MAC3A and MAC3B, Two Core Subunits of the MOS4-Associated Complex, Positively Influence miRNA biogenesis

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Li, Shengjun; Liu, Kan; Zhou, Bangjun; Li, Mu; Zhang, Shuxin; Zeng, Lirong; Zhang, Chi; and Yu, Bin, "MAC3A and MAC3B, Two Core Subunits of the MOS4-Associated Complex, Positively Influence miRNA biogenesis" (2018). *Faculty Publications from the Center for Plant Science Innovation*. 182.  
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MAC3A and MAC3B, Two Core Subunits of the MOS4-Associated Complex, Positively Influence miRNA biogenesis

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Short title: MAC3A and MAC3B in miRNA biogenesis

One-sentence summary: The MOS4-associated complex promotes miRNA accumulation by positively modulating pri-miRNA transcription, stability and processing.

The author responsible for distribution of materials integral to the findings presented in this manuscript in accordance with the policy described in the Instructions for Authors (www.plantcell.org) is: Bin Yu (byu3@unl.edu)

ABSTRACT

MAC3A and MAC3B are conserved U-box containing proteins in eukaryotes. They are subunits of the MOS4-associated complex (MAC) that plays essential roles in plant immunity and development in Arabidopsis. However, their functional mechanisms remain elusive. Here we show that Arabidopsis thaliana MAC3A and MAC3B act redundantly in microRNA (miRNA) biogenesis. Lack of both MAC3A and MAC3B in the mac3b mac3b double mutant reduces the accumulation of miRNAs, causing elevated transcript levels of miRNA targets. mac3a mac3b also decreases the levels of primary miRNA transcripts (pri-miRNAs). However, MAC3A and MAC3B do not affect the promoter activity of genes encoding miRNAs (MIR genes), suggesting that they may not affect MIR transcription. This result together with the fact that MAC3A associates with pri-miRNAs in vivo indicates that MAC3A and MAC3B may stabilize pri-miRNAs. Furthermore, we find that MAC3A and MAC3B interact with the DCL1 complex that catalyzes miRNA maturation, promote DCL1 activity and are required for the localization of HYL1, a component of the DCL1 complex. Besides MAC3A and MAC3B, two other MAC subunits, CDC5 and PRL1, also function in miRNA biogenesis. Based on these results, we propose that MAC functions as a complex to control miRNA levels through modulating pri-miRNA transcription, processing and stability.

INTRODUCTION

MicroRNAs (miRNAs), ~21-nucleotide in size, are endogenous non-coding RNAs that mainly repress gene expression at post-transcriptional levels (Baulcombe, 2004; Axtell, 2013). They are
generated from the imperfect stem-loop residing in the primary miRNA transcripts (pri-miRNAs) (Voinnet, 2009), most of which are produced by DNA-dependent RNA polymerase II (Xie et al., 2005). In plants, the RNase III enzyme DICER-LIKE 1 (DCL1) slices pri-miRNAs at least two times in the nucleus to release a miRNA-containing duplex (Baulcombe, 2004; Axtell, 2013; Zhang et al., 2015). Then, the small RNA methyltransferase HUA ENHANCER1 (HEN1) methylates the miRNA duplexes to protect them from degradation and untemplated uridine addition (Zhai et al., 2013; Ren et al., 2014). Following methylation, the miRNA strand is incorporated into the effector called ARGONAUTE 1 (AGO1) with the assistance from HEAT SHOCK PROTEIN 90 and CYCLOPHILIN 40 and recognizes target transcripts through sequence complementarity (Baumberger and Baulcombe, 2005; Vaucheret, 2008; Smith et al., 2009; Earley and Poethig, 2011). AGO1 cleaves target mRNAs or inhibits their translation, and therefore, represses gene expression.

Pri-miRNAs may be co-transcriptionally processed since DCL1 associates with MIR loci (Fang et al., 2015a). In the past decades, protein factors that regulate miRNA biogenesis through influencing pri-miRNA transcription, processing and stability have been identified in plants. The transcriptional co-activator MEDIATOR (Kim et al., 2011), the CYCLIN-DEPENDENT KINASES (CDKs) (Hajheidari et al., 2012), the transcription factor NEGATIVE ON TATA LESS 2 (NOT2) (Wang et al., 2013), the DNA binding protein CELL DIVISION CYCLE 5 (CDC5) (Zhang et al., 2013) and ELONGATOR (Fang et al., 2015a) are required for optimized Pol II activity at the MIR promoters. Following transcription, the forkhead domain-containing protein DAWDLE (DDL) (Yu et al., 2008) and the WD-40 protein PLEIOTROPIC REGULATORY LOCUS 1 (PRL1) (Zhang et al., 2014) bind pri-miRNAs to prevent their degradation.

To efficiently and accurately process pri-miRNAs, DCL1 forms a complex with the double stranded RNA (dsRNA)-binding protein HYPERSONASTIC LEAVES1 (HYL1), the Zinc-finger protein SERRATE (SE) and the RNA-binding protein TOUGH (TGH) (Fang and Spector, 2007; Fujioka et al., 2007; Song et al., 2007; Dong et al., 2008; Ren et al., 2012). The formation of the DCL1 complex requires NOT2 (Wang et al., 2013), ELONGATOR (Fang et al., 2015a), MODIFIER OF SNC1, 2 (MOS2, an RNA-binding protein) (Wu et al., 2013) and the DEAH-box
helicase PINP1 (Qiao et al., 2015). How MOS2 and PINP1 participate in the assembly of the DCL1 complex remains unclear, since they do not interact with the DCL1 complex (Wu et al., 2013; Qiao et al., 2015). Efficient loading of pri-miRNAs to the DCL1 complex requires TGH (Ren et al., 2012), the THO/TREX complex that is involved in nuclear RNA transport (Francisco-Mangilet et al., 2015), and the ribosome protein STV1 (Li et al., 2017). Notably, several additional proteins including the CAP-BINDING PROTEINs (CBPs) (Gregory et al., 2008; Laubinger et al., 2008), NOT2, ELONGATOR, DDL, CDC5 and PRL1 also associate with the DCL1 complex to enhance pri-miRNA processing. In addition, SICKLE (SIC, a proline-rich protein) (Zhan et al., 2012), RECEPTOR FOR ACTIVATED C KINASE 1 (RACK1) (Speth et al., 2013), STABILIZED1 (STA1, a pre-mRNA processing factor 6 homolog) (Ben Chaabane et al., 2013), REGULATOR OF CBF GENE EXPRESSION 3 (RCF3, also known as HOS5 and SHI1) (Chen et al., 2015; Karlsson et al., 2015) and GRP7 (a glycine-rich RNA-binding protein) (Koster et al., 2014) also regulate miRNA biogenesis. However, they do not associate with DCL1. Moreover, phosphorylation and dephosphorylation of HYL1 are crucial for pri-miRNA processing (Manavella et al., 2012). In addition, protein factors that act in miRNA biogenesis are also transcriptionally and post-transcriptionally regulated. For instance, DCL1 transcription is modulated by the histone acetyltransferase GCN5 (Kim et al., 2009), STA1 (Ben Chaabane et al., 2013) and the transcription factor XAP5 CIRCADIAN TIMEKEEPER (XCT) (Fang et al., 2015b). Notably, HYL1 protein levels are maintained by the SNF1-RELATED PROTEIN KINASE 2 (Yan et al., 2017) and the E3 ubiquitin ligase CONSTITUTIVE PHOTOMORPHOGENIC 1 (COP1) (Cho et al., 2014) through unknown mechanisms. Recently, KETCH1 (KARYOPHERIN ENABLING THE TRANSPORT OF THE CYTOPLASMIC HYL1)-mediated transportation of HYL1 from the cytoplasm to the nucleus was shown to be crucial for miRNA biogenesis (Zhang et al., 2017). Interestingly, pri-miRNA structures also influence the DCL1 activity (Mateos et al., 2010; Song et al., 2010; Werner et al., 2010; Bologna et al., 2013; Zhu et al., 2013). For instance, the internal loop below the miRNA/miRNA* within the stem-loop is important for the processing of some pri-miRNAs.

Among proteins associated with the DCL1 complex, CDC5 and PRL1 are two core subunits of the MOS4-associated complex (MAC) (Monaghan et al., 2009). MAC is a conserved complex that associates with the spliceosome (Deng et al., 2016). Its homolog complexes in human and
yeast are known as the CDC5-SNEV^{Prp19-Pso4} (PRP19) complex and the Nineteen complex (NTC), respectively (Palma et al., 2007). Both PRP19 and NTC function in splicing, DNA repair, cell cycle and genome stability (Chanarat and Strasser, 2013). MAC contains three additional core subunits, MAC3A, MAC3B and MOS4, and at least 13 accessory proteins with diversified functions (Monaghan et al., 2009). Deficiency in MAC impairs plant immunity and development (Monaghan et al., 2009). However, related mechanisms still need investigation. We have previously shown that CDC5 and PRL1 have overlapping roles in regulating DCL1 activity, but distinct functions in pri-miRNA transcription and stability (Zhang et al., 2013; Zhang et al., 2014). These results raise the possibility that other MAC components may also have diversified effects on miRNA biogenesis. Among core MAC components, MAC3A and MAC3B are two homologous U-box type E3 ubiquitin ligases (~82% identity and 90% similarity) (Monaghan et al., 2009). E3 ligase activity of MAC3B has been demonstrated \textit{in vitro} (Wiborg et al., 2008).

We previously showed that a loss-of-function mutation in MAC3A does not affect miRNA accumulation (Zhang et al., 2014). However, this result may reflect the redundant function of MAC3B with MAC3A.

In this study, we found that lack of both MAC3A and MAC3B reduces the accumulation of miRNAs and impairs the localization of HYL1 in the D-body. MAC3A associates with the DCL1 complex and pri-miRNAs and promotes pri-miRNA processing. MAC3A and MAC3B are also required for accumulation of pri-miRNAs. However, unlike CDC5, MAC3A neither interacts with Pol II nor affects \textit{MIR} transcription. These results suggest that MAC3A/3B may stabilize pri-miRNAs and act as a co-factor to promote D-body formation and pri-miRNA processing. In addition, we show that MAC3A is a phosphorylation-dependent E3 ligase and its E3 ligase activity is required for miRNA biogenesis. We propose that MAC may act as a complex to promote miRNA biogenesis and different MAC components may have distinct and cooperative effects on pri-miRNA transcription, stability and processing.
RESULTS

MAC3A and MAC3B are required for miRNA biogenesis

The fact that CDC5 and PRL1, two core components of MAC are required for miRNA biogenesis suggests that other MAC components may also function in miRNA biogenesis. However, we previously showed that a single mac3a mutation does not affect miRNA accumulation in *Arabidopsis thaliana* (Zhang et al., 2014). To evaluate if this result might reflect redundancy between MAC3A and MAC3B in miRNA biogenesis, we generated a mac3a mac3b double mutant through crossing mac3a (Salk_089300) to mac3b (Salk_050811) (Monaghan et al., 2009). Compared with Col (wild-type plant, WT), mac3a mac3b displayed pleiotropic development defects (Figure 1). For instance, the root length of mac3a mac3b is much shorter (Figure 1A and I). Moreover, the size of the mac3a mac3b was smaller (Figure 1B). Reduced cell number was likely responsible for the smaller size of mac3a mac3b, since the size of palisade cells from mac3a mac3b was comparable to that from Col (Figure 1C, 1D and 1J). In addition, mac3a mac3b leaves had three to four branch points (4–5 branches) on average, while most trichomes of Col had two branch points (three branches)(Figure 1E, 1F and 1K). Furthermore, the silique length of mac3b mac3b was shorter than that of Col (Figure 1G and 1L). Moreover, the amounts of aborted seeds were higher in the siliques of mac3a mac3b than those of WT (Figure 1H and 1M), suggesting that MAC3A and MAC3B also affect fertility.

The pleiotropic growth defects of mac3a mac3b are consistent with the effect of miRNAs on plant development; we therefore examined the accumulation of miRNAs in mac3a mac3b and Col through RNA gel blot. The abundance of all nine examined miRNAs was reduced in mac3a mac3b relative to Col (Figure 2A). RT-quantitative PCR (RT-qPCR) analyses further confirmed that miRNA levels were decreased in mac3a mac3b (Figure 2B). We also examined the effect of MAC3A and MAC3B on trans-acting siRNAs (ta-siRNAs), which is another class of sRNAs that represses gene expression at post-transcriptional levels (Peragine et al., 2004; Allen et al., 2005; Yoshikawa et al., 2005; Axtell et al., 2006). Similar to miRNAs, ta-siR255 was reduced in abundance in mac3a mac3b (Figure 2A). However, the effect MAC3A and 3B on ta-siR255 might be indirect, since the production of ta-siRNAs depending on miRNAs, whose abundance was reduced in mac3a mac3b.
We further compared miRNA profile from inflorescences of mac3a mac3b with that of WT through deep sequencing. The abundance of many miRNAs was reduced in mac3a mac3b relative to WT (Supplemental Figure 1A and Supplemental Data Set 1), suggesting that MAC3A MAC3B may have a global effect on miRNA accumulation. We also compared the effect of mac3a mac3b on miRNA accumulation with that of dcl1-9 (a weak allele of dcl mutants) and cdc5. As expected, cdc5 and dcl1-9 reduced the abundance of most miRNAs (Supplemental Figure 1B, 1C and Supplemental Data Set 1). Among significantly down-regulated miRNAs (P<0.1), DCL1, CDC5 and MAC3A/MAC3B showed overlapping effects on many of them (Supplemental Figure 1D). However, some miRNAs were differentially affected by DCL1, CDC5 and MAC3A/MAC3B (Supplemental Figure 1D). These results suggest that these proteins may have overlapping and distinct roles in miRNA biogenesis.
Next, we evaluated the influence of mac3a mac3b on the transcript levels of ARF4, ARF8, CKB3, CUC1, MYB33, PHO2, PHV, PPR, and SPL9/10/13, which are targets of tasiR-ARF, miR167, miR397, miR159, miR399, miR166, miR400, miR156, respectively. The levels of these target transcripts were increased in mac3a mac3b compared with Col (Figure 2C), suggesting that MAC3A and 3B are required for optimal activity of miRNAs and ta-siRNAs.

To determine if the lack of MAC3A and MAC3B was responsible for the observed phenotypes, we expressed a genomic copy of MAC3A fused with a GUS gene at its 3’ end under the control of its native promoter (proMAC3A:MAC3A-GUS) in mac3a mac3b. The expression of this transgene rescued the developmental defects of mac3a mac3b (Supplemental Figure 2A). In
addition, fusion constructs MAC3B-GFP (pro35S:MAC3B-GFP) or MYC-MAC3A (pro35S:MYC-MAC3A) under the control of the 35S promoter also complemented the developmental defects of mac3a mac3b (Supplemental Figures 2B and 2J). Consistent with this observation, miRNA and target transcript levels in the complementation lines were comparable to those in Col (Supplemental Figure 2K and 2L). We also examined the expression pattern of MAC3A in mac3a mac3b harboring proMAC3A:MAC3A-GUS through GUS histochemical staining. MAC3A was universally expressed and displayed high expression levels in primary root tip, lateral root, and young leaves (Supplemental Figure 2C-2I). These results demonstrate that MAC3A and MAC3B act redundantly to control development and miRNA accumulation of Arabidopsis.

MAC3A and MAC3B do not affect MIR transcription

We have previously shown that CDC5 and PRL1 regulate pri-miRNA levels through modulating pri-miRNA transcription and stability, respectively (Zhang et al., 2013; Zhang et al., 2014). This led us to test if pri-miRNA levels were also altered in mac3a mac3b. As expected, all examined pri-miRNAs were reduced in abundance in mac3a mac3b compared with Col (Figure 3A). We suspected that as in cdc5, this reduction could be caused by alteration in transcription. Thus, we evaluated the effect of mac3a mac3b on MIR promoter activity. The MIR promoter reporter construct, pMIR167a:GUS (Zhang et al., 2014), was crossed into mac3a mac3b. Histochemical staining and RT-qPCR analyses revealed that the expression levels of GUS in mac3a mac3b were similar to those in WT (Figure 3B and 3C), indicating that MAC3A and MAC3B may have no effect on MIR promoter activity. Furthermore, we tested the interaction between MAC3A and the second largest subunit of Pol II (RPB2) through co-immunoprecipitation assay (Co-IP) in the mac3a mac3b expression pro35S:MAC3A-GFP. In MAC3A-GFP precipitates, we did not detect the presence of RPB2 (Figure 3D), suggesting that unlike CDC5 and PRL1, MAC3A does not associate with RPB2. We also examined the occupancy of Pol II at the MIR promoters through chromatin immunoprecipitation (ChIP) assays in mac3a mac3b and Col performed using anti-RPB2 antibody. qPCR analysis did not detect an obvious difference of Pol II occupancy at various MIR promoters between mac3a mac3b and Col (Figure 3E). Taken together, these results suggest that MAC3A and MAC3B do not affect MIR transcription.
MAC3A and MAC3B associate with the DCL1 complex

To further understand how MAC3A and MAC3B affect miRNA biogenesis, we examined the effect of mac3a mac3b on the expression of DCL1, DDL, SE, HYL1, CBP20/80 and HEN1, which are known to function in miRNA biogenesis. The transcript levels of HYL1, and CBP20/80 were slightly increased, while the abundance of DDL transcripts was marginally reduced (Supplemental Figure 3A). In addition, the levels of DCL1, HEN1 and SE did not show significant change. Immunoblot analyses further showed that the protein levels of SE and DCL1 were not changed in mac3a mac3b whereas the HYL1 protein was slightly increased in abundance (Supplemental Figure 3B). Moreover, we also examined the effect of mac3a mac3b on the splicing of DCL1, DDL, HEN1, HYL1 and SE using RT-PCR with primers targeting a
subset of introns (Supplemental Figure 3C). MAC3A and MAC3B did not have an obvious
effect on the splicing of these introns (Supplemental Figures 3C and 3D). However, it is not clear
if MAC3A and MAC3B affect the splicing of other introns in these examined genes.

Since MAC3A and MAC3B are components of the MAC, we suspected that like CDC5 and
PRL1, MAC3A and MAC3B might also interact with the DCL1 complex. We performed a
bimolecular fluorescence complementation (BiFC) assay to test this possibility. In the leaves of
N. benthamiana transiently co-expressing MAC3A or MAC3B fused with the C-terminal
fragment of cyan fluorescent protein (cCFP) with CDC5, PRL1, DCL1 or SE fused with the N-
terminal fragment of Venus (nVenus), yellow fluorescence signals were observed (shown in
green color; Figure 4A and Supplemental Figure 4). BiFC signals of MAC3A or MAC3B with
PRL1, DCL1 and SE were localized at the discrete bodies (Figure 4A and Supplemental Figure
4). Interestingly, the interaction between MAC3A/3B and CDC5 produced not only discrete
signals but also diffused ones, agreeing with the role of MAC in mRNA splicing (Figure 4A and
Supplemental Figure 4). Co-expression cCFP-MAC3A or cCFP-3B with nVenus-HYL1 resulted
in weak and diffused YFP signals (Figure 4A and Supplemental Figure 4), consistent with the
observation that CDC5 and PRL1 do not co-immunoprecipitate with HYL1 (Zhang et al., 2014).

Next, we used co-IP to confirm the interaction of MAC3A with CDC5, PRL1, DCL1 and SE.
We first co-expressed MYC-MAC3A with CDC5-YFP, PRL1-YFP or YFP and performed IP
with anti-YFP antibodies. MYC3A was detected in CDC5-YFP and PRL1-YFP precipitates, but
not in YFP precipitates (Figure 4B and 4C), confirming the interaction of MAC3A with CDC5
and PRL1. We next co-expressed MAC3A-GFP or GFP with MYC-DCL1 or MYC-SE and
tested the interaction of co-expressed proteins. MAC3A-YFP, but not YFP, co-IPed with MYC-
DCL1 and MYC-SE (Figure 4D and 4E). Furthermore, RNAse A treatment did not disrupt the
interaction of MAC3A with DCL1 and SE (Figure 4B-4D). These results suggest that MAC3A
and MAC3B associate with the DCL1 complex in an RNA-independent manner.

mac3a mac3b reduces pri-miRNA processing in vitro

The association of MAC3A and MAC3B suggests that they may modulate DCL1 activity.
We used an *in vitro* pri-miRNA processing assay to test this possibility. As previously described, we first generated a radiolabeled pri-miR162b (*MIR162b*) composed of the stem-loop of...
miR162b with 6-nt arms at each end using in vitro transcription (Figure 5A). Processing of miR162b was then tested in the protein extracts from young flowers of mac3a mac3b or Col. The production of miR162b from MIR162b was reduced in the protein extracts of mac3a mac3b relative to Col (Figure 5B). At 50 min and 100 min time points, the levels of miR162 generated in mac3a mac3b were ~20% of those produced in Col (Figure 5C). These results suggest that MAC3A/3B may be required for the optimal activity of the DCL1 complex.

MAC3A binds pri-miRNAs in vivo

The WD domain of MAC3A and MAC3B is known to mediate protein–protein interaction. However, it can also interact with RNAs (Lau et al., 2009). Thus, it is possible that MAC3A and
MAC3B could bind pri-miRNAs. To test this hypothesis, we performed an RNA immunoprecipitation assay (RIP) on seedlings of the mac3a mac3b complementation line harboring the MYC-MAC3A transgene (Ren et al., 2012). Following cross-linking, nuclear isolation, and immunoprecipitation, we examined the presence of pri-miRNAs in MAC3A IPs using RT-PCR. All examined pri-miRNAs, but not the control EIF4A RNAs, were enriched in the MAC3A IPs (Figure 5D). By contrast, pri-miRNAs were not detected in the no-antibody controls (Figure 5D). These results suggest that MAC3A/3B associates with pri-miRNAs in vivo.

Next, we tested if MAC3A could directly bind pri-miRNA in vitro using the RNA pull-down assay (Ren et al., 2012). In this assay, MBP and recombinant MAC3A fused with maltose-binding protein (MBP) at its N-terminus (MBP-MAC3A) were expressed in E. coli, purified with amylose resin, and then incubated with [32P]-labeled MIR162b (Supplemental Figure 5A and Figure 5E). After washing, neither MBP-MAC3A nor MBP retained MIR162b (Figure 5E). MBP-MAC3A also did not interact with a ~100-nt single-stranded RNA (ssRNA), which was generated through in vitro transcription using a N-terminal fragment of the UBIQUITIN 5 (N-UBQ5), or a dsRNA generated through annealing of sense and anti-sense strands of N-UBQ5 (Figure 5E). Because MAC3A activity needs phosphorylation (see below), we treated the recombinant MAC3A protein with extracts from Col (see below) to modify the protein and then tested its interaction with MIR162b. The modified MAC3A also did not interact with RNAs (Supplemental Figure 5B). These results suggest that MAC3A is not an RNA-binding protein.

MAC3A and MAC3B are required for the localization of HYL1 in D-bodies

The interaction of MAC3A/B with the DCL1 complex also prompted us to test the effect of mac3a mac3b on the formation of the D-body. We crossed a HYL1-YFP transgenic line, which has been used as a reporter for the D-body (Wang et al., 2013; Wu et al., 2013; Qiao et al., 2015), into mac3a mac3b and examined the percentage of cells containing D-bodies in the root tips and elongation region. As previously reported (Wu et al., 2013), the HYL1-containing D-bodies existed in most cells (~84%, Figure 6A, 6B and Supplemental Figure 6A and 6B) in WT. By contrast, D-bodies were observed in only ~26% of cells in mac3a mac3b. This result demonstrates that MAC3A and MAC3B are required for correct HYL1 localization, indicating their potential role in facilitating D-body formation.
MAC3A is a U-Box ubiquitin E3 ligase whose activity depends on phosphorylation

Both MAC3A and MAC3B contain an N-terminal ligase U-box domain that confers E3 ubiquitin ligase activity and recruits the E2 conjugating enzyme, a coiled-coil region that exists in all Prp19 homologs, mediates the tetramerization of Prp19 and interacts with CDC5L and SFP27 in metazoans, and a C-terminal WD domain composed of seven WD repeats that is required for substrate recruitment (Figure 7A). Homologous of MAC3A and MAC3B exist in all plants, while their copy numbers vary among different genomes (Supplemental Figure 7 and Supplemental Data Set 2).
Because MAC3A has considerable sequence difference from MAC3B, we tested if it is a ubiquitin E3 ligase using MBP-MAC3A (Supplemental Figure 5A). We examined the E3 ligase activity in the presence of ubiquitin, the ubiquitin-activating enzyme (E1) SlUBA and the ubiquitin-conjugating enzyme (E2) UBC8 (Zhou et al., 2017). However, MBP-MAC3A displayed only weak activity (Figure 7B). We suspected that like some other E3 ligases, MAC3A activity might depend on post-translational modification (Wang et al., 2015). Thus, we treated MBP-MAC3A and MBP protein with total protein extracts from inflorescences of \(mac3a\) \(mac3b\) to avoid contamination from endogenous MAC3A/3B, since MAC3A potentially interacts with MAC3B. The treatment greatly improved MAC3A activity (Figure 7B). Notably, Alkaline Phosphatase (Calf intestinal phosphatase, CIP) treatment of MAC3A after incubation with \(mac3a\) \(mac3b\) protein extracts completely eliminated MAC3A activity (Figure 7B). These results demonstrate that MAC3A is a \textit{bona fide} ubiquitin E3 ligase and that its activity depends on protein phosphorylation.
The ubiquitin ligase activity of MAC3A is required for miRNA and pri-miRNA accumulation

Since MAC3A is a ubiquitin ligase, we next asked if its function in miRNA biogenesis requires this activity. Based on the fact that the U-box domain of Prp19-like family is conserved in eukaryotes (Ohi et al., 2003), we generated two mutant versions of MAC3A in U-box domain through site-directed mutagenesis. In one mutant, the conserved amino acids of Tyrosine (Y) at position 23 and Glutamic acid (E) at position 24 were replaced with Glycine (G) and Alanine (A) (MAC3AMut1), respectively, while in the other one, the conserved amino acids of Histidine (H) at position 31 and Aspartic acid (D) at position 34 were replaced with Alanines (AA) (MAC3AMut2) (Figure 8A). These two mutations disrupted the ubiquitin ligase activity of MAC3A (Supplemental Figure 8A). To evaluate the effect of MAC3AMut1 and MAC3AMut2 on miRNA biogenesis, we generated stable transgenic lines in mac3a mac3b expressing MAC3AMut1 or MAC3AMut2 under the control of 35S promoter. The expression MAC3AMut1 or MAC3AMut2 did not rescue the developmental defects of mac3a mac3b (Supplemental Figure 8B and Figure 8B). Agreeing with this observation, the accumulation of both pri-miRNAs and miRNAs in mac3a mac3b was not recovered by MAC3AMut1 or MAC3AMut2 (Figure 8C and 8D). These results suggest that the ubiquitin ligase activity of MAC3A is required for miRNA biogenesis.
MAC3A and MAC3B are conserved U-box type ubiquitin E3 ligases. In plants, MAC3A and MAC3B play important roles in plant immunity and development, and their counterparts in other organisms are required for splicing. In Arabidopsis, the MAC also associates with the spliceosome. However, only a few genes display moderated splicing in defects in mac3a mac3b (Monaghan et al., 2010; Xu et al., 2012). Consequently, how MAC3A and MAC3B regulate development and immunity remains elusive. In this study, we show that the accumulation of miRNA is reduced in mac3a mac3b. Furthermore, MAC3A and MAC3B associate with the DCL1 complex and pri-miRNAs. These results suggest that MAC3A and MAC3B are important players in miRNA biogenesis, in addition to their role in splicing. Impaired miRNA biogenesis...
may partially explain the pleiotropic developmental defects of mac3a mac3b, since miRNAs target many genes that are required for proper development.

There are at least three possible explanations for the decreased pri-miRNA levels in mac3a mac3b. First, mac3a mac3b may have reduced MIR transcription. The facts that mac3a mac3b does not show altered MIR promoter activity and that MAC3A does not co-IP with Pol II suggest that MAC3A and MAC3B may not affect MIR transcription. However, we cannot rule out the possibility that MAC3A and MAC3B influence MIR elongation or termination. Second, enhanced pri-miRNA processing in mac3a mac3b may also decrease pri-miRNA accumulation. However, reduced pri-miRNA processing is observed in mac3a mac3b, arguing against this possibility. Third, mac3a mac3b may have reduced stability of pri-miRNAs (Figure 9). We give this option more weight, given the observations that MAC3A associates with pri-miRNAs in vivo and interacts with PRL1, which protects pri-miRNAs from degradation. It is reasonable to speculate that MAC3A may stabilize pri-miRNAs through modulating the function of PRL1.

Indeed, it has been observed that the interaction between PRP19 (a MAC3A ortholog) and the RNA-binding protein CWC2 is required for the stabilization of small nuclear RNAs (snRNAs) related to splicing in yeast (McGrail et al., 2009; Vander Kooi et al., 2010).
MAC3A/MAC3B interacts with DCL1 and SE but appears to have weak or no association with HYL1. Interestingly, a lack of MAC3A and MAC3B impairs the localization of HYL1 at the D-body. How does this happen? One possibility is the decreased pri-miRNAs in *mac3a mac3b* may affect the formation of D-body. However, loss-of-function mutants *mos2* and *pinp1*, in which the levels of pri-miRNAs are increased or unaltered, respectively, also display impaired HYL1 localization or D-body assembly, arguing against this possibility. In human, PRP19-mediated ubiquitination regulates the protein–protein interaction of the spliceosome, which is important for the spliceosome assembly (Das et al., 2017). In addition, PRP19 also promotes the recruitment of ATRIP (a kinase) to the DNA damage site through modifying DNA replication protein A (Marechal et al., 2014). Thus, it is possible that MAC3A and MAC3B may influence the recruitment of HYL1 through modifying proteins involved in D-body assembly. Alternatively, they may co-transcriptionally facilitate the recruitment of the D-body to the
processing site of pri-miRNAs (Figure 9). The association of MAC3A/3B with the DCL1 complex is consistent with these hypotheses.

Pri-miRNA processing is also reduced in mac3a mac3b. This cannot be attributed to altered expression of genes involved in miRNA biogenesis, as the levels of these genes are either slightly increased or unaltered in mac3a mac3b. We have shown the CDC5 promotes DCL1 activity through its interaction with the regulatory domains of DCL1 (Zhang et al., 2013), while PRL1 functions an accessory factor to facilitate CDC5 function in modulating DCL1 activity (Zhang et al., 2014). By analogy, MAC3A and MAC3B may function as components of the MAC to directly or indirectly enhance the DCL1 activity (Figure 9). Alternatively, impaired HYL1 localization or D-body formation may affect the DCL1 activity.
In summary, we find that MAC3A and MAC3B, two core components of the MAC, act redundantly in miRNA biogenesis. They associate with the DCL1 complex, positively modulate pri-miRNA accumulation, facilitate HYL1 localization at the D-body and enhance DCL1 activity. More importantly, we show that MAC3A is a phosphorylation-dependent ubiquitin ligase and that this ligase activity is required for miRNA biogenesis. This result indicates that certain signals may modulate MAC3A activity through phosphorylation and thereby regulate miRNA accumulation. The involvement of four MAC core components in miRNA biogenesis suggests that the MAC functions as a complex to promote miRNA biogenesis.

Besides core components, the MAC also contains at least 13 accessory components. The core and accessory components of the MAC are proteins with diversified functions, such as transcription factors, RNA-binding proteins, ubiquitin ligase, helicases, chromatin protein, WD proteins, protein–protein interaction regulators, coiled-coil domain-containing proteins and zinc-finger-domain-containing proteins. Moreover, the accessory components are dynamically associated with the core complex, and sub-complexes with different functions are often formed. Thus, it is likely that various MAC components act individually and coordinately in miRNA biogenesis through influencing pri-miRNA transcription, processing, and stability and/or likely have a role in the assembly of D-body (Figure 9), which resembles the diversified function of PRP19 in splicing. Consistent with this notion, CDC5 and PRL1 contribute differently to pri-miRNA accumulation but act as a complex to regulate DCL1 activity. It will be interesting to further determine the functional mechanism of these proteins as individual components and as a complex in miRNA biogenesis. The functions of the PRP19 complex from metazoans in splicing, transcription, chromatin stability and lipid droplet biogenesis have been well documented (Chanarat and Strasser, 2013). However, its function in metazoan miRNA biogenesis is unknown. Given the fact that all four MAC components associate with SE, an ortholog of ARS2, which is a key component of miRNA biogenesis in metazoa, it will not be surprising if the PRP19 complex plays a role in metazoan miRNA biogenesis.
METHODS

Plant materials and growth conditions

SALK_089300 (mac3a) (Monaghan et al., 2009) and SALK_050811 (mac3b) were obtained from the Arabidopsis Biological Resources Center (ABRC). They are in the Columbia (Col) genetic background. Transgenic lines containing a single copy of proMIR167a:GUS or pro35S:HYL1-YFP were crossed to mac3a mac3b. In the F2 generation, WT plants or mac3a mac3b harboring proMIR167a:GUS or pro35S:HYL1-YFP were selected through PCR-based genotyping for mac3a, mac3b, GUS or GFP. ~15 WT or mac3a mac3b plants were pooled for GUS transcript level analyses. All plants were grown at 22°C with 16 hour light (cool white fluorescent lamps, 25-W Sylvania 21942 FO25/741/ECO T8 linear tube) and 8 hour dark cycles.

Plasmid construction

A DNA fragment containing 2066 bp promoter and 3841bp coding region of MAC3A was PCR amplified using DNAs from Col as templates with the primers of proMAC3A-4F and MAC3Acds-1R. The resulting PCR product was cloned into pENTR/D-TOPO vector and subsequently cloned into pMDC163 binary vector to generate the proMAC3A:MAC3A-GUS plasmid. The MAC3A full-length cDNA was RT-PCR amplified with the primers of MAC3Acds-1F and MAC3Acds-1R, cloned into pENTR/D-TOPO vector and subcloned into pEarleyGate203 or pMDC83 to generate the pro35S:MYC-MAC3A construct or the pro35S:MAC3A-GFP construct. The MAC3B full-length cDNA was amplified with the primers of MAC3Bcds-1F and MAC3Bcds-1R by RT-PCR and cloned into pENTR/D-TOPO vector and subcloned into pMDC83 to generate the pro35S:MAC3B-GFP construct. To construct cCFP-MAC3A or cCFP-MAC3B, MAC3A cDNA or MAC3B cDNA was PCR amplified using the primer pair MAC3A-3F/2R or MAC3B-3F/2R, respectively, and cloned into pSAT4-cCFP-C vector. Then, the pro35S:cCFP-MAC3A fragment or the pro35S:cCFP-MAC3B fragment was released by I-SceI restriction enzyme digestion and subcloned to pPZP-RCS2-ocs-bar-RI vector. The constructs cCFP-SE, nVenus-DCL1, nVenus-HYL1, nVenus-SE, and nVenus-AGO1 were described previously (Ren et al., 2012). To construct MBP-MAC3A, the MAC3A cDNA sequence was amplified with primer MAC3A-5F(Not1) and MAC3A-5R(Sal1) and subsequently inserted into the pMAL-C5X vector. Site-mutagenesis of MAC3A was performed according the protocol of QuikChange II Site-Directed Mutagenesis Kit (Agilent). The primers are list in Supplemental
Plant complementation

The proMAC3A:MAC3A-GUS, pro35S:MYC-MAC3A, pro35S:MAC3AMut1-GFP, pro35S:MAC3AMut2-GFP, and pro35S:MAC3B-GFP plasmids were transformed into mac3a mac3b using Agrobacterium-mediated floral dip method, respectively. The transgenic plants harboring proMAC3A:MAC3A-GUS, pro35S:MAC3AMut1-GFP, pro35S:MAC3AMut2-GFP, or pro35S:MAC3B-GFP were selected on MS medium containing hygromycin (30 μg/mL). pro35S:MYC-MAC3A transformants were selected by spraying seedlings with 120 mg/L BASTA solution.

E3 ubiquitin ligase activity assay

MBP- tagged fusion proteins were expressed in E. coli strain BL21 (DE3) and purified with Amylose Resin (E8021S; NEB) by following the protocol provided by the manufacturer. The purified proteins were further desalted and concentrated using the Amicon Centrifugal Filter (Millipore). The concentration of purified protein was determined using protein assay agent (Bio-Rad).

The in vitro ubiquitination assay was performed as described (Zhou et al., 2017). Briefly, the components of 3 μg FLAG-ubiquitin, 40 ng E1 (GST-SlUBA1), 120 ng 6xHIS-AtUBC8 with 4 μg MBP, MBP-MAC3A, MBP-MAC3AMut1, or MBP-MAC3AMut1 proteins were added to a 30 μL reaction buffer [50 mM Tris-HCl pH7.5, 5 mM ATP, 5 mM MgCl2, 2 mM dithiothreitol (DTT), 3 mM creatine phosphate, 5 μg/ml creatine phosphokinase]. To detect the influence of protein modification on MAC3A activity, the recombinant proteins were treated as previously described with modifications (Wang et al., 2015). Briefly, 4 μg MBP, MBP-MAC3A, MBP-MAC3AMut1, or MBP-MAC3AMut1-bound amylose resin were incubated with the total protein extracts from mac3a mac3b for one hour at room temperature followed by extensively washing for three times. Following treatment, half of protein-bound resin was treated with calf intestinal alkaline phosphatase (CIP; NEB) for 30 min, while the other half was incubated with reaction buffer without CIP. After washing, protein-bounded resins were used to perform ubiquitin assay as described above. The reaction was terminated by addition of SDS sample loading buffer with
100 mM DTT. FLAG-ubiquitin and MBP-MAC3A were then detected with a mouse monoclonal anti-FLAG M2-peroxidase-conjugated antibody (A8592, Sigma-Aldrich) and anti-MBP antibody (E8030, NEB), respectively.

**Co-IP Assay**

To test the interaction between MAC3A and RPB2, anti-RPB2 antibody was used to perform IP on the protein extracts from inflorescences of transgenic plants harboring pro35S:MAC3A-GFP (Ren et al., 2012). After IP, MAC3A-GFP and RPB2 were detected by immunoblot using an anti-GFP monoclonal antibody (B230720, Biolegend) and anti-RPB2 antibodies (ab10338, Abcam). To examine the co-IP of MAC3A with CDC5, and PRL1, MYC-MAC3A was co-expressed with YFP, CDC5-YFP or PRL1-YFP in *N. benthamiana* as described (Ren et al., 2012). To examine the co-IP of MAC3A with DCL1 and SE, MAC3A-YFP was co-expressed with MYC-DCL1 or MYC-SE in *N. benthamiana*. IP was performed on protein extracts using anti-GFP or anti-MYC antibodies coupled to protein G agarose beads as described (Ren et al., 2012). After IP, proteins were detected with immunoblotting using monoclonal antibodies against YFP (B230720, Biolegend) or MYC (06-340, Millipore).

**ChIP assay**

ChIP was performed using 14-d-old seedlings from Col-0 and mac3a mac3b as described (Kim et al., 2011). Three biological replicates were performed. Anti-RPB2 antibody (ab10338, Abcam) was used for immunoprecipitation. qPCR was performed using primers listed in Supplemental Table 1.

**Dicer Activity Assay**

*In vitro* MIR162b processing assay was performed as described (Qi et al., 2005; Ren et al., 2012). DNA templates used for *in vitro* transcription were generated through PCR with primers listed in Supplemental Table 1. *In vitro* transcription of MIR162b, N-UBQ5 and anti-sense N-UBQ5 were performed using T7 RNA polymerase in the presence of [α-32P] UTP, ATP, CTP, GTP and unlabeled UTP. MIR162b was processed in reaction buffer (100 mM NaCl, 1 mM ATP, 0.2 mM GTP, 1.2 mM MgCl₂, 25 mM creatine phosphate, 30 μg/ml creatine kinase and 4 U
RNase inhibitor) containing 30 μg protein at 25 °C. After the reaction was stopped at 50 or 100 minutes, RNAs were extracted and separated on a PAGE gel. ImageQuant was used to quantify the radioactive signals detected by a PhosphorImager.

**Morphological analyses and GUS histochemical staining**

Morphological and cellular analyses were performed according to the previously reported methods (Li et al., 2012). GUS staining was performed as described (Zhang et al., 2013). Briefly, tissues from plants of *mac3a mac3b* harboring *proMAC3A:MAC3A-GUS* or plants harboring *promIR167a:GUS* were incubated with staining solution at 37 °C for 5 hours. 70% ethanol was used for tissue clearing before imaging.

**BiFC Assay**

BiFC assay was performed as described (Zhang et al., 2013). Paired cCFP and nVenus fusion proteins were co-expressed in *N. benthamiana* leaves. After 40 h expression, a confocal microscope (Fluoview 500 workstation; Olympus) was used to detect YFP and chlorophyll autofluorescence signals at 488 nm with a narrow barrier (505–525 nm, BA505-525; Olympus).

**RNA gel blot and RT-qPCR analyses**

RNA gel blotting was performed as described (Ren et al., 2012). ~15 μg total RNAs extracted from inflorescences were resolved on 16% PAGE gel and transferred to nylon membranes. 32P-labelled antisense DNA oligonucleotides were used to detect small RNAs. Radioactive signals were detected with a Phosphorimager and quantified with ImageQuant. Inflorescences of plants grown on three different growth rooms at the same condition (22°C with 16 hour light and 8 hour dark cycles) were harvested as three replicates. The levels of pri-miRNAs, miRNA target transcripts and GUS mRNA were determined using RT-qPCR. 1 μg total RNAs from inflorescences were used to generate cDNAs using the SuperScript III reverse transcriptase (Invitrogen) and an oligo dT18 primer. cDNAs were then used as templates for qPCR on an iCycler apparatus (Bio-Rad) with the SYBR green kit (Bio-Rad). The primers used for PCR are listed on Supplemental Table 1.

**RNA immunoprecipitation (RIP) analyses**
RIP was performed according to (Wierzbicki et al., 2008; Ren et al., 2012). ~ 2g seedlings of transgenic plants harboring the pro35S:MYC-MAC3A transgene were used to examine the association of MAC3A with pri-miRNAs \textit{in vivo}. After crosslinking with 1% formaldehyde for 10 min, glycine was added to quench the reaction for 10 min. Nuclei were extracted and lysed in the buffer (50 mM Tris-HCl, pH8.0, 10 mM EDTA, 1% SDS) by sonication for five times. After debris was removed by centrifugation at 16,000g for 10 min, equal amounts of proteins from various samples were diluted with RIP dilution buffer and incubated with anti-GFP antibodies conjugated to protein-G agarose beads. The immunoprecipitates were then eluted with elution buffer (100 mM NaHCO₃, 1% SDS) at 65 °C. Following reverse crosslinking with proteinase K (Invitrogen) and 200 mM NaCl at 65 °C, RNAs were extracted and used as templates for RT-PCR analyses. All the primers are listed in Supplemental Table 1.

\textit{In vitro} RNA pull-down assay

\textit{In vitro} RNA pull-down assay was performed as described (Ren et al., 2012). The amylose resin beads containing MBP or MBP-MAC3A were incubated with [³²P]-labeled probes at 4°C for 1 hour. After the beads were washed for 4 times, RNAs were extracted and resolved on PAGE gels. Radioactive signals were detected with a PhosphorImager and quantified by ImageQuant.

Small RNA sequencing

Inflorescences of Col, \textit{mac3a mac3b} and \textit{cdc5-1} grown on two separate growth rooms at the same condition (22°C with 16 hour light and 8 hour dark cycles) were harvested as two biological replicates and used for RNA extraction and small RNA library preparation following standard protocol. The data set was deposited into the National Center for Biotechnology Information Gene Expression Omnibus (Col accession #: GSM2829820, GSM2829821, \textit{mac3a mac3b} accession # GSM2829822, GSM2829823; Col accession #: GSM2805383, GSM2805384, \textit{cdc5-1} accession #: GSM2805385, GSM2805386). The sequencing data (Col access #: GSM2257315, GSM2257316, GSM2257317; \textit{dcl1} accession #: GSM2257321, GSM2257322, GSM2257323) generated by Wu et al., (Wu et al., 2016) were used to analyze the effect of DCL1 on miRNA accumulation. After sequencing, miRNA analysis was performed after removing reads aligned to t/r/sn/snoRNA according to Ren et al (Ren et al., 2012). Normalization was done using the total numbers of perfectly aligned reads (Nobuta et al., 2010).
The mean values of miRNA abundance from biological replicates were compared by using EdgeR with trimmed mean of M values (TMM) normalization method (Robinson et al., 2010). Down-regulated miRNAs with confidence (P<0.1; fold < 0.7) were used to identify the overlapping effect of mac3a mac3b, cdc5-1 and dcl1-9. The Venn diagram was plotted with the VennDiagram from the R package (Chen and Boutros, 2011).

**Accession Numbers**

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: MAC3A (AT1G04510), MAC3B (AT2G33340), CDC5 (AT1G09770), PRL1 (AT4G15900), DCL1 (AT1G01040), SE (AT2G27100), HYL1 (AT1G09700), DDL (AT3G20550), CBP20 (AT5G44200), CBP80 (AT2G13540), HEN1 (AT4G20910), AGO1 (AT1G48410), ARF4 (AT5G60450), ARF8 (AT5G37020), CKB3 (AT3G60250), CUC1 (AT3G15170), MYB33 (AT5G06100), PHV (AT1G30490), PHO2 (AT2G33770), PPR (AT1G62670), SPL9 (AT2G42200), SPL10 (AT1G27370), SPL13 (AT5G50570), UBIQUITIN5 (AT3G62250). Protein sequences of MAC3 homologs in other species can be obtained in National Center for Biotechnology Information under the following accession numbers: AAN13133 (MAC3A, AT1G04510, Arabidopsis thaliana), FJ820118 (MAC3B, AT2G33340, Arabidopsis thaliana), XP_009143870 (Brassica rapa), XP_009141306 (Brassica rapa), XP_004247768 (Solanum lycopersicum), XP_003555746 (Glycine max), XP_003535988 (Glycine max), XP_015614850 (Oryza sativa), KXG38386 (SORBI_3001G226000, Sorghum bicolor), ONM06005 (ZEAMMB73_Zm00001d032763, Zea mays), AOK6571 (ZEAMMB73_Zm00001d014078, Zea mays), XP_001701820 (Chlamydomonas reinhardtii), NP_055317 (HsPRP19, Homo sapiens), NP_598890 (MmPRP19, Mus musculus), CAB10135 (SpPRP19, Shizosaccharomyces pombe), and CA997487 (ScPRP19, Saccharomyces cerevisae). Small RNA deep sequencing datasets are available from the National Center for Biotechnology Information Gene Expression Omnibus under the following reference numbers: Col accession #: GSM2829820, GSM2829821, mac3a mac3b accession # GSM2829822, GSM2829823; Col accession #: GSM2805383, GSM2805384, cdc5-1 accession #: GSM2805385, GSM2805386; Col access #: GSM2257315, GSM2257316, GSM2257317; dcl1 accession #: GSM2257321, GSM2257322, GSM2257323.

**Supplemental Data**
Supplemental Figure 1. Small RNA sequencing analyses of mac3a mac3b, cdc5 and dcl1-9. (Supports Figure 2)

Supplemental Figure 2. Expression of MAC3A and MAC3B complements the defects of mac3a mac3b. (Supports Figure 2)

Supplemental Figure 3. Effect of MAC3A and MAC3B on the expression levels and splicing of genes involved in miRNA biogenesis. (Supports Figure 4)

Supplemental Figure 4. Interaction of MAC3B with CDC5, PRL1 and the DCL1 complex detected by BiFC analysis. (Supports Figure 4)

Supplemental Figure 5. RNA-binding activity of MAC3A. (Supports Figure 5)

Supplemental Figure 6. HYL1-YFP localization in root tips in Col and mac3a mac3b mutant. (Supports Figure 6).

Supplemental Figure 7. Phylogenetic analysis of MAC3A orthologs. (Supports Figure 7)

Supplemental Figure 8. Ubiquitin ligase activity of MAC3A is required for miRNA biogenesis. (Supports Figure 8)

Supplemental Table 1. The sequences of oligonucleotides.

Supplemental Data Set 1. miRNA profile change in mac3a mac3b, cdc5-1, and dcl1-9 relative to wild-type plants as determined by small RNA sequencing. (Supports Supplemental Figure 1)

Supplemental Data Set 2. Text file of the alignment used for the phylogenetic analysis shown in Supplemental Figure 7. (Supports Figure 7)

Supplemental Data Set 3. Results of statistical analyses (Supports Figures 1, 2, 3, 5, 6, 8, and Supplemental Figures 2, 3)

Acknowledgements

This work was supported by the Nebraska Soybean Board (Award #1727 to B.Y), the National Science Foundation (Awards OIA- 1557417 to B.Y and IOS-1645659 to L.Z), and the Pioneer Hundred Talents Program of Chinese Academy of Sciences (to S.L).

Author Contributions

S.L. and B.Y. designed the experiments and prepared the manuscript. S.L., B.Y., K.L., B.Z., M.L., S.Z., L.Z. and C.Z. performed the experiments. S.L., C.Z. and B.Y. analyzed the data.
Competing Financial Interests

The authors declare no competing financial interests.


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MAC3A and MAC3B, Two Core Subunits of the MOS4-Associated Complex, Positively Influence miRNA Biogenesis
Shengjun Li, Kan Liu, Bangjun Zhou, Mu Li, Shuxin Zhang, Lirong Zeng, Chi Zhang and Bin Yu
*Plant Cell*; originally published online February 5, 2018;
DOI 10.1105/tpc.17.00953

This information is current as of February 8, 2018

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