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1 Short title: CARP9 and HYL1 interact to promote AGO1 stability

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- 3 Title: The intrinsically disordered protein CARP9 bridges HYL1 to AGO1 in the nucleus
- 4 to promote micro RNA activity
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- 12
- 13 One-sentence summary

An intrinsically disordered protein_interacts with HYPONASTIC LEAVES1 and
ARGONAUTE1 in a post-miRNA processing complex to promote ARGONAUTE1 stability
and miRNA activity.

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18 Author contributions: AHT, DAR and PAM conceived the research. AHT, DAR, DAC and LG carried out the experimental work. ALA analyzed sequencing data. FR and JEM performed the evolutionary and conservation analysis. AHT, DAR, PAM interpreted results and wrote the paper. All authors read, edited, and approved the final version of the manuscript.

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23 ABSTRACT

24 In plants, small RNAs (sRNA) are loaded into ARGONAUTE (AGO) proteins to fulfill their 25 regulatory functions. Micro RNAs (miRNAs), one of the most abundant classes of endogenous 26 sRNAs, are preferentially loaded into ARGONAUTE1 (AGO1). Such loading, long believed to 27 happen exclusively in the cytoplasm, was recently proposed to also occur in the nucleus. Here 28 we identified CONSTITUTIVE ALTERATIONS IN THE SMALL RNAS PATHWAYS9 29 (CARP9), a nuclear-localized, intrinsically disordered protein, as a factor promoting miRNA 30 activity in Arabidopsis (Arabidopsis thaliana). Mutations in the CARP9-encoding gene led to a 31 mild reduction of miRNAs levels, impaired gene silencing, and characteristic morphological 32 defects, including young leaf serration and altered flowering time. Intriguingly, we found that 33 CARP9 was able to interact with HYPONASTIC LEAVES1 (HYL1), but not with other 34 proteins of the miRNA biogenesis machinery. In the same way, CARP9 appeared to interact with mature miRNA, but not with pri-miRNA, positioning it after miRNA processing in the 35 36 miRNA pathway. CARP9 was also able to interact with AGO1, promoting its interaction with 37 HYL1 to facilitate miRNA loading in AGO1. Plants deficient in CARP9 displayed reduced levels of AGO1-loaded miRNAs, partial retention of miRNA in the nucleus, and reduced levels
of AGO1. Collectively, our data suggest that CARP9 might modulate HYL1AGO1 crosstalk,
acting as a scaffold for the formation of a nuclear post pri-miRNA processing complex that
includes at least HYL1, AGO1 and HSP90. In such a complex, CARP9 stabilizes AGO1 and
mature miRNAs, allowing the proper loading of miRNAs in the effector complex.

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44 INTRODUCTION

45 Post-transcriptional regulation of gene expression mediated by miRNAs controls numerous 46 processes during plant development and response to the environment (Li et al., 2017). In 47 Arabidopsis (Arabidopsis thaliana), RNA polymerase II transcribes MIRNA loci to primary 48 miRNAs (pri-miRNAs), which are processed into mature ~21 nt miRNA duplexes by multiple 49 and sequential cuts (Bologna et al., 2013; Zhu et al., 2013; Moro et al., 2018). The RNase-III endonuclease DICER-LIKE 1 (DCL1), together with the RNA binding protein HYPONASTIC 50 51 LEAVES 1 (HYL1) and the zinc finger protein SERRATE (SE), recognizes and processes pri-52 miRNAs into mature miRNA duplexes (Kurihara and Watanabe, 2004; Dong et al., 2008). In 53 addition, numerous accessory proteins were identified lately to regulate the miRNA biogenesis 54 at different stages (Rogers and Chen, 2012; Achkar et al., 2016; Manavella et al., 2019). HYL1 55 was proposed to remain bound to the mature miRNA duplexes and interact with HUA 56 ENHANCER 1 (HEN1), acting as a scaffold to ensure miRNA methylation (Li et al., 2005; 57 Yang et al., 2010; Baranauske et al., 2015). Mature miRNAs, potentially still bound by to 58 HYL1, are then loaded into an AGO protein, with AGO1 as the main effector of the miRNA pathway in plants, to fulfill their functions (Fang and Qi, 2016). Loading of miRNA duplexes 59 into AGO1 and the assembly of the RNA-Induced Silencing Complex (RISC) were initially 60 61 thought to be cytoplasmic exclusive processes (Bologna and Voinnet, 2014). This was inferred 62 from metazoan cells where miRNA loading into the RISC occurs in the cytosol, and the fact that 63 the homolog of the human EXPORTIN-5 of Arabidopsis, known as HASTY (HST), showed 64 impaired miRNA accumulation (Park et al., 2005). A recent report demonstrated that AGO1 is 65 at least partially loaded with miRNA duplexes in the nucleus and then exported to the cytosol as 66 an AGO1:miRNA complex (Bologna et al., 2018). It is not clear if the nuclear AGO1 loading is 67 dependent on HYL1. However, some evidences suggested that this is a certain possibility: 1) 68 AGO1 colocalizes with HYL1 in nuclear speckles (Fang and Spector, 2007); 2) HYL1 is 69 required for the proper miRNA loading and strand selection by AGO1 (Eamens et al., 2009; 70 Manavella et al., 2012); and 3) HYL1 remains bound to mature miRNAs after processing (Yang 71 et al., 2010). ENHANCED MIRNA ACTIVITY1 (EMA1) and TRANSPORTIN1 (TRN1) were 72 also shown to interact with AGO1 and modulate miRNAs loading into AGO1 (Wang et al., 73 2011; Cui et al., 2016). The finding of the nuclear loading of AGO1 not only created a new

74 model for miRNAs export but also rationalized the recently identified nuclear functions of
75 AGO1 (Dolata et al., 2016; Schalk et al., 2017; Liu et al., 2018).

76 Using a forward genetic screening, we identified CONSTITUTIVE ALTERATIONS IN 77 THE SMALL RNAS PATHWAYS 9 (CARP9), a predicted intrinsically disordered protein 78 (IDP), as a new partner of the miRNA pathway. Mutations in CARP9 produced morphological 79 alterations, a mild reduction in the miRNA accumulation, and impaired gene silencing. We 80 found that CARP9 interacts with HYL1 in discrete nuclear speckles promoting HYL1-AGO1 81 interaction. Our data suggest that CARP9 did not participate in pri-miRNAs processing, but 82 instead it is associated with HYL1 and mature miRNAs in a post-miRNA-processing complex. 83 In such complex, we found that CARP9 also interacts with AGO1 and HEAT SHOCK PROTEIN 90 (HSP90). CARP9 mutants presented low levels of AGO1-associated miRNAs, a 84 reduction of AGO1 levels, and an apparent depletion of miRNAs in the cytoplasm. All this 85 evidence allow us to suggest that CARP9 could be acting as a scaffold protein, connecting 86 87 HYL1 to AGO1 in a post pri-miRNAs processing complex, ensuring AGO1 stability thus 88 leading to the proper loading of the AGO1:miRNA complexes, likely before its export to the 89 cytosol.

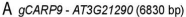
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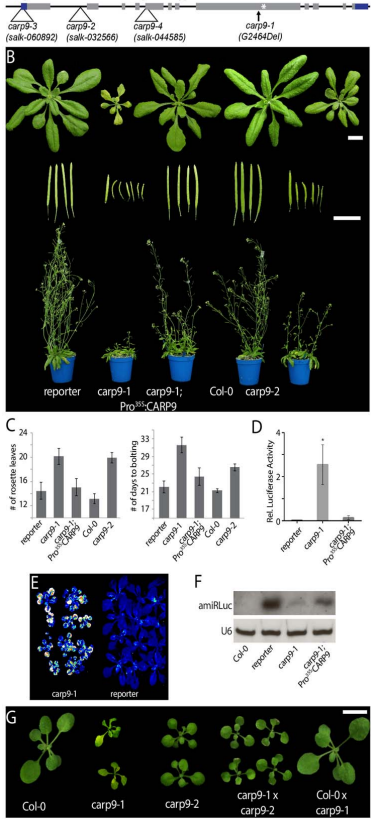
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92 RESULTS

93 Identification of CARP9 from a miRNA-activity screen

94 During the last years, we characterized several miRNA deficient mutants isolated from a genetic screening based on the silencing of a luciferase reporter by an artificial miRNA (amiRLUC) 95 96 (Manavella et al., 2012; Francisco-Mangilet et al., 2015; Karlsson et al., 2015; Re et al., 2020). 97 Here, using mapping by sequencing (Sun and Schneeberger, 2015), we localized the causal 98 mutation in one of the isolated plants, named constitutive alterations in the small RNAs 99 pathways 9 (CARP9), to a small region of chromosome 3 (Supplemental Figure S1A). Within 100 this region, we detected a single nucleotide deletion (G2464Del, AT3G21290, Chr3:7486786, 101 TAIR10) resulting in a premature stop codon and an aberrant C-terminal region of 34 amino 102 acids (Figure 1A). Compared to the reporter plants, this mutant allele (named *carp9-1*) showed a 103 reduced stature, irregular young leaves edges, pale green color, shorter and twisted siliques, a 104 delay in flowering time, and observable defects in the architecture of flowers, including shorter 105 stamens (Figure 1B, 1C, and Supplemental Figure S1B and S1C). Transformation of the mutant with a WT copy of CARP9 cDNA, either expressed under the 35S promoter or its native 106 107 regulatory region, fully reverted the morphological phenotype of the mutant (Figure 1B, 1C, and Supplemental Figure S1B and S1C). In the overexpressing lines, we selected lines with low 108 109 expression levels for the analyses. Besides, the carp9-1 mutant showed elevated luciferase 110 activity with a severe reduction of the amiRLUC levels (Figure 1D, 1E and 1F). All 111 morphological defects of carp9-1 were also observed, to a lesser degree, in a mutant with a T-DNA insertion within the first intron of this gene named carp9-2 (Figure 1A, 1B, 1C, and 112 Supplemental Figure S1B). To further confirm that the mutation in AT3G21290 was causing the 113 observed phenotype, we crossed capr9-1 and carp9-2 mutants and analyzed the progeny. The 114 115 compound heterozygotes offspring showed similar phenotypes to the parental lines confirming 116 that the mutation in CARP9 was the cause of the observed phenotypes (Figure 1G). In the case of a third allele, carp9-3 (SALK_060892), bearing a T-DNA insertion in the 5'-UTR region of 117 CARP9 approximately 100 bp upstream the ATG codon (Figure 1A), we were not able to 118 119 recover any homozygous line suggesting that plants are not viable when this mutation is 120 homozygous. Interestingly, we did not find signs of embryonic abortion or impaired germination/growth in carp9-3 +/- progeny (Supplemental Figure S1D and S1E). Nevertheless, 121 all genotyped seedlings were either WT or heterozygous for carp9-3 mutation suggesting a 122 123 problem with the male gametes. Supporting this scenario, we were not able to detect the carp9-3 124 insertion in the offspring of WT plants fertilized with pollen of *carp9-3* heterozygous plants. 125 Similarly, we failed to detect the homozygous insertion in the SALK 044585 allele (carp9-4), 126 which is annotated to have an insertion in CARP9 fifth exon (Figure 1A). Altogether, the 127 analysis of these alleles suggests that CARP9 is essential for the plant development and points 128 carp9-1 and carp9-2 as hypomorphic alleles. In this sense, carp9-1 may still produce a





129 truncated but active protein, while carp9-2 partially reduces CARP9 transcription as indicated

130 by RT-qPCR analysis (Supplemental Figure S1F).

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132 CARP9 displayed impaired activity of the miRNA pathway

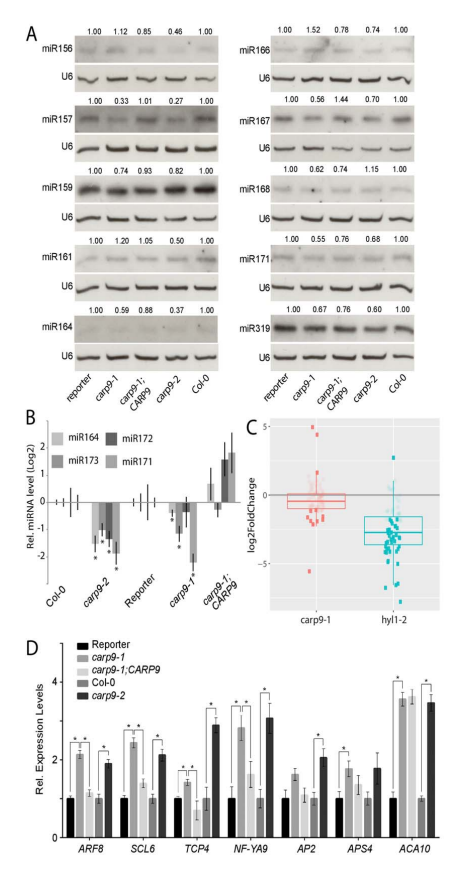
As carp9-1 was isolated as a miRNA deficient mutant, we explored the endogenous levels of 133 134 several miRNAs by RNA blots and RT-qPCR. The quantification of miRNAs revealed a mild 135 reduction of several tested miRNAs in the single mutants, as well as in the carp9-1/carp9-2 136 compound heterozygous plants (Figure 2A, 2B, and Supplemental Figure S2A). As expected, 137 the overexpression of CARP9 in carp9-1 restored miRNA levels (Figure 2A and 2B). Aiming to 138 explore the genome-wide profile of miRNA accumulation in the *carp9* mutant, we performed 139 Illumina small RNA sequencing of *carp9-1* mutants. The results were consistent across 140 replicates (Supplemental Figure S2B) and supported our northern blot results. We observed an 141 overall reduction of miRNAs and miRNA*s accumulation, although to a lesser degree than hyl1-2 plants, used as controls of impaired miRNA production (Figure 2C, Supplemental Figure 142 143 S2C, Supplemental Table S1). Only the nuclear-acting miR845 (Borges et al., 2018), miR391, 144 and miR827 appeared over accumulated in *carp9-1* (Supplemental Table S1). 145 Coincidently with impaired activity of the miRNA pathway in the mutants, we observed an

145 Concidently with impaired activity of the miRNA pathway in the mutants, we observed an
 146 over-accumulation of several miRNA-targeted mRNAs in the single mutants and *carp9-* 147 *1/carp9-2* compound heterozygous plants (Figure 2D, Supplemental Figure S2D and S2E). As
 148 expected, the *35S:CARP9* plants showed reduced miRNA-target transcripts to WT levels
 149 (Figure 2D, and Supplemental Figure S2D).

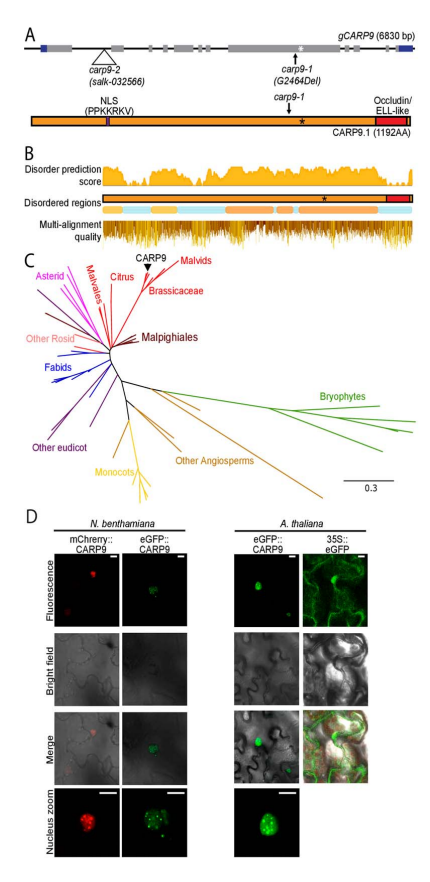
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151 *CARP9* encodes an intrinsically disordered protein of unknown function conserved among 152 land plants

153 CARP9 encodes an 1192 amino acid long protein of unknown functions that includes a 154 predicted nuclear localization signal (NLS) at position 238-245 (Figure 3A). A single isoform is 155 reported in the TAIR10 genome version (Arabidopsis Genome, 2000), while a second splicing 156 variant is annotated in the Araport11 release (Cheng et al., 2017), but we were not able to detect 157 it by RT-PCR and represent less than the 2% of splicing junctions in RNA-seq experiments 158 (Supplemental Figure S3A). An analysis of the protein sequence searching for conserved 159 homology domains revealed only the presence of an Occludin/ELL-like domain (pfam E-value 160 score: 8.71e-27, (El-Gebali et al., 2019) in the C-terminal region, covering 8.4% of the total 161 protein length (amino acids1088 to 1187, Figure 3A). Most of the eukaryotic RNA polymerase 162 II Elongation Factors (ELL) contain this kind of domains and are thought to mediate protein 163 interactions (Li et al., 2005; Van Itallie and Anderson, 2018), but they are mostly unstudied in plants yet (Shilatifard et al., 1996). Besides this domain, a structure prediction indicated that 164 165 CARP9 is an intrinsically disordered protein with 61.1% of disordered regions (Figure 3B) 166 (Potenza et al., 2015).



167 Interestingly, proteins with long disordered regions followed by an occluding domain appeared



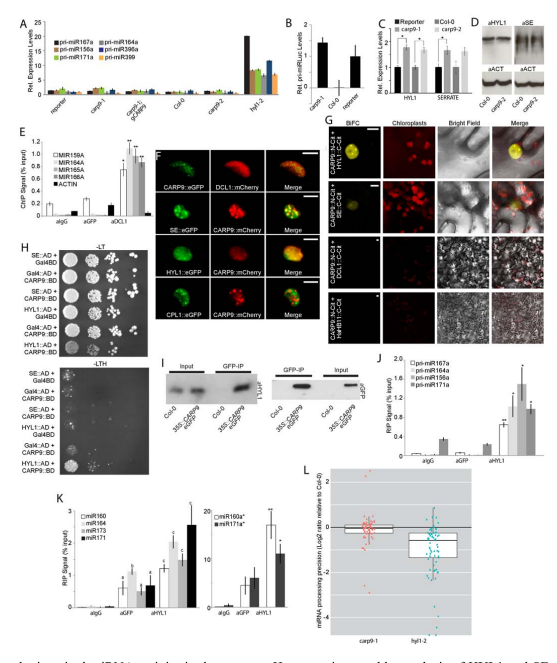
to be exclusive of plant species, according to the InterPro database (Mitchell et al., 2019). In this

sense, we performed a BLASTp search of CARP9 in plant genomes. This search yielded that 169 170 this gene is well conserved among embryophytes species (Figure 3C). We could also track 171 CARP9 orthologs in all embryophytes, including bryophytes, while they are absent in algae species, suggesting that CARP9 has evolved within land plants. Using the Maximum Likelihood 172 173 method, we performed a phylogenetic analysis with these proteins (Figure 3C and Supplemental 174 Figure S3B). The obtained phylogenetic tree is entirely consistent with the evolution of plant 175 species and we did not identify conserved duplication events for this gene. In most of the 176 species, CARP9 orthologs correspond to a single-copy gene. Such a single copy of the gene 177 may explain why some of the studied alleles are not viable in homozygosis as redundancy may 178 be absent for this gene. However, we cannot exclude partial functional redundancy with other 179 IDPs. Proteins alignment showed that the conserved regions include both the Occludin/ELL 180 domain and the disordered region (Figure 3B). Moreover, CARP9 paralogs are also predicted as 181 disordered in all species. This suggests that both parts of the protein, the Occludin/ELL domain, 182 and the disordered region, might be necessary for their molecular function. IDPs are important 183 for molecular recognition (Tompa et al., 2015) and work as a scaffold for many molecular 184 interactions (Cortese et al., 2008). They are also particularly abundant in cellular membrane-less 185 organelles such as nuclear bodies (Uversky et al., 2015). Furthermore, intrinsically disordered 186 regions are essential in protein-RNA interactions, mediating both specific and non-specific 187 interactions (Varadi et al., 2015; Jarvelin et al., 2016). Recently, Arabidopsis FLL2, also a 188 disordered protein, was reported to be located in nuclear bodies promoting liquid-liquid phase separation (Fang et al., 2019). The fusion of CARP9 with the fluorescent proteins eGFP or 189 190 mCherry, followed by confocal microscopy, revealed that the protein localizes in the nucleus, 191 and particularly in nuclear bodies of unknown nature (Figure 3D and 3E). Such nuclear entities 192 could perfectly reflect specific liquid organelles or the well-known dicing-bodies, subnuclear 193 speckles of miRNA processing (Fang and Spector, 2007).

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195 CARP9 interacts with HYL1 but does not affect miRNA processing

196 The reduced miRNA levels in carp9 mutants and the presence of an ocluddin/ELL domain in 197 CARP9 led us to propose a putative role as an elongation factor controlling the transcription of 198 genes encoding miRNA biogenesis factors or even of MIRNA genes. To test this hypothesis, we 199 first quantified by RT-qPCR transcript levels of pri-miRNAs and genes encoding core 200 components of the miRNA biogenesis machinery. In contrast to hyl1-2 mutants, used as a 201 positive control for defective processing where pri-miRNAs over accumulate, carp9-1 and 202 carp9-2 presented normal levels of the miRNA precursors as well as pri-miRLUC (Figure 4A 203 and 4B). Among all tested genes encoding miRNA related proteins, we observed a subtle but 204 significant increase in HYL1 and SE transcript levels (Figure 4C). This result is opposed to our hypothesis of CARP9 acting as an elongator factor but compatible with a feedback response to 205



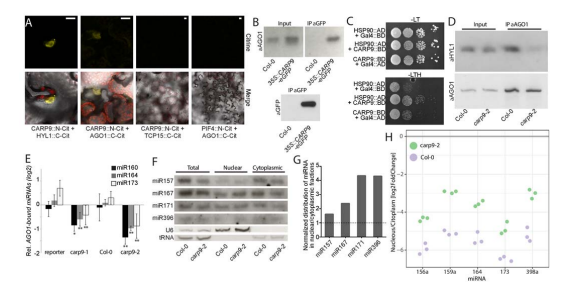
the impaired miRNA activity in the mutants. However, immunoblot analysis of HYL1 and SE 206 207 showed that these protein levels remained stable in the mutants (Figure 4D). It has also been 208 shown that DCL1 is recruited to MIRNA genes by the action of elongation factors (Fang et al., 209 2015). Thus, it is possible that CARP9 does not affect MIRNA genes transcription itself but allows the recruitment of the processing machinery to the loci. We tested the capacity of 210 CARP9 to interact with MIRNA loci by ChIP-qPCR using an eGFP-tagged version of the 211 protein. Unlike DCL1, used as a positive control, CARP9 did not appear associated with the 212 213 tested MIRNA loci (Figure 4E). Altogether, these results suggested that CARP9 is not acting as 214 an elongator factor to control miRNA activity. Alternatively, it is also possible that CARP9 is

directly implicated in the processing of pri-miRNA, based on its particular sub-nuclear 215 216 localization in speckles similar to dicing bodies (Figure 3D and 3E). Interestingly, transient 217 expression of fluorescent-tagged CARP9 followed by confocal microscopy showed 218 colocalization with SE and HYL1 in the same nuclear speckles (Figure 4F). However, it did not, 219 or only partially, colocalized with DCL1 and CTD-PHOSPHATASE-LIKE 1 (CPL1), which 220 de-phosphorylate HYL1 to enhance pri-miRNA processing (Figure 4F). Coincidently, 221 bimolecular fluorescence complementation (BiFC) assays showed fluorescent reconstitution 222 when CARP9::N-CITRINE was confronted with HYL1 and SE fused to the C-CITRINE, but not 223 with DCL1 (Figure 4G and Supplemental Figure S4). We used the nuclear transcription factor 224 HaHB11 (Cabello et al., 2016) as a negative control for the assay. Since colocalization and 225 BiFC assays indicate protein proximity but not necessarily interactions, we performed a yeast two-hybrid assay (Y2H) to evaluate direct protein-protein interaction between CARP9, HYL1, 226 227 and SE. Only HYL1 among the tested proteins was able to interact with CARP9 (Figure 4H). 228 This interaction was further confirmed by a Co-IP experiment using eGFP tagged CARP9 and 229 an antibody against the endogenous HYL1 (Figure 4I). The fact that HYL1 is known to interact 230 with SE (Lobbes et al., 2006), may explain the positive signal detected between SE and CARP9 231 in BiFC assays. RIP-qPCR assays using a GFP-tagged CARP9 showed no association of this protein with pri-miRNAs, which appeared associated to HYL1 (Figure 4J). Conversely, we 232 233 detected abundant mature miRNAs associated with CARP9 in the IP samples (Figure 4K). This 234 association of CARP9 with mature miRNAs is probably through its interaction with HYL1, 235 which is known to interact with miRNAs (Yang et al., 2010). The fact that HYL1 binds both 236 pri-miRNA and mature miRNAs (Figure 4J and 4K, and Yang et al., 2010) but CARP9 only 237 appeared to interact with mature miRNA suggests that CARP9-HYL1 interaction occurs after 238 pri-miRNA processing. Supporting this scenario, miRNA-processing precision, a feature 239 impaired in hyll-2 mutants and associated to its function as DCL1 accessory protein, was not 240 affected in *carp9-1* plants (Figure 4L). MiRNA processing precision was calculated by scoring 241 the ratio of total miRNA-matched small RNA to the pool of imprecisely processed small RNAs, 242 defined as those only partially matching the mature miRNA sequence. A late action of CARP9 243 in the pathway is also in line with the lack of interaction of CARP9 with DCL1 and CPL1, 244 which are expected to act on early stages of the miRNA processing. This is also in agreement 245 with the unchanged levels of pri-miRNAs in the mutant compared to WT (Figure 4A), which 246 tend to accumulate in mutants impair in pri-miRNA processing (Ben Chaabane et al., 2013), 247 suggesting that CARP9 is not active during miRNA biogenesis.

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249 CARP9 interacts with AGO1, affects its stability and miRNA loading

The potential post-miRNA-processing interaction of CARP9 with HYL1, and its nuclear localization, suggested that this protein might affect one of the nuclear steps of the miRNA

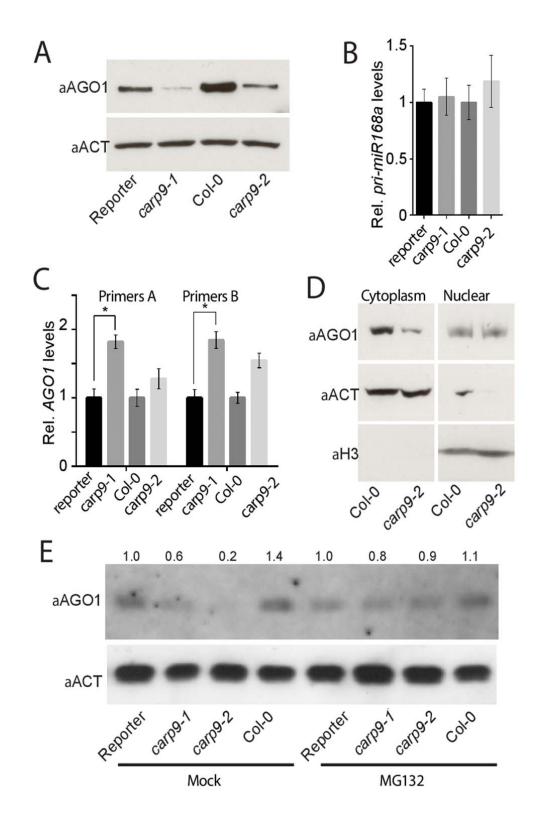


pathway downstream of biogenesis. This could be the case of the recently reported nuclear 252 253 loading of some miRNAs into AGO1 (Bologna et al., 2018). To explore this possibility, we first tested the capacity of CARP9 to interact with AGO1 by BiFC and Co-IP assays. Both 254 experiments showed that CARP9 can interact with AGO1, and this interaction occurs in the 255 256 nucleus, as observed in the microscopy assays (Figure 5A, 5B, Supplemental Figure S5A, and S5B). The nuclear transcription factors TCP15 and PIF4 (Ferrero et al., 2019) were used as 257 258 negative controls for the assay. Interestingly, Y2H assays showed that CARP9 also interacts 259 with the known AGO1-partner HSP90, which can locate in the nucleus (Bologna et al., 2018) 260 (Figure 5C). Contrary, BiFC assays revealed that CARP9 do not interact with SQUINT, which according to our data, interacts with AGO1 exclusively in the cytoplasm (Supplemental Figure 261 262 S5C). As CARP9 interacts with HYL1, and this protein was shown to interact with AGO1 in the nucleus (Fang and Spector, 2007), it is possible that CARP9, as many intrinsically disordered 263 264 proteins, acts as a scaffold for AGO1-HYL1 interaction. To test this hypothesis, we performed 265 AGO1-HYL1 co-IP experiments in WT and carp9-2 mutants treated with the proteasome 266 inhibitor MG132. MG132 treatments equalize AGO1 levels, which are altered in carp9 mutants 267 (see below), allowing us a correct interpretation of the results. The experiment showed that 268 AGO1-HYL1 interaction is partially impaired in the mutant background suggesting that CARP9 269 facilitates the formation of a post-processing miRNA complex containing AGO1, HYL1, and 270 likely HSP90 (Figure 5D and Supplemental Figure S5D).

Altogether this data suggests that CARP9 may act stabilizing HYL1-AGO1 interaction and perhaps facilitating the loading of miRNAs into AGO1. To test whether AGO1 loading is affected in *carp9* mutants, we immunoprecipitated AGO1 in WT and mutant plants treated with MG132 and scored the levels of associated miRNAs by RT-qPCR (RIP-qPCR). We observed a significant depletion of AGO1-associated miRNAs in the mutants when compared to WT 276 plants, except for miR822 that is canonically loaded in the cytoplasm ((Rajagopalan et al., 277 2006), Figure 5E, Supplemental Figure S5E and S5F), suggesting that CARP9 is participating in 278 the nuclear loading of miRNAs into AGO1 or at least stabilizing AGO1-miRNA association. 279 This observation may explain the mild reduction in miRNA accumulation as most likely *carp9* 280 mutation only affect the portion of miRNAs loaded in the nucleus but not the cytoplasmic 281 loading. The impaired loading of miRNAs into AGO1 in carp9 mutants was also evident when 282 we measured mature miRNAs accumulation in nuclear and cytoplasmic fractions. RNA blot and 283 RT-qPCR assays revealed that the nuclear portion of mature miRNAs is enriched in *carp9-2*, 284 compatible with deficient AGO1 nuclear loading, and subsequent cytoplasm exportation of this 285 miRNA fraction (Figure 5F, 5G, 5H, and Supplemental Figure S5G).

Intriguingly, we detected increased levels of AGO1 in the plants overexpressing eGFP:CARP9 286 287 construct (Figure 5B and Supplemental Figure S5B) and a reduction in carp9-1 and carp9-2 288 mutants when compared to their respective controls (Figure 6A). Such changes in AGO1 levels 289 cannot be attributed to the known regulation of AGO1 by miR168 (Vaucheret et al., 2004), as 290 we did not observe an increment of miR168 or MIR168 transcript levels (Figure 1E, 6B, and 291 Supplemental Table S1). In contrast to AGO1 protein, AGO1 transcripts levels, measured by 292 RT-qPCR with primers designed toward the 3'end of the mRNA (primers set A) or flanking the miR168 target site (primers set B), were higher in the mutants than in control plants probably as 293 294 a feedback response to the protein reduction or as a consequence of the lower miR168 activity 295 (Figure 6C, Supplemental Table S1). Nuclear/cytoplasm protein fractionation of WT and carp9-296 2 plants showed that the nuclear fraction of AGO1 is not reduced in the mutant, which is 297 consistent with a possible cytoplasmic proteolytic degradation of AGO1 (Figure 6D). Treatment 298 of the mutant plants with the proteasome inhibitor MG132 partially reverted AGO1 reduction 299 confirming that this protein is being degraded in the mutants (Figure 6E and Supplemental 300 Figure S6). Such reduced levels of AGO1 can be the cause, but also the consequence of the deficient miRNA loading observed in the mutants as unloaded AGO1 could become unstable 301 302 and get degraded. Nevertheless, the most parsimonious explanation to our observations is that 303 CARP9 is acting as a scaffold protein promoting the formation of a nuclear post-processing 304 miRNA complex containing at least AGO1, HYL1 and HSP90. In the absence of CARP9 this 305 hypothetical complex would be disrupted thus altering AGO1 stability, the proper miRNA 306 loading and therefore the stability of miRNA duplex itself.

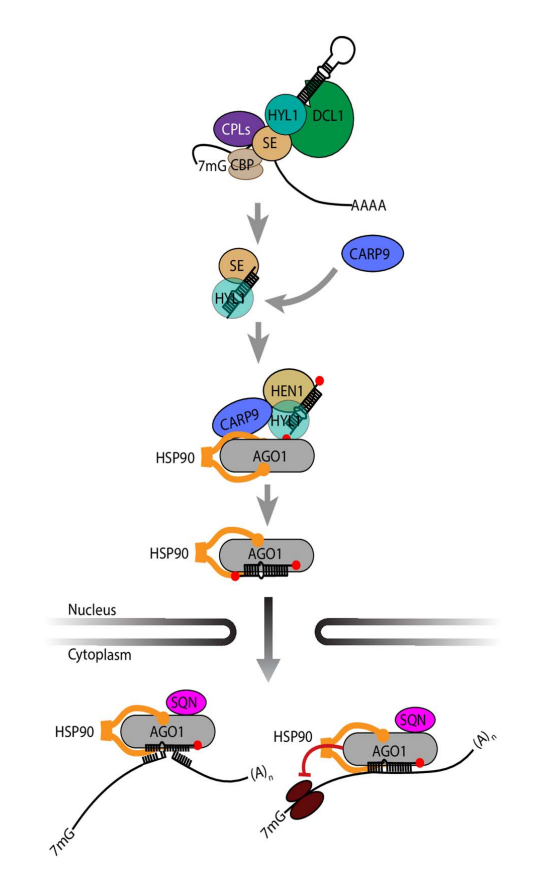
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309 DISCUSSION

During the last years, our knowledge of the miRNA pathway in plants has grown exponentially. 310 311 New functions were assigned to well-known proteins of the pathway, such as the chromatin 312 association of AGO1 and DCL1 (Fang et al., 2015; Dolata et al., 2016; Liu et al., 2018), the role of these same proteins in DNA damage repair (Wei et al., 2012; Schalk et al., 2017), the 313 314 transcriptional and epigenetic regulation of genes by SE (Ma et al., 2018; Speth et al., 2018), 315 and the independence of some pri-miRNA of HYL1 at low temperatures (Re et al., 2019), 316 among other examples. A recent report demonstrated that a portion of miRNAs can be directly 317 loaded into AGO1 inside the nucleus and exported as a complex (Bologna et al., 2018). This 318 evidence contrasts the previous conception that miRNAs were exported to the cytoplasm by 319 HASTY in order to be loaded into AGO1. The functional relevance of AGO1 nuclear loading is 320 becoming important since AGO1 involvement in several nuclear processes were recently 321 described (Dolata et al., 2016; Schalk et al., 2017; Liu et al., 2018). However, it remains unknown whether this nuclear loading of AGO1 is a passive process or if components of the 322 323 processing machinery assist it. It has been previously shown that proper loading of AGO1, 324 particularly the miRNA strand selection, is impaired in HYL1 mutants, suggesting that this 325 processing factor might participate actively during AGO1 loading (Eamens et al., 2009; 326 Manavella et al., 2012). Interestingly HEN1, which 2'-O-methylates mature miRNA, interacts 327 with HYL1, but not with the processing factor SE (Baranauske et al., 2015). This potentially 328 also place HEN1 in an hypothetical post-processing complex together with AGO1 and HYL1. 329 The capacity of HYL1 to efficiently bind mature miRNAs in vitro (Yang et al., 2010), suggests 330 that, after processing, mature miRNA might remain bound to HYL1 until loaded into AGO1. In this context CARP9 appeared to act as a nexus among these proteins facilitating the proper 331 function of such post-processing complex (Figure 7). 332

333 In this study, we describe CARP9, an intrinsically disordered protein conserved among land 334 plants, as a nuclear protein participating in miRNA activity. Interestingly, we found that CARP9 335 interacts with HYL1 and mature miRNAs but not with the miRNA processing machinery or 336 miRNA precursors, suggesting that this protein functions after the pri-miRNA processing steps. 337 This idea is strongly supported by a miRNA-processing precision analysis that showed that 338 carp9-1 mutants, opposite to hyl1-2, displayed a normal processing activity (Figure 4L). This 339 discovery also supports previous reports proposing that HYL1 remains bound to the mature 340 miRNA duplexes once they are produced (Figure 5, (Baranauske et al., 2015). In this context, it 341 is possible that after pri-miRNA processing, HYL1 transfers the mature miRNA duplexes to 342 AGO1 in the nuclear speckles where they colocalize, a scenario compatible with the AGO1 343 loading defects in hyl1 mutants (Fang and Spector, 2007; Eamens et al., 2009; Manavella et al., 344 2012). Furthermore, HYL1 may facilitate AGO1 loading in the cytoplasm, where it is also 345 located without reported functions (Cho et al., 2014; Achkar et al., 2018), . However, and to the 346 best of our knowledge, there is no evidence of direct interaction between HYL1 and AGO1 in



347 the cytoplasm. Besides its interaction with HYL1, we found that CARP9 also interacts with

AGO1 contributing to the miRNA loading process (Figure 5). It is common that IDPs, as 348 349 CARP9, act as chaperones assisting the folding of other proteins, allowing interactions and even 350 preventing their aggregation (Tompa and Kovacs, 2010). In such a scenario, CARP9 could act as a scaffold promoting AGO1-HYL1 interaction, providing stability to AGO1 and to the 351 352 mature miRNA duplexes and consequently promoting loading of miRNA into AGO1. As HYL1 353 is also found in the cell cytoplasm, where it undergoes protein degradation during the night 354 (Cho et al., 2014; Achkar et al., 2018), it is possible that this RNA binding protein escort AGO1 355 during miRNA nuclear export, a process that would exclude CARP9 as we found it to be 356 exclusively nuclear.

It was interesting to notice that *carp9* mutant alleles displayed reduced AGO1-bound miRNAs, 357 358 miRNA levels, and morphological defects compatible with miRNA deficient mutants. However, 359 these defects are milder than the observed in mutants of core proteins of the pathway, such as 360 HYL1 and AGO1. Redundancy with proteins with high sequence similarity is unlikely to 361 explain this observation as CARP9 appeared as a single gene in most of the genomes of plants 362 considered here, including Arabidopsis and by the fact that plants homozygous for *carp9* null 363 alleles appeared not to be viable. It is more likely that a reduced activity of CARP9 in the 364 mutant alleles explains the observations, as the premature stop codon in *capr9-1* would allow a 365 large portion of the protein to be translated and the intronic T-DNA insertion of carp9-2 366 possibly represents a knock-down allele as indicated by the reduced CARP9 transcripts in this 367 mutant (Supplemental Figure S3). Besides, as CARP9 is likely to only affect the nuclear loading 368 of miRNAs, the large pool of cytoplasmic loaded miRNAs are not expected to be affected, thus restricting the effect of the protein to only a subpopulation of miRNAs. 369

It is notable the high degree of conservation of CARP9 among land plants suggesting a crucial 370 371 role of this protein for plant homeostasis. The extraordinary level of amino acid conservation in 372 the intrinsically disordered regions, which commonly tend to diverge rapidly during evolution 373 rapidly, reinforce the idea of an evolutionary pressure to conserve this protein. Such inference 374 might explain why we failed to identify homozygous T-DNA mutants in the coding sequence of 375 this gene. The identification of strong but viable loss-of-function alleles of CARP9 could help to 376 dimension the importance of this protein in plant development and miRNA processing in the 377 future.

CARP9 represents a new component of the miRNA pathway that links the post-processing machinery with the miRNA effector complex. Several questions remain open; for example, HEN1 was shown to interact with HYL1 post-processing (Baranauske et al., 2015), opening the question of whether AGO1 is loaded with already methylated miRNA or if such process occurs after loading for the fraction of nuclear-loaded miRNAs. EMA1 and TRN1 were also shown to interact with AGO1 modulating miRNAs loading (Wang et al., 2011; Cui et al., 2016). However, it remains to be addressed whether these proteins also participate in the nuclear AGO1-loading process interacting with HYL1 and CARP9 or have a different role. Similarly, we showed that CARP9 interacts with AGO1 and HYL1 to facilitate loading, but the biochemical activity of CARP9 during this process is unclear. Perhaps the most intriguing question lays in the observed destabilization of AGO1 in *carp9* mutants. How this process is triggered and accomplished are questions we still need to answer. In particular, it will be essential to understand how AGO1 degradation, CARP9-HYL1 interaction, and miRNA loading are interconnected.

392 We found three significant effects of *carp9* mutations; a mild reduction in miRNAs levels, a 393 reduction in AGO1 loading, and a reduction in AGO1 protein levels. Interestingly, each of these 394 observations can be explained in light of the other ones. In this sense, miRNA reduction and 395 impaired AGO1 loading can be the consequence of AGO1 destabilization. It is also possible that 396 impaired AGO1 loading destabilizes AGO1 and miRNAs. Finally, if CARP9 is directly involved in stabilizing the miRNA duplex after processing, we could also expect to see impaired 397 398 AGO1 loading and stability. Perhaps the first scenario is the less likely as AGO1 loading is 399 impaired in *carp9* even when MG132 treatments block protein degradation. Nevertheless, we 400 cannot exclude any of these possibilities, and further studies will be necessary to discriminate 401 these alternatives and to understand the role of CARP9 in HYL1-AGO1 crosstalk precisely.

402

403 MATERIALS AND METHODS

404 Plant Material and Growth Conditions

405 Arabidopsis (Arabidopsis thaliana) ecotype Columbia (Col-0), reporter and mutant plants were 406 grown at 23°C on soil in long-day photoperiod (LD, 16 hours of light/8 hours of dark). 407 Alternatively, plants were grown on plates containing 2.2 g/L of Murashige and Skoog (MS) 408 medium (pH 5.7) and 0.6% (w/v) agar in LD conditions. Seeds were disinfected with 10% (v/v) 409 bleach and 0.1% (w/v) SDS and stratified in 0.1% (w/v) agar for 3 days at 4° C before sowing. 410 Mutant lines carp9-2 (SALK 032566); capr9-3 (SALK 060892), carp9-4 (SALK 044585), 411 carp9-5 (WiscDsLox358B04) and hyl1-2 (SALK 064863) were obtained from ABRC stock 412 center. Col-0 miRNA activity reporter lines (reporter) used for EMS mutagenesis, and thus as 413 capr9-1 control, were previously described (Manavella et al., 2012). Transgenic lines were grown on MS plates with 50 mg/mL kanamycin. 414

415

416 DNA Constructs and Plant Transformation

The *CARP9*, *HSP90*, *SQN* coding region sequences (CDS), with or without stop codon, were
amplified and cloned into pEntr/D-TOPO or pCR8GW-TOPO entry vector (Thermo Fisher
Scientific). The *CARP9* promoter region (2558 bp upstream the transcription start site) was

amplified by PCR, fused to CARP9 cDNA fragment by PCR, and cloned into pEntr/D-TOPO 420 421 entry vector. EGFP and mCherry fusion constructs were obtained by recombination of the entry 422 clones into modified pGREEN vectors under the control of the CaMV 35S promoter. An untagged cDNA copy of CARP9 under a 35S promoter was used to rescue carp9-1 mutants. 423 424 Yeast two-hybrid constructs were obtained by cloning the specific cDNAs into pEntr/D-TOPO, 425 followed by recombination into the pDEST32 or pDEST22 vectors (Life Technologies). Refer 426 to Supplemental Table S2 for a detailed list of constructs used in this work. Arabidopsis 427 transgenic seedlings were selected using 50 mg/ml kanamycin on plates. At least 15 428 independent pooled T1 seedlings were used for quantitative measurement of transgenic lines. 429 Transient infiltration of *Nicotiana benthamiana* leaves was performed as described previously 430 (de Felippes and Weigel, 2010). We were unable to directly transform *carp9* mutants by floral 431 dip. Thus all carp9 transgenic plants were obtained by crossing the mutants with transgenic Col-0 plants and then recover the homozygosis on the mutant alleles. In the case of the experiments 432 performed with overexpression lines we only used plants with low expression levels of CARP9 433 434 and pooled lines to minimize variability.

435

436 RNA Analysis

437 Total RNA was extracted from 15-days-old plants using TRIzol reagent (Life Technologies). For RNA blots, 1-5 μ g of total RNA were resolved in 17 % (v/v) polyacrylamide gels under 438 439 denaturing conditions (7 M urea) and then transferred to HyBond-N+ charged nylon membranes 440 (Amersham) by semidry electroblotting (Tomassi et al., 2017). RNA was covalently fixed to 441 membranes in a UV Crosslinker. Membranes were hybridized overnight with DNA 442 oligonucleotide probes labeled with second-generation DIG Oligonucleotide 3'-End Labeling Kit (Roche); the signal was detected using CSPD ready-to-use solution, by exposure to 443 444 Amersham hyperfilm ECL (GE Healthcare Life Sciences). ImageJ was used to analyze the band 445 intensity of small RNA blots as integrated pixel density, using the intensity of U6 bands to 446 normalize sample loading. Reverse transcriptase reactions were performed using 1 µg of DNaseI-treated total RNA (Thermo Fisher Scientific) using the RevertAid RT Reverse 447 448 Transcription Kit (Thermo Fisher Scientific). Quantitative RT-qPCRs, were performed using 449 biological replicates of pooled three independent seedlings. and ACTIN2/8 450 (At3g18780/At1g49240) were used as a housekeeping loading control. Stem-Loop RT-qPCRs for miRNAs quantification were performed as previously described (Kramer, 2011). Averages 451 from biological triplicates and SE were calculated from $2^{-\Delta\Delta Ct}$ values, and the error displayed as 452 453 two times SM. Each replicate was treated as independent samples for statistical analysis. 454 Statistical differences between samples were determined by an unpaired, two-tailed, t-test 455 analysis and corrected with Holm-Sidak method for multiple pair comparisons. All 456 quantifications were repeated twice in independent experiments. See Supplemental Table S3 for

457 oligonucleotide primer and probe details.

458

459 Small RNA sequencing

460 Small RNA libraries were prepared as indicated by the TruSeq small RNA library prep kit (Illumina) using biological triplicates of the reporter lines and hyl1-2 mutants and duplicates of 461 462 carp9-1 mutants. 50 ng of small RNAs purified with the ZR small-RNA PAGE Recovery Kit 463 (Zymo Research) were used as input for the library preparation. Small RNA libraries size selections were performed using the BluePippin System (SAGE Science). Single-end Illumina 464 465 sequencing was performed with a HiSeg3000 apparatus. Small RNA reads were first processed 466 to remove 3' adapters using cutadapt (version 1.9.1) and then mapped using bowtie (version 467 1.1.2). Reproducibility was tested by computing the Spearman correlation of the miRNA counts 468 per million between all samples, converting this correlation to a distance (1-Spearman Rho) and 469 performing a hierarchical clustering of the samples, with the "complete" agglomeration method, 470 to show the degree of similarity between them. The references used were the databases for 471 hairpin and mature miRNAs for A. thaliana from miRBase (release 21), in the latter mature 472 miRNAs with identical sequences were collapsed into single miRNAs. Additionally, reads were 473 mapped to the A. thaliana genome using the same software. For the differential expression 474 analysis of the miRNAs, only reads mapping to the full-length mature miRNAs were 475 considered, and primary alignments of reads mapping to the sense strand were counted (filtering 476 with "samtools view-F 272"). Counts per miRNA were used as input for baySeq (version 2.8.0) 477 to perform the differential expression analysis. For this, miRNAs with low expression levels 478 (less than 10 counts in all samples) were discarded, and size factors were set according to the 479 total number of reads mapping to the genome for each sample. Graphics and statistical analyses 480 were performed in the R statistical programming environment (R Core Team, 2018) with the 481 ggplot package. MiRNA processing precision was calculated by scoring the ratio of total 482 miRNA-matched small RNA to the pool of imprecisely processed small RNAs, defined as those 483 only partially matching the mature miRNA sequence. All data reported in this paper is available 484 at the European Nucleotide Archive (ENA), PRJEB37499.

485

486 **Protein Analysis**

For immunoblot analysis, proteins were extracted from 15-days-old pooled plants (n=5) with
100 μl extraction buffer (50 mM Tris pH 7.5, 150 mM NaCl, 1 mM EDTA, 10% (v/v) Glycerol,
1 mM DTT, and one tablet Complete Protease Inhibitor Cocktail [Roche]) per 100 mg of grind

490 tissue. Proteins were resolved in 8% (w/v) SDS-PAGE gels (running buffer: 25 mM Tris-Base;

192 mM Glycin; 0.1% (w/v) SDS) and transferred using a standard wet tank blotting (blotting 491 492 buffer: 2.5 mM Tris-Base; 19.2 mM Glycine; 10% (v/v) methanol) to PVDF membrane 493 (Amersham). Thermo Scientific PageRuler Prestained Protein Ladder was used to determine the 494 molecular weight of the bands and to confirm transfer efficiency. AGO1, HYL1, and SE were 495 detected using a polyclonal antibody targeting the endogenous Arabidopsis protein (Agrisera 496 AS09527, AS06136, and AS09532; dilution 1:10,000 each). ACTIN 8 (Agrisera AS132640; 497 dilution 1:10,000), HISTONE3 (Agrisera AS10710; dilution 1:10,000), or coomassie blue 498 staining, were used as loading controls in different experiments. HBR-conjugated polyclonal 499 anti-rabbit IgG, (Agrisera AS09602; dilution 1:20,000), was used to detect primary antibodies. 500 Signal was detected using ECL Plus Western Blotting Substrate (Thermo Fisher Scientific). Experiments were repeated at least twice. For Co-IP assays, eGFP-CARP9 was 501 502 immunoprecipitated with an anti-GFP antibody (ABCAM ab290, dilution 1:1000) from samples extracted from transgenic A. thaliana flowers transformed with 35S::eGFP:CARP9 including 503 504 WT Col-0 flowers as a negative control, and AGO1 was immunoprecipitated with an anti-505 AGO1 antibody (Agrisera, dilution 1:1000) from samples extracted from pooled (n=5) 15-days-506 old carp9-1, carp9-2, reporter and WT Col-0 plants, using Sure BeadsTM Protein-A magnetic 507 beads (BioRad) following the manufacturer's instruction. eGFP:CARP was then detected in the 508 input and IP fraction by immunoblot using an anti-GFP antibody (ABCAM ab290, dilution 509 1:10000). HYL1 and AGO1 were detected in the input and IP fraction using the antibodies 510 previously described. Yeast two-hybrid assays were performed with the ProQuest Two-Hybrid System (Thermo Fisher Scientific). Selection medium containing 2.5 mM of 3-AT (3-amino-511 1,2,4-triazole) was used to detect interactions reducing autoactivation. CARP9 fusions to the N-512 terminal and C-terminal fragments of Citrine, eGFP or mCherry were used for BiFC assays, and 513 514 protein localization in transiently transformed N. benthamiana leaves. Fluorescent protein visualization and imaging were performed using a Leica TCS SP8 confocal microscope. The 515 excitation wavelengths were 488 nm, 514 nm, and 552 nm, and emission was collected at 500-516 517 530 nm, 525-560 nm, and 600-630 nm for eGFP, mCitrine, and mCherry, respectively. For 518 Luciferase activity assays, twelve 5mm discs cut from mature leaves were individually 519 embedded in 100 mM D-Luciferin potassium salt solution and bioluminescence quantified using 520 a Fluoroskan AscentTM FL plate Luminometer (Thermo Scientific). Means and SE was 521 calculated (n=12) and p-values of less than 0.05 in a two-tailed, unpaired t-test Holm-Sidak-522 corrected, were considered significant. Alternatively, Luciferase activity was detected in plants 523 sprayed with 100 mM D-Luciferin potassium salt solution using an Orca 2-BT cooled CCD 524 camera with 5 minutes integration time (Hamamatsu Photonics).

525

526 Nuclear-Cytoplasmic Fractionation

527 Nuclear/cytoplasmic fractionation was performed following a protocol previously described 528 (Wang et al., 2011). Briefly, samples of pooled 15-days-old plants (2.5 g) were ground on ice 529 with 2 ml/g of lysis buffer (20 mM Tris-HCl pH 7.5, 20 mM KCl, 2 mM EDTA, 2.5 mM MgCl₂, 25% (v/v) glycerol, 250 mM Sucrose, and 5 mM DTT) supplemented with 200 µM 530 531 PMSF (only for proteins fractionations). The homogenate was filtered through a layer of Miracloth, and the flow-through was centrifuged at 1,500 g for 10 minutes. The supernatant 532 533 (cytoplasmic fraction) was centrifuged at 10,000 g and 4° C for 10 minutes, and collected. The 534 pellet was washed 3-5 times with 5 mL of NRBT buffer (20 mM Tris-HCl pH 7.5, 25% (v/v) 535 glycerol, 2.5 mM MgCl₂, and 0.2% (v/v) Triton X-100) and then resuspended with 500 μ L of NRB2 (20 mM Tris-HCl pH 7.5, 0.25 M Sucrose, 10 mM MgCl₂, 0.5% (v/v) Triton X-100, and 536 537 5 mM β -mercaptoethanol) supplemented with Complete Protease Inhibitor Cocktail (Roche, 538 only for protein fractionations). The resuspension was carefully pipetted on top of 700 μ L of NRB3 buffer (20 mM Tris-HCl pH 7.5, 1.7 M Sucrose, 10 mM MgCl₂, 0.5% (v/v) Triton X-539 540 100, and 5 mM β -mercaptoethanol, supplemented with Complete Protease Inhibitor Cocktail). The obtained sucrose gradient was centrifuged at 16,000 g and 4° C for 1-3 minutes. For protein 541 542 extraction, the pellet was resuspended in 100 μ L lysis buffer and sonicated in a Bioruptor Pico 543 water bath (10 cycles of 30 seconds on/30 seconds off pulses at high intensity) (Diagenode). 544 After centrifugation at 16,000 g for 10 min at 4° C, the supernatant was collected as the nuclear 545 fraction. For RNA extraction, the pellet was resuspended in 200 µL lysis buffer, and then 1 mL 546 of TRIzol reagent (Life Technologies) was added, as to the cytoplasmic fraction, following then 547 by a standard protocol for RNA extraction. As a quality and loading controls for the fractionation, ACTIN 8 and a tRNA probe (for RNA quantifications) were used as cytoplasmic 548 markers, while Histone H3 and U6 RNA (for RNA quantifications) were used as nuclear 549 550 markers.

551

552 RNA and Chromatin Immunoprecipitation Assays (RIP and ChIP)

553 RIP assays to detect mature miRNA or pri-miRNAs bound to eGFP::CARP9, HYL1, and 554 AGO1 were performed using four independent biological replicates following a reported 555 protocol with a few modifications and scaled down to 50% of the volumes (Carbonell, 2017). 556 Immunoprecipitation was performed to 4 g of tissue (flowers) using anti-GFP (AS152987, 557 dilution 1:250), antiHYL1 (AS06136, dilution 1:500) and antiAGO1 (AS09527, dilution 1:500). 558 In the case of AGO1 RIP experiments, 15-days-old UV-crosslinked plants, grown on MS agar 559 plates and treated with MG132 for 24 hours, were used as starting material. In all cases, Sure BeadsTM Protein-A magnetic beads (BioRad) were used for the immunoprecipitation. RNA was 560 561 finally extracted from the IP fraction by a regular TRIzol extraction. Associated RNA 562 quantifications were performed as described in the "RNA analysis" section of material and

methods. Chromatin immunoprecipitation assays of eGFP::CARP9 and DCL1 associated loci 563 564 were performed using antiGFP (AS152987, dilution 1:250) and antiDCL1 (AS122102, dilution 565 1:200). We first performed nuclei enriched of samples obtained from 3 g of seedlings following the same protocol described above for cell fractioning assay. Extracted nuclei were then 566 567 resuspended in 500 µL Nuclei Lysis Buffer (50 mM Tris-HCl pH 8, 10 mM EDTA, 1 % (w/v) 568 SDS, 1 mM PMSF, 1 % (w/v) Complete Protease Inhibitors (Roche)). Chromatin was sheared 569 using a Bioruptor Pico (Diagenode; 10 cycles 30" ON, 30" OFF). After fragmentation, nuclear 570 debris was pelleted and the supernatant diluted ten-fold with ChIP Dilution Buffer (1.1 % (v/v) Triton X-100, 1.2 mM EDTA, 16.7 mM Tris-HCl pH 8, 167 mM NaCl). Immunoprecipitation 571 was carried out at 4 °C overnight with 100µl of washed Sure Beads[™] Protein-A magnetic beads 572 (BioRad) and the appropriate amount of antibody. After washing five times, the beads were 573 resuspended in 100 µl of TE, and 1 µl of 20 mg/ml Proteinase K added. Samples were incubated 574 at 43 °C for 1h, and the protease inactivated by incubation at 95 °C for 10 min. Samples were 575 576 centrifuged 1 min at maximum speed and 1.5 µl used for qPCR reactions. Negative controls 577 were performed with samples in which specific antibodies were not included. Values were 578 expressed as a % of the input signal for the same measured miRNA.

579

Phylogenetic analysisCARP9-like protein sequences of representative plant species were 580 581 retrieved from Phytozome (Goodstein et al., 2012) using AtCARP9 full amino acid sequence as 582 a query for BLASTP search with default parameters. Proteins sequences with low sequence 583 similarity were discarded (query coverage <30%, E-value>1.1-10). The resulting sequences 584 were aligned using the MAFFT G-INS-1 iterative method (Katoh and Standley, 2013) and 585 automatically trimmed using TrimAI webserver (Capella-Gutierrez et al., 2009) with 0.9 gap 586 threshold fraction. Splicing variants were manually removed. Phylogenetic analysis was 587 performed using the Maximum likelihood (ML) method with IQTree default parameters 588 (Trifinopoulos et al., 2016) using a Shimodaira–Hasegawalike approximate likelihood ratio test. 589 The consensus tree was obtained with all compatible groups and visualized using FigTree v1.4.3 590 (http://tree.bio.ed.ac.uk/software/figtree/).

591

592 CARP9 conservation analysis

To test conservation of CARP9 in algae species, protein sequences of AtCARP9 and *Marchantia polymorpha* CARP9 were used as a query for BLASTP search against NCBI nonredundant database. Domain architecture searches were made using InterProScan (Mitchell et al., 2019). Amino acid disorder score and regions were obtained from MobiDB (Piovesan et al., 2018) using full protein sequences. Alignment quality was computed with Jalview (Waterhouse et al., 2009) using the same alignment obtained for phylogenetic analysis. NLS

- 599 was predicted using cNLS Mapper (<u>http://nls-mapper.iab.keio.ac.jp</u>) with 7.0 as the cut-off 600 score.
- 601

602 Accession numbers

603 Sequence data from this article can be found in the GenBank data libraries under accession

604 numbers AT3G21290 (CARP9); AT1G09700 (HYL1); AT1G48410 (AGO1); (HSP90);

- 605 AT2G15790 (SQN); AT2G27100 (SE); AT1G01040 (DCL1); AT4G21670 (CPL1);
- 606 AT1G69690 (TCP15); AT2G43010 (PIF4); AT5G56010 (HSP90).
- 607

608 Supplemental data

- 609 Supplemental Figure S1. Phenotype of *carp9* mutants and rescued plants.
- 610 Supplemental Figure S2. Impact of CARP9 mutation on the miRNA pathway.
- 611 Supplemental Figure S3. Conservation of CARP9 among plants.
- 612 Supplemental Figure S4. CARP9 interacts with HYL1.
- 613 Supplemental Figure S5. CARP9 interacts with AGO1.
- 614 Supplemental Figure S6. CARP9 mutations impair AGO1 stability.
- 615 Supplemental Table S1. Differentially accumulated miRNAs as shown by small RNAseq data
- analysis of carp9-1 mutants.
- 617 Supplemental Table S2. Transgenes and plasmids.
- 618 Supplemental Table S3. DNA oligonucleotide primers and probes.
- 619

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- 628

629 FIGURE LEGENDS

- Figure 1. Characterization of *carp9* mutants. A, Gene structure of *CARP9* showing single
 nucleotide deletion of the *carp9-1* allele and T- DNA insertion sites in *carp9-2*, *carp9-3*, *carp9-3*
- 632 4, and carp9-5. Solid boxes and lines represent exons and introns, respectively; gray boxes
- 633 represent coding sequence while blue boxed shows 5'- and 3'-UTR regions. A white asterisk (*)
- 634 marks the position of the stop codon that results from the frameshift caused by the *carp9-1*

single nucleotide deletion (G2464Del). B, Phenotypic characterization of carp9 mutants, control 635 636 lines (reporter and Col-0), and *carp9-1* mutants complemented by the overexpression of the 637 CARP9 cDNA (carp9-1; Pro³⁵⁵:CARP9). 21-days-old plants, fully expanded siliques, and 40days-old plants are displayed. White bars on the right represent 1 cm. Plants were imaged 638 639 individually, digitally extracted, and mounted on a single black background panel to facilitate 640 comparison and observation. C, Analysis of the flowering time of control, carp9 mutant, and 641 complemented lines grown in long-day photoperiod as measured by the number of rosette 642 leaves or the number of days to bolting. Error bars show means \pm SE (n \geq 15). D, Bioluminescence activity, quantified by a luminometer, of 12 leave discs belonging to 20-days-643 old *carp9-1* mutants and reporter plants. Error bars show means \pm SE (n \geq 12). p-values of less 644 than 0.05 (*) in a two-tailed, unpaired t-test Holm-Sidak-corrected were considered significant. 645 646 E, Bioluminescence activity as measured with a CCD camera, in 20-days-old carp9-1 mutants 647 and reporter plants. Luminescence intensity is color scaled from low (blue) to high (white). Two pots containing 12 plants each were imaged individually, digitally extracted, mounted in a 648 649 single black background panel, and displayed in the figure. E, RNA blots for detecting 650 amiRLuc. U6 was used as a loading control. F, Phenotypic features of 18-days-old carp9 651 mutants, control lines, and carp9-1/capr9-2 compound heterozygous mutants. White bars on the 652 right represent 1 cm. Plants were imaged individually, digitally extracted, and mounted in a 653 single black background panel to facilitate comparison and observation.

Figure 2. CARP9 mutants present impaired miRNA activity. A, RNA blots for detecting 654 655 endogenous miRNAs. U6 was used as a loading control. The relative abundance of each 656 miRNA, indicated above each band, was calculated by measuring the band intensity using 657 ImageJ and relativized to the corresponding control plant (reporter for carp9-1 and 658 complemented plants, and Col-0 for carp9-2). B, MiRNA levels, as measured by RT-qPCR, in mutants and control lines. Error bars show means $\pm 2xSE$ (n \geq 4), p-values of less than 0.05 (*) in 659 660 a two-tailed, unpaired FDR-corrected t-test were considered significant. C, Mean expression 661 levels of individual miRNAs in *carp9-1* and *hyl1-2* plants relative to Col-0 plants. Horizontal 662 segments indicate the median of the expression levels. Each dot corresponds to a single miRNA 663 or collapsed miRNA family. Dark and light dots show differentially (FDR adjusted p-value < 664 0.05) and not-differentially accumulated miRNAs, respectively. D, Expression of miRNA 665 targets in control and mutant plants as measured by RT-qPCR. Error bars show means $\pm 2xSE$ $(n\geq 4)$, p-values of less than 0.05 (*) in a two-tailed, unpaired t-test Holm-Sidak-corrected were 666 667 considered significant.

Figure 3. Conservation analysis of CARP9 across the plant kingdom. A, On top, gene
structure of *CARP9* as shown in Figure 1A. On the bottom, the CARP9 protein structure, on
purple is marked a putative NSL signal and its amino acid sequence. The occluding/ELL-like

domain is marked in red. B, Top and middle, disordered score and regions in AtCARP9 amino 671 672 acid sequence according to MobiDB (Piovesan et al., 2018). Bottom, amino acid alignment 673 quality of CARP9-like genes using Jalview software (Waterhouse et al., 2009), positions are 674 based on AtCARP9 full sequence. C, Phylogeny of CARP9-like genes in embryophytes. The 675 unrooted consensus tree was generated using the Maximum Likelihood method. Colors 676 represent different lineages of plant species, referenced in the figure. AtCARP9 is highlighted 677 with a black arrow. Supplemental Figure S3B shows the fully annotated tree. D, Confocal 678 microscopy images showing the nuclear localization of eGFP- and mCherry- tagged versions of 679 CARP9 in N. benthamiana transiently transformed leaves (left) and stably transformed 680 Arabidopsis plants (right). Scale bars represent 5 µm.

681 Figure 4. CARP9 interacts with HYL1 and mature miRNAs but not with the miRNA 682 processing machinery. A, B, and C, Expression of pri-miRNA, pri-miRLuc, HYL1 and SE in 683 control and mutant plants as measured by RT-qPCR. D, HYL1, and SE quantification by 684 immunoblot in samples extracted from *carp9-2* and reporter plants. The detection of ACTIN 685 was used as a loading control. E, ChIP experiment using either anti-GFP, anti-DCL1, or anti-686 IgG antibodies in plants that express a GFP tagged version of CARP9 to detect MIRNAs loci 687 associated with the proteins. Primers used for the amplification are listed in the Supplemental Table S3 and based on a previous report (Fang et al., 2015). ACTIN gene was used as control 688 689 not targeted by CARP9 nor DCL1. F, Confocal microscopy images simultaneously showing the 690 localization of CARP9 with DCL1, SE, HYL1, and CPL1 in transiently transformed N. 691 benthamiana leaves. Scale bars represent 5 µm. G, BiFC assay in N. benthamiana cells showing 692 CARP9 interaction with HYL1 and SE. Negative interactions are displayed in a wider magnification to show the negative interactions better. Positive interactions in a wider 693 magnification are shown in Supplemental Figure S4. Scale bars represent 5 µm. H, Interaction 694 695 of CARP9 with HYL1, but not with SE, as detected by yeast two-hybrid assays. GAL4 696 activation domain (AD); GAL4 DNA binding domain (BD); -LT, medium without leucine and 697 tryptophan; -LTH, selective medium without leucine, tryptophan, and histidine. Each column 698 shows a 1:10 serial dilution. I, CARP9-HYL1 interaction detected by Co-IP assays. Leaves of A. thaliana plants transformed with Pro35S::CARP9-eGFP were immunoprecipitated using an 699 700 anti-GFP antibody. Interacting HYL1 was identified using an antibody targeting the endogenous 701 protein. J and K, pri-miRNAs (J) and mature miRNAs or miRNA*s. (K) associated with 702 CARP9, or HYL1, as quantified by RIP-RTqPCR in samples extracted from plants expressing a 703 GFP tagged version of CARP9 and immunoprecipitated with either an anti-GFP, anti-HYL1 or 704 anti-IgG antibodies. Values are given as a percentage to the qPCR signal detected in the input 705 samples. (L) Precisely processed miRNA reads at all highly expressed MIRNA loci. Each dot 706 represents an individual miRNA; horizontal black bars indicate medians. miRNA levels in all samples are expressed as a ratio to the precisely processed miRNAs in Col-0 plants grown at 23°C in LD photoperiod. In all panels involving RT-qPCR experiments, error bars show means $\pm 2xSE$ (n≥4), p-values of less than 0.05 (*) or 0.01 (**) in a two-tailed, unpaired t-test were considered significant.

711 Figure 5. CARP9 interacts with AGO1 to modulate its nuclear miRNA-loading. A, BiFC 712 assay in Nicotiana benthamiana cells showing CAP9 interaction with AGO1, and with HYL1 as 713 controls. The nuclear transcription factors PIF4 and TCP15 were used as negative controls and 714 displayed in a wider magnification. Scale bars represent 5 µm. B, Co-IP assays. Protein samples extracted from Col-0 WT plants or plants transformed with a 35S::CARP9-eGFP were 715 716 immunoprecipitated using an anti-GFP antibody, AGO1-CARP9 interaction was then detected 717 using an anti-AGO1 antibody. C, Interaction of CARP9 with HSP90 as detected by yeast two-718 hybrid assays. GAL4 activation domain (AD); GAL4 DNA binding domain (BD); -LT, 719 medium without leucine and tryptophan; -LTH, selective medium without leucine, tryptophan, 720 and histidine. Each column shows a 1:10 serial dilution. D, Co-IP assays. Protein samples 721 extracted from Col-0 WT or carp9-2 plants were immunoprecipitated using an anti AGO1 722 antibody, AGO1-HYL1 interaction was then detected using an anti-HYL1 antibody. AGO1 was 723 detected to test IP efficiency. E, Relative amount of mature miRNA bound to AGO1 as 724 measured by Stem-loop RT-qPCR of samples immunoprecipitated using an anti-AGO1 725 antibody. Co-IPed miRNAs were normalized to the levels of the same miRNA in the input 726 samples. For both mutants, the miRNA levels were then expressed as relative to their 727 corresponding control. Error bars show means $\pm 2xSE$ (n=4), p-values of less than 0.05 (*) or 0.01 (**) in a two-tailed, unpaired t-test were considered significant. No-antibody samples and 728 729 ago1-36 mutant plants were used as negative controls for the IP experiment, not showing 730 detectable signal in the assayed conditions. F, RNA blots for detecting miRNAs in different cell 731 fractions. Quantification of U6 and tRNAs were used as a loading control and to monitor the purity of the nuclear/cytoplasmic fractions. G, Quantification of the miRNA distribution 732 733 measured in (F). Band intensity was quantified by ImageJ and normalized by the corresponding 734 loading control. Distributions of miRNAs in the nuclear/cytoplasmic fractions were then 735 expressed as relative to Col-0 (marked as a dashed line). H. Quantification of the miRNAs 736 distribution in nucleus vs. cytoplasm fractions as measured by RT-qPCR. Each dot represents an 737 independent replicate. Significant differences were tested with an Anova test: between miRNAs 738 p-value = 0.00484; between genotypes p-value = 0.00013.

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Figure 6. AGO1 stability is compromised in *CARP9* mutants. A, AGO1 levels quantified by
immunoblot in samples extracted from *carp9* mutants and control plants. Levels of ACTIN were
measured as a loading control. B and C, Expression of pri-miRNA168 (B), and *AGO1* (C) as

measured by RT-qPCR. AGO1 transcript levels were measured using two sets of primers; a pair amplifying the 3'-end of the transcript (Primers A) and a pair flanking the miR168 recognition site in the AGO1 mRNA (Primers B). Error bars show means $\pm 2xSE$ (n=4), p-values of less than 0.05 (*) in a two-tailed, unpaired t-test were considered significant. D, AGO1 levels, as measured by immunoblots, in cytoplasmic or nuclear cell fractionated samples. ACTIN and Histone 3 (H3) were used to verify the purity of the fractions. E, Immunoblot quantification of AGO1 levels in mutant and control plants treated with the proteasome inhibitor MG132.

751 Figure 7. A model for the role of CARP9 as a scaffold in a post-pri-miRNA processing and

nuclear AGO1 loading complex. After the processing of pri-miRNAs by DCL1, the mature duplex remains associated with HYL1 and SE, which later is replaced by HEN1. CARP9 is recruited to this post-processing complex by its binding to HYL1. Nuclear AGO1 is then associated with the complex and HSP90 interacting with CARP9, a process leading to the enhances AGO1 stability and loading of AGO1 with the mature miRNAs. Loaded AGO1 is then exported to the cytosol to silence their target genes.

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