Regulation of pri-miRNA processing by the hnRNP-like protein AtGRP7 in Arabidopsis

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ABSTRACT

The hnRNP-like glycine-rich RNA-binding protein AtGRP7 regulates pre-mRNA splicing in Arabidopsis. Here we used small RNA-seq to show that AtGRP7 also affects the miRNA inventory. AtGRP7 overexpression caused a significant reduction in the level of 30 miRNAs and an increase for 14 miRNAs with a minimum log2 fold change of ±0.5. Overaccumulation of several pri-miRNAs including pri-miR398b, pri-miR398c, pri-miR172b, pri-miR159a and pri-miR390 at the expense of the mature miRNAs suggested that AtGRP7 affects pri-miRNA processing. Indeed, RNA immunoprecipitation revealed that AtGRP7 interacts with these pri-miRNAs in vivo. Mutation of an arginine in the RNA recognition motif abrogated in vivo binding and the effect on miRNA and pri-miRNA levels, indicating that AtGRP7 inhibits processing of these pri-miRNAs by direct binding. In contrast, pri-miRNAs of selected miRNAs that were elevated or not changed in response to high AtGRP7 levels were not bound in vivo. Reduced accumulation of miR390, an initiator of trans-acting small interfering RNA (ta-siRNA) formation, also led to lower TAS3 ta-siRNA levels and increased mRNA expression of the target AUXIN RESPONSE FACTOR4. Furthermore, AtGRP7 affected splicing of pri-miR172b and pri-miR162a. Thus, AtGRP7 is an hnRNP-like protein with a role in processing of pri-miRNAs in addition to its role in pre-mRNA splicing.

INTRODUCTION

Post-transcriptional regulation makes an important contribution to co-ordinating eukaryotic gene expression programs. This control at the RNA level can be executed by proteins or small RNAs. RNA-binding proteins (RBPs) specifically interact with defined sequence motifs in mRNAs to control pre-mRNA splicing, nuclear export or RNA decay (1–3). An abundant class of RBPs comprises the heterogeneous nuclear ribonucleoproteins (hnRNPs), originally identified on the basis of their association with nascent pre-mRNAs (4). More recently, hnRNPs have been shown to mediate many steps in RNA maturation (5,6). Among small regulatory RNAs are the microRNAs (miRNAs) that interact with complementary sites in mRNAs to determine their availability for translation. MiRNAs and their targets are collectively referred to as miRNA modules (7). MiRNAs are 21–24 nt-long single-stranded RNAs generated from precursors that are transcribed by RNA polymerase II (8–10). In Arabidopsis, these pri (primary)-miRNAs are stabilized by DAWDLE (DDL), a forkhead-associated domain protein that likely recruits processing factors (11). The pri-miRNAs contain imperfect double-stranded foldback structures and endonucleolytic cleavage by the RNAseIII family protein DICER-LIKE1 (DCL1) first releases the stem–loops, giving rise to pre (precursor)-miRNAs. These are further processed by DCL1 into duplexes of the miRNA guide strand and the miRNA* passenger strand. The miRNA/miRNA* duplexes have 2-nt overhangs at the 3′ end that are methylated at the 2′-OH groups by HUA ENHANCER 1 (12), and poly(U) tails are added. These features prevent degradation of the miRNAs (12,13).

After export to the cytoplasm the miRNA/miRNA* duplexes dissociate and the miRNA guide strand binds to ARGONAUTE1 (AGO1) or other members of the AGO family. The resulting RNA-induced silencing complexes
guide the AGO proteins to complementary sites in the miRNA targets, leading to their down-regulation by AGO cleavage or inhibition of translation (14,15). Although the miRNA* strand initially was thought to simply decay, accumulating evidence points to a functional role for some miRNA*s (16,17).

The double-stranded RBP HYponastic LEAVes1 (HYL1), the zinc finger protein SERRATE (SE) and the G-patch domain protein TOUGH (TGH) contribute to pri-miRNA processing (18). Together with DCL1, these proteins form the so-called Microprocessor complex that is tethered to the pri-miRNA via interaction with the cap binding proteins CBP20 and CBP80 (19–21). SE and HYL1 promote accurate pri-miRNA processing by DCL1 (22,23), and TGH enhances DCL1 activity without influencing the processing accuracy (18). Thus, several classes of RBPs are involved in miRNA processing. As such a function has not yet been described for a plant hnRN protein, we investigated whether the glycine-rich hnRN-like protein AtGRP7 (Arabidopsis thaliana glycine-rich RBP 7) has an effect on miRNA metabolism. AtGRP7 consists of an RNA recognition motif (RRM) and a namesake glycine stretch. It responds to abiotic stress, is controlled by the circadian clock and participates in pathogen defense (24–29).

AtGRP7 was the first hnRN-like protein in plants that was shown to regulate alternative splicing by direct binding to pre-mRNAs (30).

Here we used small RNA sequencing to show that AtGRP7 affects the miRNA inventory. We identified 44 miRNAs with altered levels in plants ecotopically expressing AtGRP7 (AtGRP7-ox). An increased level of several primary transcripts was found at the expense of the mature miRNAs. Furthermore, AtGRP7 interacted with these pri-miRNAs in vivo and this interaction was lost upon mutation of a single conserved arginine residue in the RRM. Moreover, AtGRP7 affected alternative splicing of pri-miR172b and pri-miR162a. Thus, we identify a novel function for a plant hnRN-like RBP in pri-miRNA processing.

MATERIALS AND METHODS

Transgenic plants

AtGRP7-ox and AtGRP7-R49Q-ox plants express the AtGRP7 wild-type (wt) coding sequence or a mutant version with Arg49 exchanged for Gln under control of the Cauliflower Mosaic Virus (CaMV) 35S promoter in the C24, Col-2 and Landsberg erecta accessions (31–34). AtGRP8-ox plants express the AtGRP8 coding region under control of the CaMV promoter in Col-2 (30).

The line AtGRP7::AtGRP7::GFP expresses an AtGRP7 green fluorescent protein (GFP) fusion under control of 1.4 kb of the AtGRP7 promoter and the AtGRP7 5′UTR, intron and 3′UTR in atgrp7-1, and the line AtGRP7::GFP expresses GFP only under control of the AtGRP7 promoter, 5′UTR and 3′UTR (30,35). To generate an AtGRP7::AtGRP7-R49Q::GFP fusion protein, the 529 bp SacI-XmnI fragment in the AtGRP7::AtGRP7::GFP construct was exchanged with the corresponding fragment carrying the R49Q mutation and the construct was introduced into the atgrp7-1 background. The AtGRP8::AtGRP8::GFP construct includes 1.9 kb of the AtGRP8 promoter and the native 5′UTR, 3′UTR and intron and was introduced into Col-2. The atgrp7-1 mutant lacks detectable AtGRP7 expression due to a T-DNA insertion in the 5′UTR but has elevated levels of the paralog AtGRP8 due to relief from repression by AtGRP7, and the atgrp7-1 81% mutant lacks AtGRP7 and has wt levels of AtGRP8 due to an AtGRP8 RNAi construct (30,36).

Plant growth

Arabidopsis seeds were surface-sterilized and sown on half-strength MS (Murashige–Skoog) (Duchefa) plates (37). Plants were grown in light/dark cycles at 20°C in Percival incubators (CLF laboratories). For RNA analysis, aerial parts of the plants were harvested. At least 10 plants were bulked for each sample in each replicate.

RNA analysis

For RNA analysis, about ten 14-day-old plants per time point were pooled and RNA was isolated using TRIzol® reagent (Invitrogen). For small RNA gel blots, 30 µg of total RNA was fractionated on a 17% polyacrylamide–8 M urea gel in 3-(N-morpholino)propanesulfonic acid buffer, transferred onto GeneScreen membrane and cross-linked with 1-ethyl-3,2-dimethylcarbodiimide and 1-methylimidazole (38). Hybridisation was performed with radiolabeled antisense oligonucleotides in PerfectHybPlus buffer (Sigma) at 38°C. Blots were analysed using a Typhoon 8000 PhosphorImager and ImageQuant software (Amersham Pharmacia Biotech, http://www5.amershambiosciences.com/). Quantification of real-time PCR (RT-PCR) products was done using DNA1000 chips on an Agilent 2100 bioanalyzer. RT-PCR and quantification of miRNAs by stem–loop RT-PCR were done as described (39,40). Primers are listed in Supplementary Table S1.

RNA immunoprecipitation

Plants expressing AtGRP7, AtGRP8 or AtGRP7-R49Q fused to GFP or GFP only were subjected to RNA immunoprecipitation (RIP) as previously described (41). Briefly, plants were subjected to formaldehyde fixation and the extract was subjected to IP with GFP-Trap™ beads (Chromotek) (IP+) and mock IP with RFP-Trap™ beads (Chromotek) (IP−). Coprecipitated RNAs were identified via RT-PCR. In parallel, transcript levels were determined in RNA isolated from the extract before addition of the beads (input).

Small RNA libraries and sequencing

AtGRP7-ox and Col-2 plants were grown for 23 days on half-strength MS medium containing 1% sucrose. Total RNA was extracted from aerial tissue using TRIzol® reagent (Invitrogen). Small RNA libraries were prepared from two biological replicates using the NEBNext® Small RNA Sample Prep Set 1 kit (NEB). Before and after adapter ligation, RNA was size-fractionated on urea-PAGE gels. PCR amplification was done with Phusion® polymerase (Finnzymes) and primers supplied in the NEBNext® Small RNA Sample Prep Kit.
RNA Sample Prep Set 1 kit. The quality of the library was checked on a DNA1000 Bioanalyzer chip and sequencing was performed on the HiSeq2000 Illumina system with a read length of 50 nt.

Bioinformatic evaluation of small RNA-seq data

Reads were mapped onto the Arabidopsis genome (TAIR10) using Bowtie (v0.12.9) without allowing mismatches (42). Mapped reads were counted using the Bedtools (v2.17.0) script coverageBed (43) with the annotation file from miRBase (http://www.mirbase.org/). For mature miRNAs we counted only reads that matched perfectly to the miRBase annotated region without any shift. The differential expression analysis was performed with Bioconductor (2.13) package DESeq (44). MiRNAs were declared as differentially expressed if the adjusted P-value (padj) was below 0.1. To determine miRNAs differentially expressed in the tough mutant (18) the read counts of the dataset (GSE38600) were used.

Processing accuracy of small RNAs

To assess the accuracy of pri-miRNA processing the mapping positions were combined with the annotations for miRNAs from miRBase. The annotated start site at the 5’end and end site at the 3’end were denoted by ASS and AES, and the mapped start and end sites were denoted by MSS and MES, respectively. For each miRNA the distance $D_{\text{start}} = \text{ASS} - \text{MSS}$ and the distance $D_{\text{end}} = \text{AES} - \text{MES}$ was calculated. The histograms of $D_{\text{start}}$ and $D_{\text{end}}$ were calculated separately for Col-2 and AtGRP7-ox.

Immunoblot analysis

Total protein was isolated as described (45). Chemiluminescence detection of the immunoblots was done with Stella (Raytest) and signals were quantified using the AIDA software. The antibodies are listed in Supplementary Table S2.

Confocal microscopy

Transient expression in Nicotiana benthamiana leaves was performed as described (46) and analyzed using a LSM 780 confocal laser scanning microscope (Zeiss). Constructs are described in Supplementary Table S3.

RESULTS

MiR398 abundance is reduced in AtGRP7-ox plants

Genome-wide transcriptome analysis of AtGRP7-ox plants had previously revealed an elevated steady-state abundance of the COPPER/ZINC SUPEROXIDE DISMUTASE 2 (CSD2) transcript encoding a superoxide dismutase that uses copper as a metal cofactor (47). CSD2 is a target of the miR398 family, which comprises identical miRNAs encoded by MIR398b and MIR398c on chromosome V, and a miRNA with a 3’ U instead of G encoded by MIR398a on chromosome II (48,49). To test whether the altered CSD2 transcript level in AtGRP7-ox plants reflected an impact of AtGRP7 on miR398, we monitored miR398 levels using low molecular weight northern blot analysis. MiR398 steady-state abundance was reduced in AtGRP7-ox lines in the Col-2, L er and C24 accessions compared to the corresponding wt plants (Figure 1).

Small RNA profiling reveals a widespread impact of AtGRP7 on the miRNAome

To test whether AtGRP7 affects miRNA levels more globally, small RNA libraries were constructed from AtGRP7-ox and wt plants for high-throughput sequencing. Among 223 miRNAs mapping to the genome with a perfect match, 53 were excluded from further analysis because they were represented by fewer than five reads. The levels of 118 of the remaining 170 miRNAs were not significantly altered. Thus, AtGRP7 overexpression does not have a general effect on miRNA abundance (Supplementary Figure S1A). We tested the validity of our analyses by examining in the same manner a published dataset for the mutant defective in TOUGH (18); our pipeline confirmed a widespread alteration in miRNA abundance in tgh (Supplementary Figure S1B). Of the 52 miRNAs that were significantly changed relative to wt in the AtGRP7-ox plants in two biological replicates, 30 miRNAs showed a log2 fold change $< -0.5$ and the level of 14 miRNAs showed a log2 fold change $> 0.5$ (Supplementary Figure S2, Supplementary Tables S4 and S5). Most of the miRNAs with elevated levels showed only a low read coverage and therefore we mainly focussed on miRNAs with reduced levels upon AtGRP7 overexpression.

Reduced miR398 levels correlate with elevated levels of CSD and CCS in AtGRP7-ox but not AtGRP7-R49Q-ox plants

To validate the RNA-seq data, steady-state abundance of selected miRNAs was monitored by stem–loop RT-PCR. In independent AtGRP7-ox lines grown on half-strength MS medium, miR398 levels reached only 25–50% of the wt level (Figure 2A, Supplementary Figure S3). In contrast, in plants overexpressing AtGRP7 with a single arginine in the RRM mutated (AtGRP7-R49Q-ox) (32) miR398 levels were similar to wt levels. This suggests that AtGRP7 contributes to reduced miR398 steady-state abundance through a mechanism requiring RNA binding. Transcript levels of the miR398 targets CSD1 localized in the cytosol, CSD2 localized in the chloroplast and CCS (COPPER CHAPERONE...
the MIR398b and c promoters to be active (54). When the medium was supplemented with 15 μM Cu2+ that inhibits MIR398b and c promoter activity (52,53), no miR398 was detected and CSD1, CSD2 and CCS accumulated to the same high level in both wt and AtGRP7-ox plants (Supplementary Figure S4). Thus, in the absence of miR398, elevated AtGRP7 levels do not provoke a further increase beyond wt.

**AtGRP7 affects miR390-dependent TAS3 ta-siRNAs**

AtGRP7-ox plants have reduced levels of miR390, the initiator of TAS3 ta-siRNA biogenesis (Figure 3A). This indicates that AtGRP7 also affects miRNAs that are loaded into AGO2 and AGO7 (16,55). We investigated whether the miR390-dependent TAS3 ta-siRNAs are likewise affected. A lower read coverage was observed for all three TAS3 loci in AtGRP7-ox plants (Supplementary Figure S5). Stem–loop RT-PCR confirmed that TAS3′D7(+)/D7′(-) levels were reduced in AtGRP7-ox but not AtGRP7-R49Q-ox plants (Figure 3B, Supplementary Figure S6). Furthermore, expression of the ta-siRNA target ARF4 was elevated in AtGRP7-ox but not AtGRP7-R49Q-ox plants (Figure 3C).

We also confirmed that the levels of both miR172 and miR172b* were significantly reduced in AtGRP7-ox plants (Supplementary Figure S7A). MiR172 targets APETALA-2 type transcription factors mostly at the translational level although slicing also occurs (56,57). Levels of SCHLAFMUETZE (SMZ), a miR172 target regulated by slicing (57), were elevated in AtGRP7-ox plants (Supplementary Figure S7B). Of the miR167 family, miR167c was significantly reduced (Supplementary Table S4) and miR167a and b were weakly reduced in AtGRP7-ox plants. This was confirmed by stem–loop RT-PCR, and the level of its target AUXIN RESPONSE FACTOR8 (ARF8) (58) was increased. For miR824, a recently evolved miRNA encoded by a single locus (59,60) that was expressed at reduced levels in AtGRP7-ox plants, the level of its target AGL16 that has been implicated in stomatal density (61) was slightly elevated in the AtGRP7-ox lines.

For miR159, RNA-seq had not revealed any difference in the miR159a, b and c isoforms in AtGRP7-ox plants whereas stem–loop RT-PCR revealed a reduced level of miR159. The levels of its targets, the MYB33 and MYB65 transcription factors that control many processes including leaf development and flowering time (62) were increased in AtGRP7-ox plants, suggesting that in this case stem–loop RT-PCR may more faithfully reflect the in vivo situation. Such discrepancies in comparing miRNA levels by RNA-seq or RT-PCR based methods have been previously discussed (63).

Among miRNAs with elevated levels in AtGRP7-ox plants compared to wt were members of the miR395 family that is involved in the response to sulfate starvation (Supplementary Figure S7A). The target ATP sulfurylase 3 (APS3), one of the isoenzymes catalyzing the first step of sulfate assimilation, was present at reduced levels in AtGRP7-ox plants (Supplementary Figure S7B). Taken together, altered miRNA levels were confirmed in independent AtGRP7-ox plants and correlated with altered target levels for a suite of the miRNA modules.
Levels of several pri-miRNAs are inversely correlated with mature miRNAs in AtGRP7-ox plants

The altered steady-state level of miRNAs in AtGRP7-ox plants could be a consequence of altered levels of pri-miRNAs or altered fates of mature miRNAs. RT-PCR analysis revealed significantly elevated levels of pri-miR398a, pri-miR398b and pri-miR398c in AtGRP7-ox but not AtGRP7-R49Q-ox plants (Figure 4A and B). Similarly, the primary transcript for miR390b that initiates TAS3 cleavage in ta-siRNA generation was significantly elevated in AtGRP7-ox but not AtGRP7-R49Q-ox plants. Furthermore, we found elevated levels of pri-miR172b and pri-miR159a in independent AtGRP7-ox lines (Figure 4B). As the miR159 precursor is processed by an unusual mechanism with the initial cut near the loop of the stem rather than at its base (64), AtGRP7 appears to affect miRNA precursors processed by the conventional base-to-loop mechanism and precursors processed in loop-to-base direction (Supplementary Tables S4 and S5). The elevated pri-miR399b levels (Figure 4B) correlate with the reduced miR399b level found by RNA-seq.

The miR395 and miR319 families are examples of miRNAs with elevated levels in AtGRP7-ox plants. Pri-miR395e levels were slightly but not significantly reduced and pri-miR319b levels were not altered in AtGRP7-ox plants (Figure 4C), suggesting that AtGRP7 affects different pri-miRNAs by different means. We also monitored the precursors of pri-miR408 and pri-miR171 that were not affected by AtGRP7 and found pri-miR408a and pri-miR171c at similar levels in wt and independent AtGRP7-ox lines.

AtGRP7 does not affect levels of pri-miRNA processing components

The accumulation of pri-miRNAs at the expense of mature miRNAs in AtGRP7-ox plants indicates that AtGRP7 contributes to pri-miRNA processing. To determine whether this is an indirect effect caused by misexpression of general processing factors, we monitored levels of pri-miRNA processing components and additional factors that affect miRNA levels in Arabidopsis. The steady-state abundance of the CBP80, HYLI, DCLI, SE, DDL, HEN1 and TGH transcripts was not consistently altered (Supplementary Figure S8A and B). This was also the case for mRNAs encoding MOS2 (MODIFIER OF SCN1, 2) that facilitates recruitment of pri-miRNAs to the Dicing complex (65), and the proline-rich protein SICKLE (SIC) which is required for accumulation of a subset of miRNAs (66). CBP20 was expressed at a higher level upon AtGRP7 overexpression in C24 but not in Col-2, and the CBP20 protein level was not elevated. The transcript encoding CDC5 (CELL DIVISION CYCLE 5) that functions as an activa-
tor of pri-miRNA transcription and promotes processing of several pri-miRNAs (67) was not significantly affected by AtGRP7. Similarly, the transcripts of the RACK1A (RECEPTOR FOR ACTIVATED C KINASE 1A), RACK1B and RACK1C proteins that interact with SE and act redundantly on miRNA accumulation (40) were not affected. Furthermore, levels of HYL1, CBP80 and SE protein were not altered in AtGRP7-ox plants (Supplementary Figure S8C and D). Thus, AtGRP7 does not have a significant impact on miRNA processing factors.

**AtGRP7 binds pri-miRNAs in vivo**

Because AtGRP7 did not appear to affect miRNA levels indirectly via a global effect on processing factors, we tested whether it directly interacts with pri-miRNAs in vivo. We performed RIP on transgenic plants expressing AtGRP7::GFP under control of its own promoter in the atgrp7−1 background. Pri-miR398b and pri-miR398c were enriched in RNP complexes precipitated with GFP-Trap® beads (IP+) from these AtGRP7::AtGRP7::GFP plants relative to mock precipitates with RFP-Trap® beads (IP−) (Figure 5A). PP2A which served as a negative control was not significantly enriched. Both pri-miR398b and pri-miR398c were not enriched in precipitates from plants expressing GFP alone. Notably, they were also not recovered from plants expressing AtGRP7::AtGRP7-R49Q::GFP (Figure 5B). The pri-miR398a level was too low to allow reliable quantification, in line with its weak expression (68). Furthermore, pri-miR172b and pri-miR159a were enriched in IP+ relative to IP− and not present in precipitates from plants expressing AtGRP7::AtGRP7-R49Q::GFP or GFP only. For pri-miR390b, the expression level was also too low to allow reliable quantification.

If direct binding of AtGRP7 to the pri-miRNA has a functional consequence, the primary transcripts of miRNAs with unaltered levels should not be targets. Indeed, no enrichment of pri-miR408a, pri-miR171b and pri-miR171c in RIP of AtGRP7::AtGRP7::GFP plants was detected (Figure 5D). Similarly, pri-miR319b was not enriched. pri-miR319 is an example of a miRNA with an elevated level in AtGRP7-ox plants but pri-miRNA level is not altered. This suggests that the effect of AtGRP7 on pri-miR319 is indirect.

**AtGRP7 colocalizes with processing components**

Binding of AtGRP7 to pri-miRNAs in vivo implies a direct role in processing. Therefore, we compared the subcellular localization of AtGRP7 to that of other processing components by transient coexpression of fluorescent protein-tagged fusion proteins in N. benthamiana leaves (Supplementary Figure S9). AtGRP7 colocalizes with DCL1, SE, DDL and CBP80 in the nucleus. In the case of DCL1 and SE the colocalization is confined to the nucleoplasm whereas DDL and CBP80 also localize to the nucleolus, as does AtGRP7. For DCL1, we detected colocalization in punctate structures in the nucleus, likely representing dicing bodies that are known sites of miRNA processing (69–71).

The paralogous AtGRP8 can compensate for loss of AtGRP7

As elevated levels of AtGRP7 impair processing of a suite of pri-miRNAs, we asked whether the absence of AtGRP7 would likewise have an effect. Levels of mature miR398 were not significantly changed in the atgrp7−1 line that lacks AtGRP7 but has higher levels of its paralog AtGRP8 than wt due to relief from repression by AtGRP7 (Supplementary Figure S10A and B). It was also not changed in the line atgrp7−1 i8 that lacks AtGRP7 and has wt levels of AtGRP8 due to an RNAi construct. Pri-miR398a and pri-miR398b levels were slightly but significantly decreased in atgrp7−1 i8 and thus changed in the opposite direction in response to elevated and reduced AtGRP7, respectively, as expected for a direct target (Supplementary Figure S10C). The level of pri-miR398c was not consistently altered either in atgrp7−1 or atgrp7−1 i8. Similarly, levels of miR172b*, miR395 and miR159 were not altered. Neither were the levels of miR390, the TAS3 5′D7(+)-ta-siRNA and ARF4 altered (Supplementary Figure S10D–F). Thus, while AtGRP7 clearly contributes to processing of a subset of pri-miRNAs, a reduced level appears to be compensated by other factors.

To test whether the residual AtGRP8 in the atgrp7−1 i8 line may mask the loss of AtGRP7, we tested the effect of AtGRP8 on miRNA levels. In the absence of a true atgrp8 loss-of-function mutant, we investigated AtGRP8-ox plants and found the levels of miR398 to be reduced to ∼35% of the wt level and miR172 and miR159 to be reduced to ∼50% (Supplementary Figure S11A). In turn, the levels of pri-miR398b, pri-miR172b and pri-miR159a were elevated (Supplementary Figure S11B). This suggests that AtGRP8 in addition to AtGRP7 promotes the accumulation of these pri-miRNAs at the expense of the mature miRNA and thus the presence of AtGRP8 in atgrp7−1 or atgrp7−1 i8 masks the mutant phenotype.

To test whether AtGRP8 is also able to interact in vivo with the pri-miRNAs it regulates, we generated transgenic plants expressing AtGRP8::AtGRP8::GFP. Pri-miR398b, pri-miR398c, pri-miR172b and pri-miR159a indeed also coprecipitated with AtGRP8 (Supplementary Figure S11C). The weaker enrichment compared to the AtGRP7 RIP likely is due to competition by the endogenous AtGRP8 and AtGRP7 proteins, as in the absence of a true atgrp8 loss-of-function mutant, the AtGRP8::AtGRP8::GFP line had to be generated in the wt background. As a control, we monitored the interaction of AtGRP8::GFP with AtGRP7 and AtGRP8 itself that had been demonstrated in vitro (28). We confirmed that AtGRP8 interacts with these transcripts also in vivo.

**Processing accuracy in AtGRP7-ox and wt plants**

Altered levels of miRNAs in AtGRP7-ox plants may be due to imprecise processing of pri-miRNAs such that the excised miRNAs would not fall exactly within the annotated mature miRNA positions. Therefore, the start positions at the 5′end and the end positions at the 3′end of each excised miRNA in wt and AtGRP7-ox plants were compared to the start and end positions annotated in miBase. The resulting differences of the mapped and annotated start sites or end sites are presented in Supplementary Figure S12. The distribution showed a strong peak at the annotated 5′ and
Figure 5. At GRP7 binds to pri-miRNAs in vivo. Plants expressing AtGRP7::AtGRP7::GFP (A, D) or AtGRP7::AtGRP7::R49Q::GFP (B) in the atgrp7−1 background and AtGRP7::GFP in Col-2 (C) were subjected to RIP. The levels of the pri-miRNAs and PP2A were determined in the GFP-Trap beads precipitate (IP+), the RFP-Trap beads precipitate (IP−) and the input fraction (IN), respectively. Pri-miR398b and c, pri-miR172b and pri-miR159a correspond to miRNAs with reduced level in AtGRP7-ox plants, pri-miR319b corresponds to a miRNA with elevated levels in AtGRP7-ox plants, and pri-miR408a and pri-miR171b and c correspond to miRNAs that are not affected. Data are based on three biological replicates. Asterisks denote a significant difference according to Student’s t-test (P < 0.05). n.s., not significant; n.d., not detectable.

3′ ends both in wt and AtGRP7-ox, suggesting that AtGRP7 does not generally affect processing accuracy.

Furthermore, we mapped all reads to the annotated pri-miRNAs and found no major products from non-standard processing outside the annotated miRNA and miRNA* positions (Supplementary Figure S13). Finally, to test whether miRNAs of unusual size accumulate in AtGRP7-ox plants, the read length distributions of the excised miRNAs were compared between wt and AtGRP7-ox plants. Most pri-miRNAs in AtGRP7-ox plants released sRNAs with a similar size distribution to that in wt (Supplementary Figure S14). Taken together, this suggests that AtGRP7 does not generally affect processing accuracy.

Whereas in animals miRNA precursors have a fold-back structure of similar size, plant miRNA precursors can adopt a wide range of structures and the fold-back structures can be up to 900 nt in length (72). To investigate whether the action of AtGRP7 on a miRNA may correlate with the length of its fold-back structure, we retrieved the sizes of 329 miRNA stem–loops from miRBase. The size distribution of precursors of the miRNAs with significantly reduced or elevated levels in AtGRP7-ox plants did not deviate from the overall size distribution (Supplementary Figure S15). Thus, the length of the fold-back structure does not appear to determine the preference of AtGRP7 for pri-miRNAs.

At GRP7 affects alternative splicing of pri-miR172b

At GRP7 overexpression leads to reduced levels of miR172b and miR172b* with a concomitant increase in pri-miR172b which contains introns both 5′ and 3′ of the stem–loop structure (Figure 6A). Because splicing of introns located 3′ of the stem–loop can impact accumulation of mature miRNAs (73,74), we asked whether AtGRP7 affects splic-
Figure 6. *AtGRP7* affects alternative splicing of pri-miRNAs. (A) Scheme of *MIR172b*. Black boxes = exons, grey box = position of the pri-miRNA, thin line = introns. The arrows denote the position of the primers used in (B). (B) RNA from the *AtGRP7*-ox lines D and G and Col-2 wt was analysed by RT-PCR. The amplification products corresponding to the intron-retained form and the fully spliced forms are indicated by arrowheads and asterisks, respectively. *PP2A* served as a control. DNA = genomic DNA. (C) The ratio of intron-retained versus spliced *pri-MIR172b* was quantified using Bioanalyzer DNA1000 chips. Shown is the mean of two reps. (D) Scheme of the non-protein-coding RNA harbouring *MIR162a*. Black boxes = exons, grey box = position of the pri-miRNA, open boxes = annotated 5’ and 3’ UTRs, thin line = introns. The arrows denote the position of the primers used in (E). (E) RNA from the *AtGRP7*-ox lines D and G and Col-2 wt was analysed by RT-PCR. The transcript forms corresponding to the amplification products are indicated. The rhombus denotes an alternative version of the 168 nt band generated by an alternative 3’ splice site 3 nt downstream of the authentic 3’ splice site. *PP2A* served as a control. DNA = genomic DNA. (F) The ratio of the alternative splice forms versus the spliced form was quantified using Bioanalyzer DNA1000 chips. Shown is the mean of two reps.

In *AtGRP7*-ox plants, the spliced *pri-MIR172b* accumulated at the expense of the intron retained form, indicating that *AtGRP7* indeed affects splicing of pri-miRNAs (Figure 6B and C). In contrast, splicing of the introns located upstream of the hairpin was not altered (not shown). Thus, an enhanced assembly of the spliceosome at the 5’ end of the downstream intron in *AtGRP7*-ox plants may interfere with pri-miRNA processing. Alternatively, a slower release of miR172b and miR172b* caused by *AtGRP7* binding may allow more efficient assembly of the spliceosome.

While in animals around 80% of all miRNAs are encoded in introns, in Arabidopsis fewer miRNA genes are intronic (75). Among those is *pri-MIR162a* located within an alternative intron of non-protein-coding RNA 78 (Figure 6D) (76). In *AtGRP7*-ox plants, the unspliced precursor that retains the *MIR162a* gene is present at higher levels than in wt plants, whereas the isoform with exon 2 skipped that lacks the entire *MIR162a* gene is present at somewhat reduced levels (Figure 6E and F). An alternative isoform that retains exon 2 but lacks part of the stem–loop is present at higher levels in *AtGRP7*-ox plants. Thus, *AtGRP7* can also impact alternative splicing of an intronic pri-miRNA, but the level of mature miR162a was not significantly altered in *AtGRP7*-ox plants. MiR162a targets *DCL1*, pointing to a complex feedback regulation between the miRNA and its target.

**DISCUSSION**

Here we show that the hnRNP-like protein *AtGRP7* affects steady-state abundance of a suite of miRNAs in Arabidopsis. Small RNA profiling of plants ectopically expressing *AtGRP7* identified subsets of miRNAs that accumulated either to lower or higher levels than in wt. Because most miRNAs with elevated levels showed only a low read coverage, we focussed on miRNAs with reduced levels.

One affected miRNA was miR398, and reduced miR398 levels correlated with increased levels of the miR398 targets CSD1, CSD2 and the copper chaperone CCS in *AtGRP7*-ox plants. Importantly, the effect of *AtGRP7* on these miR398 targets was dependent on miR398, as upon deple-
ation of miR398 by high Cu\textsuperscript{2+} concentrations CSD and CCS levels in \textit{AtGRP7-ox} plants were indistinguishable from wt.

For a suite of miRNAs with reduced levels in \textit{AtGRP7-ox} plants, a reciprocal increase in pri-miRNA levels was observed in independent \textit{AtGRP7-ox} lines, implicating \textit{AtGRP7} in processing of a number of pri-miRNAs into mature miRNAs. To test whether \textit{AtGRP7} was required for pri-miRNA processing, we monitored miRNA and pri-miRNA levels in lines lacking \textit{AtGRP7}. In the \textit{atgrp7–1} mutant, the levels of the tested miRNAs and pri-miRNAs were virtually unaltered relative to wt plants. This line lacks \textit{AtGRP7} but has elevated levels of the paralogous protein \textit{AtGRP8} due to relief from repression by \textit{AtGRP7}. Also in the \textit{atgrp7–1} \textit{8f} line that lacks \textit{AtGRP7} and expresses \textit{AtGRP8} at wt levels due to an RNAi construct most tested miRNAs and pri-miRNAs remained at wt levels. Only for pri-miR398a and b were levels significantly reduced relative to wt and thus changed in opposite directions upon loss-of-function and \textit{AtGRP7} overexpression, respectively, as predicted for a direct target.

This limited phenotype in the loss-of-function lines indicates that \textit{AtGRP7} is not essential for the regulation of pri-miRNA processing. Thus, the impact of \textit{AtGRP7} on miRNA metabolism observed in the \textit{AtGRP7-ox} plants may be unrelated to its function in wt plants and rather be attributed to non-physiological \textit{AtGRP7} levels. However, we consider this unlikely for three reasons. Firstly, the accumulation of pri-miRNAs at the expense of mature miRNAs and the reciprocal increase in the miRNA targets were observed upon overexpression of \textit{AtGRP7} but not upon expression to the same high level of a mutant variant of \textit{AtGRP7} with a single amino acid exchange (\textit{AtGRP7-R\textsuperscript{49}Q}) (Figure 2D). This is a strong hint that the impact of \textit{AtGRP7} on miRNA metabolism is not an artefact of high protein levels but rather a specific function of the protein requiring its RNA-binding activity. The importance of R\textsuperscript{49} for \textit{AtGRP7} function has been demonstrated before: during infection of Arabidopsis plants with \textit{Pseudomonas syringae}, R\textsuperscript{49} is ADP-ribosylated by the bacterial effector HopU1 (36,77), interfering with \textit{AtGRP7} binding to defense-related transcripts as part of the bacterial virulence strategy (29,39). Furthermore, the increased sensitivity of \textit{atgrp7–1} to virulent \textit{P. syringae} is complemented by a 3 kb genomic \textit{AtGRP7} fragment but not the same fragment with R\textsuperscript{49} mutated (29). Secondly, in \textit{in vivo} binding of \textit{AtGRP7} to miRNA precursors supports our hypothesis that increased accumulation of several pri-miRNAs in \textit{AtGRP7-ox} plants is due to impaired processing. Importantly, the \textit{AtGRP7::AtGRP7-GFP atgrp7–1} line used for RIP expresses \textit{AtGRP7-GFP} under control of the native promoter and noncoding regions in the absence of endogenous \textit{AtGRP7} and thus has wt levels of \textit{AtGRP7}, ruling out any effect due to overexpression compared to wt levels. \textit{In vivo} binding was observed for precursors of miRNAs with reduced levels in \textit{AtGRP7-ox} plants but not for those of miRNAs expressed at wt levels. Moreover, the pri-miRNAs were not enriched in immunoprecipitates from \textit{AtGRP7::AtGRP7-R\textsuperscript{49}Q-GFP} plants. This indicates that direct binding of the pri-miRNAs has functional significance for the impact of \textit{AtGRP7} on pri-miRNA processing. Thirdly, in a previous microarray analysis only 0.4% of the transcripts present on the ATH1 microarray changed >1.5-fold in \textit{AtGRP7-ox} lines relative to wt, arguing against a nonspecific misregulation of a large body of genes in \textit{AtGRP7-ox} plants (47).

Based on these observations, we rationalized that the limited miRNA phenotype of the \textit{atgrp7} mutant lines may be due to redundancy with other regulatory proteins, an obvious candidate being the paralogous \textit{AtGRP8} protein. In the absence of a true \textit{atgrp7} loss-of-function line, we monitored the levels of several miRNA and pri-miRNAs that are regulated by \textit{AtGRP7} in \textit{AtGRP8-ox} plants. Indeed, several pri-miRNAs accumulated at the expense of the mature miRNAs in the \textit{AtGRP8-ox} plants. Moreover, an \textit{AtGRP8-GFP} fusion protein expressed under control of the native promoter and the noncoding regions showed \textit{in vivo} binding to the precursors of these miRNAs. Thus, we conclude that \textit{AtGRP7} and \textit{AtGRP8} inhibit processing of a suite of pri-miRNAs by direct binding, and that a lack of \textit{AtGRP7} does not promote processing of these pri-miRNAs due to redundancy with \textit{AtGRP8} and, perhaps, other regulatory proteins that remain to be identified.

In line with a direct role in pri-miRNA processing, \textit{AtGRP7} colocalizes with DCL1, HYL1, CBP80 and SE in the nucleus. Furthermore, steady-state levels of components known to be involved in miRNA biogenesis were not altered in \textit{AtGRP7-ox} plants, consistent with a previous microarray analysis (47) and arguing against an indirect effect. Several RBPs with different types of RNA-binding domains have been shown to co-operate with DCL1, including HYL1 with a double-stranded RNA-binding domain (22,78,79), SE with a zinc finger (19,80), TOUGH with a G patch domain (18) and the CAP binding complex (CBC) subunits CBP20 and CBP80 (19,20). These proteins have a more global impact on pri-miRNA processing whereas \textit{AtGRP7} affects processing of a limited set of pri-miRNAs. Furthermore, these DCL1-interacting proteins promote processing whereas \textit{AtGRP7} appears to mostly act as an inhibitor.

\textit{AtGRP7} is the first plant hnRNP-like protein shown to regulate miRNA biogenesis. A role for an hnRNP in the processing of a miRNA has first been described for human hnRNP A1 (81). HnRNP A1 binds specifically to pri-miRNAs, plant miRNA precursors display stem–loops of highly variable size that may pose additional challenges to the processing machinery (72). Comparison of the length of the stem–loop of pri-miRNAs annotated in
miRBase with the size distribution of precursors with significantly altered levels in AtGRP7-ox plants revealed no difference. Processing of the pri-miRNAs mostly involves an initial cleavage at the base of the stem and proceeds to the loop. Recently, several pri-miRNAs have been shown to be processed with the first cut at the loop, proceeding to the base (64). Moreover, depending on the number of cuts by DCL1 to release the mature miRNA, sequential base-to-loop and sequential loop-to-base mechanisms have been defined if more than one cut by DCL1 is required to release the mature miRNA (72). Among the pri-miRNAs binding to AtGRP7 \textit{in vivo} was pri-miR159a which is processed via an initial cut at the loop (64). Thus, AtGRP7 appears to affect miRNA precursors processed by the conventional base-to-loop mechanism or in loop-to-base direction. Moreover, among the differentially expressed miRNAs are several that are generated by sequential cuts (indicated in Supplementary Tables S4 and S5).

Altered levels of miRNAs in AtGRP7-ox plants may also be due to imprecise processing of pri-miRNAs such that the excised miRNAs would not fall exactly within the annotated mature miRNA. However, the processing accuracy was not altered upon AtGRP7 overexpression, different from what has been observed in hyll and se mutants (85). In the future, the establishment of crosslinking and immunoprecipitation techniques in plants to determine \textit{in vivo} binding sites will help to define pri-miRNA features recognized by AtGRP7.

Recently, a reciprocal interaction between splicing of introns in pri-miRNAs that are located downstream of the stem–loop and accumulation of the corresponding mature miRNA has been described (73,74). We find that splicing of the pri-miR172b intron located downstream of the stem–loop is favored in AtGRP7-ox plants compared to wt with concomitantly reduced levels of miR172b and miR172b*. Because AtGRP7 affects alternative splicing of pre-miRNAs, it may promote removal of the intron and lead to reduced miR172b levels. On the other hand, impaired processing of the stem–loop may allow more efficient splicing of the intron, as observed for dcl1 mutants (73). However, the role of AtGRP7 in inhibiting processing is not contingent on removal of introns, as it also affects steady-state levels of miRNAs derived from intron-less pri-miRNAs, e.g. pri-miR167a and pri-miR172e. In turn, AtGRP7 affects alternative splicing of pri-miR162a (30) but the level of mature miR162a is not significantly altered in AtGRP7-ox plants. Thus, the common features of pri-miRNAs regulated by AtGRP7 remain to be determined.

Taken together, AtGRP7 is the first hnRNP-like protein in plants with a dual role in alternative splicing of pre-miRNAs and maturation of pri-miRNAs. This resembles the function of SE and the CBC (19,86). For AtGRP7, regulation of some of its splicing substrates is due to direct binding \textit{in vivo}, not yet shown for SE or the CBC (30). AtGRP7 not only directly impacts miRNAs through alternative splicing but also indirectly via pri-miRNA processing and consequently miRNA levels, supporting the view of extensive cross-talk between regulation by RBPs and miRNAs.

**SUPPLEMENTARY DATA**

**Supplementary Data** are available at NAR Online.

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