring cells (NC) at 7 and 14 dpi for both ARF8A and ARF8B lines 327 (Figure 3a-d). The strong activity of both the ARF8A and ARF8B promoters observed in galls 328 *in vivo* confirms the upregulation in galls observed on transcriptomic analysis.

329

Generation of *SlARF8A* and *SlARF8B* mutants by the CRISPR/Cas9 gene-editing
system.

332 We investigated the function of *SlARF8A* and *SlARF8B* during plant-nematode interaction, by 333 generating tomato Micro-Tom *slarf8* KO mutants with CRISPR/Cas9 gene-editing 334 technology. SIARF8A and SIARF8B single and double mutants were obtained with a sgRNA 335 complementary to a region identical in both genes (Figure 4a). The transformed lines were 336 screened and six independent R0 lines were generated and validated for the presence of the 337 construct in their genome. All mutants had mutations in the targeted region and the features 338 of the mutant plants were similar. In the R1 and R2 generations, the presence of mutation was 339 confirmed in more progeny lines. Three Cas9-free and homozygous mutant lines containing 340 single (SlARF8A or SlARF8B) or double mutation (SlARF8A and SlARF8B) were selected in 341 the following experiments. These three mutants types can be classified as SlARF8A single 342 mutation (*slarf8acr*), *SlARF8B* single mutation (*slarf8b-cr*) and *SlARF8A&B* double mutation 343 (*slarf8a&b-cr*) (Figure 4b). As shown in figure 4, the deletion mutations led to a frame shift 344 mutation followed by an early stop codon leading to the expression of truncated SIARF8 345 proteins that do not contain the ARF family functional domains B3, III and IV. For slarf8a-346 cr, a deletion of 2 nt was detected in the sgRNA1 targeted region, leading to a 13 amino acid 347 (aa) protein rather than the 845 aa WT protein; for *slarf8b-cr*, an 11nt deletion was observed 348 in the sgRNA1 target region, leading to a 9 aa protein sequence rather than 843 aa in WT 349 protein; and, for *slarf8a&b-cr*, there was a 2 nt deletion on *SlARF8A* and a 4 nt deletion on 350 *SlARF8B* resulting in a 13 aa SlARF8A protein and a 20 AA SlARF8B protein.

351

352 ARF8 auxin-related transcription factors are involved in tomato-RKN interactions

353 We investigated the role of both ARF8A and ARF8B in gall development, by analyzing the effect of CRISPR deletions within ARF8 coding sequences on M. incognita infection. The 354 arf8a^{CR-2}, arf8b^{CR-11} and arf8ab^{CR-2,4} double-mutant CRISPR lines had root phenotypes 355 356 identical to that of the WT (Figure S3 and Table S10). The rate of infection of these 357 CRISPR lines after inoculation with *M. incognita* was determined by counting the galls on 358 infected roots and the egg masses produced by the adult females at the root surface at the end 359 of the RKN lifecycle. A significant large decrease, by approximately 50%, in the numbers of galls and egg masses was observed for the $arf8a^{CR-2}$, $arf8b^{CR-11}$ and $arf8ab^{CR-2,4}$ lines relative 360 361 to WT plants (Figure 5a and Table S10). Thus, ARF8 disruption decreases suceptibility to infection, thereby demonstrating that ARF8A and ARF8B play a key role in the plant-RKN 362 interaction. We investigated the reasons for the lower susceptibility of the $arf8a^{CR-2}$, $arf8b^{CR-2}$ 363 ¹¹ and $arf8ab^{CR-2,4}$ lines, by measuring the area covered by giant cells directly with a confocal 364 365 microscope, after gall clearing with BABB (Cabrera et al. 2017). A comparison of the mean surface areas covered by giant cells in each gall showed that giant cells from the CRISPR lines were approximately 30% smaller than those from control plants (**Figure 5b-c**). These

lines were approximately 30% smaller than those from control plants (Figure 5b-c). These

results demonstrate that the expression of *ARF8A* and *ARF8B* is required for correct giant cell

369 development during the tomato-RKN interaction.

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371 Identification of *ARF8A*- and *ARF8B*-regulated genes in galls

372 For the identification of genes regulated by ARF8A and ARF8B in galls, mRNA from 14 dpi galls of arf8a^{CR-2}, arf8b^{CR-11} mutants and WT Micro-Tom tomatoes were sequenced. 373 Transcript levels in the galls from WT and mutant plants were compared in DESeq2 and 374 EdgeR statistical analyses. These two methods identified 189 and 66 genes, respectively, as 375 differentially expressed between galls from WT and $arf8a^{CR-2}$ or $arf8ab^{CR-2,4}$ mutants (Table 376 377 **S11a and b**). Several auxin-inducible genes have already been identified as candidate genes 378 downstream from ARF8s in tomato floral tissue (Liu et al. 2014): SMALL AUXIN-379 UPREGULATED RNAs (SAUR; Solyc07g042490.1.1) and two EXPANSINS (Solyc08g077900.3.1 and Solyc08g077910.3.1) were found to be repressed in arf8a^{CR-2}tomato 380 381 galls. Only 16 DEGs were common to both mutants (Table S11c). These genes are located 382 directly or indirectly downstream from ARF8 in tomato galls. No defense marker genes, such 383 orthologs of salicylic acid-mediated response marker PATHOGENESIS-RELATED as 384 PROTEIN-1 (PR1), jasmonic acid-mediated defense maker PROTEINASE INHIBITOR 2 (PIN2) or PLANT DEFENSINS, were found to be induced in $arf8a^{CR-2}$ or $arf8ab^{CR-2,4}$ galls. 385 This absence of defense marker gene induction in the galls of $arf8a^{CR-2}$ and $arf8ab^{CR-2,4}$ 386 mutants indicates that the decrease in galls and egg masses observed is due to a loss of 387 388 susceptibility rather than the induction of a plant defense mechanism. Together with the 389 requirement of ARF8s for correct giant cell development, this findings supports a key role for 390 ARF8s in feeding site formation.

391

392 **Discussion**

393 Identification of miRNA/mRNA target pairs involved in gall formation

RKN of the genus *Meloidogyne* are highly polyphagous sedentary plant parasites that can induce the formation of giant feeding cells in most crop species. The formation of feeding cells by RKN has been shown to result from a massive reprogramming of plant gene expression induced by the nematode (Jammes *et al.*, 2005; Fuller *et al.*, 2007; Ibrahim *et al.*, 2011; Damiani *et al.*, 2012; Yamaguchi *et al.*, 2017; Kaur *et al.*, 2017). In this study, we 399 identified 4187 protein-coding genes, corresponding to 12.3% of all annotated tomato genes, 400 as differentially expressed in tomato galls 7 and 14 dpi with M. incognita relative to 401 uninfected roots. This proportion is consistent with previous transcriptomic analyses in 402 Arabidopsis and tomato (Yamaguchi et al. 2017; Portillo et al. 2013). We investigated the 403 regulators of this reprogramming of gene expression, by analyzing the expression of 404 microRNAs and mRNAs in galls and uninfected tomato roots and using degradome 405 sequencing to identify transcripts cleaved under the guidance of microRNAs. Finally, a gene 406 regulation network for gall development was built by integrating all these -omics data. 407 Twelve of the 153 transcripts identified as targeted by microRNAs in tomato galls in 408 degradome analysis were considered the most robust candidates, based on their expression 409 profiles, which were inversely correlated with those of the corresponding microRNA. Some 410 of these 12 miRNA/mRNA pairs have already been reported to be involved in the plant-411 nematode interaction in Arabidopsis: the miR408/UCCLACYANINE (blue copper protein) 412 and the miR398/copper superoxide dismutase (Noureddine et al. 2022), and the auxin-413 regulated miR172/APETALA2 pair (Díaz-Manzano et al. 2018). Moreover, other pairs as 414 also appear to be interesting candidates based on the processes in which they are involved. 415 This is the case, for example for the miR164/NAC transcription factor involved in lateral root 416 development (Guo et al. 2005). In Arabidopsis root, the auxin-mediated induction of miR164 417 induces the silencing of NO APICAL MERISTEM-1 (NAC1) transcripts, thereby affecting 418 transmission of the auxin signal and regulating lateral root growth.

419

420 *ARF8s* are regulated at posttranscriptionally by miR167 in tomato roots

421 Among the 12 most robust miRNA/mRNA pairs identified as putatively involved in the 422 formation of galls, the ARF8A and ARF8B targets of miR167 were selected for further 423 functional analyses based on their role in auxin signaling, as auxins are a class of hormones 424 controlling root development and architecture (De Smet et al., 2007; Quint & Gray, 2008; 425 Majda & Robert, 2018). ARF8 is an auxin-responsive factor (ARF). The transcription factors 426 of this family regulate the activation or repression of auxin-induced genes by binding to the 427 auxin response elements (AuxREs) in their promoters (reviewed in Guilfoyle and Hagen, 428 2007; Chandler, 2016; Li et al., 2016). The ARF gene family is conserved across the plant 429 kingdom and is well described in various plant species, including A. thaliana (23 genes) 430 (Hagen & Guilfoyle, 2002), S. lycopersicum (22 genes) (Zouine et al., 2014), and Oryza 431 sativa (25 genes) (Wang et al., 2007). The ARF family has been implicated in the regulation 432 of plant developmental processes, such as embryo morphogenesis (Rademacher *et al.*, 2011),

433 the formation of lateral roots in response to low levels of nitrogen (Gifford *et al.* 2008), the 434 formation of adventitious roots (Lee et al., 2019), leaf structure and senescence (Wilmoth et 435 al., 2005), flower development (Ellis et al., 2005) and fruit initiation (Liu et al., 2014). Like 436 ARF5, 6, 7 and 19, ARF8 has been described as a transcriptional activator (reviewed in 437 Guilfoyle and Hagen, 2007). In Arabidopsis, a partial redundancy between ARF8 and ARF6 438 has been reported, with both these activators silenced by miR167 (Reeves *et al.* 2012). The 439 Arabidopsis arf6arf8 double mutant has defective flower development, as the flower is 440 entirely sterile (Nagpal, 2005), whereas the arf8 mutant presents defects of pollination and 441 fertilization (Tian et al., 2004; Vernoux et al., 2011). A role for ARF8 has been reported in 442 the formation of lateral roots in Arabidopsis and soybean (Gifford et al., 2008; Wang et al. 443 2015) and in the formation of adventitious roots (Gutierrez et al., 2009). ARF6 and ARF8 444 were recently implicated in cambium establishment and maintenance (Ben-Targem et al., 445 2021). The *arf6arf8* double mutant displays low levels of xylem occupancy and an absence of 446 fiber accumulation until very late stages of plant growth.

447

448 In A. thaliana and tomato, ARF8 genes are regulated posttranscriptionally by miR167 (Wu et 449 al., 2006; Liu et al., 2014). Transcriptomic analyses of galls showed that ARF8B is 450 overexpressed in tomato galls at 7 and 14 dpi, whereas ARF8A is overexpressed at 14 dpi. 451 The infection of tomato lines expressing the GUS reporter gene under the control of the 452 ARF8A or ARF8B promoter revealed high levels of activity for both ARF8 promoters in giant 453 cells and neighboring cells at 7 and 14 dpi, confirming the overexpression observed in the 454 transcriptomic analyses. Gall degradome analysis identified ARF8A and ARF8B transcripts as 455 cleaved by members of the miR167 family. Four tomato MIR167 genes encode mature 456 proteins with identical sequences and are downregulated in tomato galls at 7 and 14 dpi. This 457 suggests that the ARF8A and ARF8B transcripts are cleaved by miR167 in uninfected tomato 458 roots, as demonstrated in A. thaliana roots. We showed that RKN infection induces the 459 inhibition of miR167 in galls, thereby decreasing the cleavage of ARF8 transcripts by 460 miR167, resulting in an overexpression of ARF8A and ARF8B in galls.

461

462 Auxin is a major factor regulating the formation of feeding cells in tomato

We used tomato lines with CRISPR deletions within the *ARF8A*, *ARF8B* and *ARF8AB* coding sequences to analyze the function of *ARF8* in plant-nematode interactions. The *arf8a*, *arf8b* and *arf8ab* lines displayed decreased susceptibility to nematode infection, with fewer gall and egg masses in mutants than in wild-type tomato plants. Moreover, the phenotyping of giant 467 feeding cells within cleared galls showed the giant cells from the three CRISPR lines to be 468 smaller than those from the wild type. These defects, associated with CRISPR mutations, 469 confirmed the involvement of *ARF8A* and *ARF8B* in the tomato response to RKN interaction 470 and that requirement for functional ARF8A and ARF8B for the correct development of 471 feeding cells.

472

473 MicroRNAs and transcription factors regulate ARF8 expression, whereas auxin peaks 474 regulate ARF8 activity: when auxin levels are high, ARF8 (class II ARFs) activates the 475 transcription of auxin-responsive genes (Tiwari et al., 2003). Auxin is known to be a major 476 factor regulating the formation of feeding cells in response to RKN signals (reviewed in 477 Gheysen and Mitchum, 2019; Oosterbeek et al., 2021). Microarray analyses of A. thaliana 478 gall transcripts have revealed an early activation of genes responsible for auxin homeostasis 479 and auxin-responsive genes, and a downregulation of repressors of auxin responses (Hammes 480 et al., 2005; Jammes et al., 2005; Barcala et al., 2010). Studies of lines expressing reporter 481 genes under the control of the synthetic auxin-responsive DR5 promoter showed that this 482 promoter is activated in galls induced by RKN (Hutangura et al., 1999; Karczmarek et al., 483 2004; Absmanner et al., 2013b). In A. thaliana galls, a strong signal was detected in both 484 giant cells and neighboring cells at 4 dpi for DR5:GUS lines (Cabrera et al., 2014). This 485 increase in auxin levels in the galls may be controlled by either the plant or the nematode. 486 Auxin-mimicking compounds have been found in nematode secretions (De Meutter et al., 487 2003, 2005).

488

489 **ARF8** is involved in plant responses to biotic and abiotic stresses

490 ARF8 transcription factors have been implicated in plant responses to microorganisms. ARF8 491 is regulated in tomato leaves in response to biotic stresses, such as flagellin treatment or 492 infection with Pseudomonas syringae (Bouzroud et al., 2018). A recent study in Arabidopsis 493 provided evidence for a suppression of miR167 expression, together with an induction of 494 ARF6 and ARF8, in response to infection with P. syringae pv. tomato DC3000 in A. thaliana 495 (Caruana et al., 2020). The P35S:MIR167 and arf6 arf8 double mutants were found to be 496 more resistant to bacterial infection than the wild type. The authors suggested that ARF6 and 497 ARF8 modulate salicylic acid defenses response to *P. syringae* infection under miR167 498 regulation. Furthermore, soybean ARF8A and ARF8B have been shown to downregulate the 499 nodulation induced by miR167 (Wang et al., 2015). All these studies suggest that the auxin-500 responsive pathway, including miR167/ARF8, is a key actor in the response to

microorganisms. Analysis of the transcriptomes of two tomato CRISPR *arf8* mutants showed
no induction of genes associated with plant defense in galls from *arf8* mutants. This finding
supports the notion that the lower levels of RKN infection observed in *arf8* mutants result

from a loss of susceptibility rather than an enhancement of plant defense.

- 505
- 506

507 ARF8, an environmental hub connecting development and stress responses

508 We have shown that ARF8 genes are posttranscriptionally regulated by miR167 in galls, but 509 the transcriptional regulation of these genes has yet to be deciphered. ARF8 was recently 510 shown to be regulated by a complex network of multiple activating and repressing 511 transcription factors in A. thaliana (Truskina et al., 2021). Interestingly, some of these 512 transcription factors, such as WUSCHEL, Squamosa Promoter Binding Protein Like-13 513 (SPL13) and WRKY33, have also been implicated in plant development, and many ARF8 514 regulators are also associated with plant responses to biotic and abiotic stresses. Based on 515 these results, Truskina et al. suggested that ARF8 may act as an environmental hub 516 connecting development and stress responses mediating auxin responsiveness. The formation 517 of RKN-induced feeding sites interferes with plant developmental processes, including, in 518 particular, the development of lateral roots (Cabrera et al., 2014, Olmo et al. 2020), which 519 suggests that the nematode may hijack this process. For example, the transcription factor 520 LBD16 and the microRNA miR390a, two key components of the auxin pathway and 521 transducers of lateral root development, are involved in gall formation in Arabidopsis and 522 tomato (Cabrera et al., 2014, Olmo et al. 2020). ARF8 has also been implicated in lateral root 523 formation in Arabidopsis (Gifford et al. 2008) and may, therefore, integrate biotic stress and 524 developmental processes during the formation of giant feeding cells. The common induction, 525 in tomato and Arabidopsis galls, of ARF8 and of the transcription factor LBD16 (Olmo et al. 526 2020) and miR390a (Diaz Manzano et al. 2016), suggests that there may be a conserved 527 auxin-mediated molecular pathway in galls. Early in the development of galls in Arabidopsis, 528 at 3 dpi, the silencing of ARF3 by miR390 via tasiRNAs, and the induction of ARF5 have 529 been shown to be required for parasitism (Cabrera et al. 2014a; Olmo et al. 2020). These 530 results suggest that there is a complex network involving ARFs and microRNAs responsible 531 for mediating auxin signaling during the development of galls induced by RKN.

532

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542 Author contributions

543 Y.N., B.F and S.J.P. designed the study, performed the experimental work and wrote the

544 manuscript. Y.N. and C.M. produced biological material for sequencing. M.dR. analyzed

NGS data. M.Z. and JA designed, generated and characterized *arf8* CRISPR-Cas9 mutants.

546 KM participated to the qPCR analyses. M.Q., J.M. and P.A. participated in the writing of the

547 manuscript. All the authors analyzed and discussed the data.

548 Data avaibility

549 s

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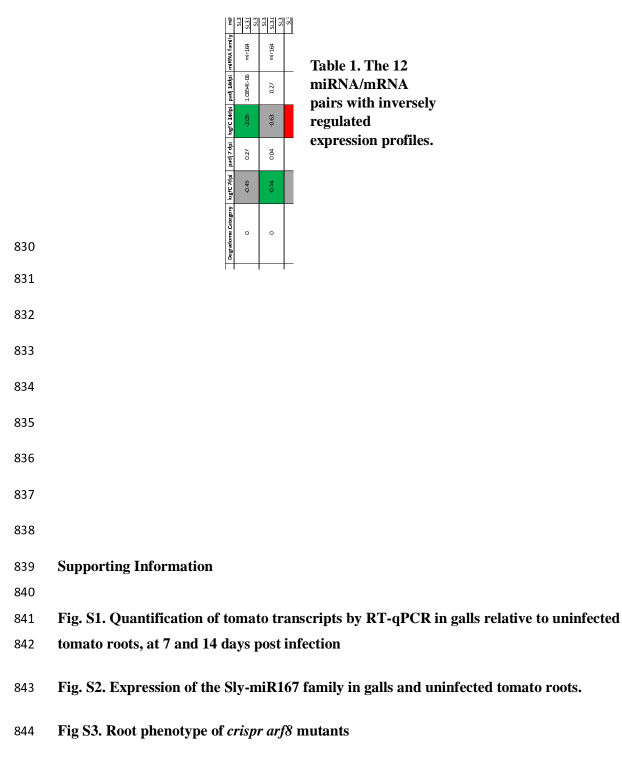
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- 845 **Table S1. Primers used for RT-qPCR.**
- Table S2. Protein-coding genes DE in galls at 7 dpi
- Table S3. Protein-coding genes DE in galls at 14 dpi

848 849	Table S4. Quantification of tomato transcripts in galls by RT-qPCR (G) relative to uninfected tomato roots (R), at 7 and 14 days post inoculation (dpi).
850	Table S5. Gene ontology (GO) analysis of the genes DE in galls at 7 and/or 14 dpi.
851 852	Table S6. The number of raw reads from the three libraries obtained from galls and uninfected roots of <i>S. lycopersicum</i> at 7 dpi and 14 dpi.
853	Table S7. De novo prediction of Solanum lycopersicum MIR genes.
854	Table S8. MicroRNAs differentially expressed in galls at 7 and/or 14 dpi.
855 856	Table S9. 153 transcripts targeted by miRNAs in galls at 7 and/or 14 dpi identified by degradome sequencing and CleaveLand analysiss.
857 858	Table S10. Infection assays in arf8 CRISPR lines infected with M. incognita and comparison to WT.
859	Table S11. Genes differentially expressed between crisprarf8 mutant galls and wild-type

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862 Figure legends

(WT) galls.

863

864 Figures and Tables

865 Table 1. The 12 miRNA/mRNA pairs with inversely regulated expression profiles. 866 Degradome analysis identified 12 genes targeted by miRNAs in galls. Solanum lycopersicum 867 (Solyc) gene expression levels were compared between galls and uninfected roots, by two 868 statistical methods (DSeq2 and EdgeR), and the expression of mature miRNAs was compared 869 by DSeq2. Gall/root fold-change differences in expression (LogFC) at 7 and 14 days post 870 infection (dpi) and the adjusted *p*-value obtained by the Benjamini-Hochberg method (adj *p*-871 value) are indicated for genes and miRNAs. The genes and mature microRNAs upregulated in 872 galls are shown in red, and those downregulated in galls are shown in green.

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Figure 1. Tomato protein-coding genes and miRNAs differentially expressed in *M. incognita*-induced galls relative to the corresponding uninfected roots, at 7 and/or 14 dpi. The numbers of genes differentially expressed at 7 days post inoculation (dpi) (pink) and/or 14 dpi (yellow) between galls and the corresponding uninfected roots are indicated inVenn diagrams.

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880 Figure 2. ARF8A and ARF8B are strongly transcribed in tomato galls induced by M. 881 *incognita*. The activity of the ARF8A and ARF8B promoters (pARF8A and pARF8B) was 882 studied in galls induced by *M. incognita* in *S. lycopersicum* lines expressing the 883 *pARF8A::GUS* or the *pARF8B::GUS* construct, at 7 and 14 days post inoculation (dpi). Blue 884 staining indicating GUS activity under the control of *pARF8A* was observed in (a) uninfected 885 root tips and (b-c) galls induced by *M. incognita* at 7 dpi (b) and 14 dpi (c). GUS activity 886 under the control of *pARF8B* was observed in (d) uninfected root tips and in (e-f) galls at 7 887 dpi (e) and 14 dpi (f). Bars: 500 µm.

888

889 Figure 3. ARF8A and ARF8B are strongly transcribed in 14 dpi nematode-induced 890 feeding sites. The activity of the ARF8A and ARF8B promoters (pARF8A and pARF8B) was 891 studied in galls induced by *M. incognita* in *S. lycopersicum* expressing the *pARF8A::GUS* (a-892 b) or the *pARF8B::GUS* (c-d) construct, at 7 and 14 days post inoculation (dpi). Gall sections 893 (5 µm) were cut after GUS staining and observed by dark-field microscopy. Red staining, 894 reflecting GUS activity, was observed in giant cells and neighboring cells in pARF8A::GUS 895 galls (a) 7 dpi and (b) 14 dpi. Strong GUS activity was observed in giant cells and 896 neighboring cells in the galls of pARF8B::GUS plants at (c) 7 dpi and (d) 14 dpi. *, giant 897 cells; nc: neighboring cells; Bars: (a,b) 100 µm (c,d) 50 µm.

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Figure 4. Tomato *slarf8-KO* **lines.** Generation of slarf8-KO mutant lines by CRISPR/Cas9.

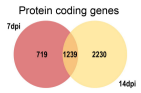
Guide RNAs (sgRNA, red bar) anchored next to the Zinc Finger Motif (ZFM) were designed
for CRISPR/Cas9 strategy. Protospacer adjacent motif (PAM) are indicated in blue. Mutations
within *slarf8* coding sequences corresponding to nucleotide deletions are shown in green.
Three types of mutants predicted to produce heavily truncated proteins were chosen for
further phenotypic characterization. These mutants are annotated as *arf8a-cr* (*arf8a* single
mutant), *arf8a-cr* (*arf8b* single mutant), *arf8ab-cr* (*arf8a* and *arf8b* double mutant). The
predicted mutated proteins are schematically illustrated (right panel).

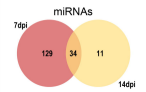
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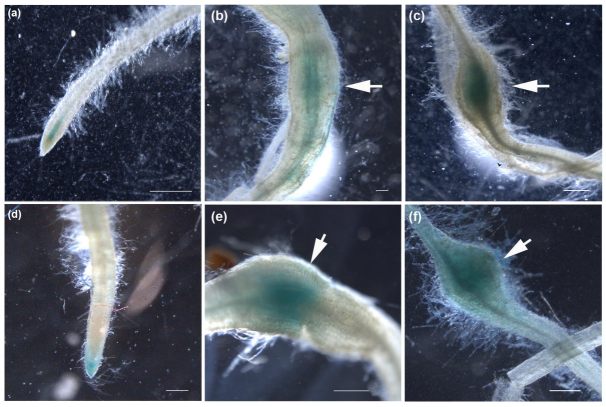
Figure 5. The single mutants $arf8a^{CR-2}$ and $arf8b^{CR-11}$ and the double mutant $arf8ab^{CR-2,4}$ 908 909 were significantly less susceptible to *M. incognita* than the wild type. (a), The 910 susceptibility of the single and double CRISPR-Cas9 mutant lines and wild-type (WT) 911 MicroTom plants to *M. incognita* was evaluated by counting the numbers of galls and egg 912 masses per plant (G/plant and EM/plant, respectively) in two independent infection assays in 913 soil. (b) The effect of deleting ARF8A and/or ARF8B on the development of giant feeding 914 cells was further evaluated by measuring the size of the feeding site produced in each mutant 915 line and comparing it to that in the WT. Galls were collected 21 days post infection (dpi) in *vitro* to measure the area (mm^2) covered by the giant cells by the BABB clearing method 916 917 (Cabrera et al., 2018). The impact of plant genotype was assessed in Mann-Whitney tests. *, 918 P < 0.05. Boxes indicate the interquartile range (25th to 75th percentile). The central lines 919 within the boxes represent the medians. Whiskers indicate the minimum and maximum usual 920 values present in the dataset. The circle outside the box represents an outlier. n, the number of 921 plants analyzed in each assay. Bars 50 µm.

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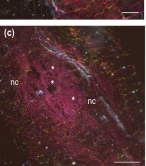






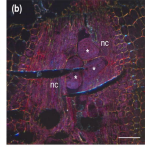
pARF8A::GUS

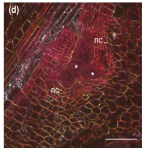
(a)

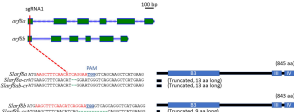


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SlorfRb-cr ATGAAGCTTTCAACATCAG-----CAGGCTCATGAAGG artSab-cr ATGAAGCTTTCAA----AGGAATGGGTCAGCAGGCTCATGAAGG



