

***IAA-Ala Resistant3*, an Evolutionarily Conserved Target of miR167, Mediates *Arabidopsis* Root Architecture Changes during High Osmotic Stress**^W

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The functions of microRNAs and their target mRNAs in *Arabidopsis thaliana* development have been widely documented; however, roles of stress-responsive microRNAs and their targets are not as well understood. Using small RNA deep sequencing and ATH1 microarrays to profile mRNAs, we identified *IAA-Ala Resistant3 (IAR3)* as a new target of miR167a. As expected, *IAR3* mRNA was cleaved at the miR167a complementary site and under high osmotic stress miR167a levels decreased, whereas *IAR3* mRNA levels increased. *IAR3* hydrolyzes an inactive form of auxin (indole-3-acetic acid [IAA]-alanine) and releases bioactive auxin (IAA), a central phytohormone for root development. In contrast with the wild type, *iar3* mutants accumulated reduced IAA levels and did not display high osmotic stress-induced root architecture changes. Transgenic plants expressing a cleavage-resistant form of *IAR3* mRNA accumulated high levels of *IAR3* mRNAs and showed increased lateral root development compared with transgenic plants expressing wild-type *IAR3*. Expression of an inducible noncoding RNA to sequester miR167a by target mimicry led to an increase in *IAR3* mRNA levels, further confirming the inverse relationship between the two partners. Sequence comparison revealed the miR167 target site on *IAR3* mRNA is conserved in evolutionarily distant plant species. Finally, we showed that *IAR3* is required for drought tolerance.

INTRODUCTION

Arabidopsis thaliana plants accumulate a multitude of microRNAs (miRNAs), which are single-stranded RNA molecules 20 to 24 nucleotides in length (Reinhart et al., 2002; Carrington and Ambros, 2003). When bound to ARGONAUTE proteins, miRNAs guide RNA-induced silencing complexes to mRNAs that harbor complementary sequences (Chapman and Carrington, 2007; Montgomery and Carrington, 2008). The RNA-induced silencing complex then inhibits gene expression through translational repression or mRNA cleavage (Llave et al., 2002; Aukerman and Sakai, 2003; Bartel, 2004; Chen, 2004). One of the challenges in the field is to identify functional miRNA-mRNA pairs and elucidate their roles in the life of a plant.

Phenotypes of mutants defective in miRNA biogenesis components, such as *dicer-like1*, *hua enhancer1*, *serrate*, and *argonaute1*, provided the first indication that miRNAs are functional in developmental processes (Jacobsen et al., 1999; Lynn et al., 1999; Golden et al., 2002; Park et al., 2002; Schauer et al., 2002; Vaucheret et al., 2004; Lobbes et al., 2006; Yang et al.,

2006; Sunkar et al., 2007). To date, functions of various miRNA-mRNA pairs have been clarified in the temporal and spatial development of flowers, leaves, and roots (Kutter et al., 2007; Chen, 2009; Poethig, 2009; Covarrubias and Reyes, 2010; Martin et al., 2010; Nonogaki, 2010; Nag and Jack, 2010; Khan et al., 2011).

Throughout their lifecycle, plants are challenged by highly variable environmental conditions, such as changes in nutrient levels and water availability. Plants exhibit adaptive adjustments in these conditions, allowing individual plants to maintain function and hence survival across a range of diverse environments (Sultan, 2000; Moldovan et al., 2010). The dynamic stress-responsive gene expressions, which underpin physiological adjustments, suggest that miRNAs are also likely to be involved in stress adaptations. Consistent with this notion, a mutation in HYPONASTIC LEAVES1 (HYL1), a component of the miRNA biogenesis machinery, caused altered perception of abscisic acid (ABA). Because ABA is involved in conditions of water limitation, such as salt and drought stress, this observation possibly implicates HYL1 in stress responses (Lu and Fedoroff, 2000; Han et al., 2004; Vazquez et al., 2004).

Potential involvement of a number of miRNAs has been suggested for responses to water-limiting conditions (e.g., salt, drought, and ABA responses). In particular, miR393, miR397b, and miR402, whose predicted targets are mRNAs encoding auxin receptor TRANSPORT INHIBITOR RESPONSE1, LACCASE, and DEMETER-LIKE PROTEIN3, respectively, were upregulated in

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response to cold, dehydration, NaCl, and ABA in *Arabidopsis*, rice (*Oryza sativa*), and *Phaseolus vulgaris* (Dharmasiri and Estelle, 2002; Sunkar and Zhu, 2004; Dharmasiri et al., 2005; Zhao et al., 2007; Liu et al., 2008; Arenas-Huertero et al., 2009). However, only a few miRNA-target mRNA pairs, such as miR169-*NFYA5*, have been functionally validated in cases of water stress (Li et al., 2008).

We hypothesized there may be additional new and functional miRNA-mRNA pairs to be identified if we analyze differential changes in miRNA and mRNA expression patterns between standard conditions and water-limiting conditions. To this end, we subjected hydroponically grown *Arabidopsis* plants to high osmotic stress and determined miRNA and mRNA expression patterns by parallel analyses via deep sequencing (for small RNAs) and microarray expression profiles (mRNA levels). We found the miR167-*IAR3* (for *IAA-Ala Resistant3*) pair to be a modifying factor for root architecture under osmotic stress conditions. We showed that *IAR3* is a regulator of root architecture changes in osmotic stress and drought tolerance. Consistent with its proposed role, the miR167-*IAR3* relationship seems to be evolutionarily conserved in higher dicot and monocot species.

RESULTS

Reduced Accumulation of miR164a/b, miR167a/b, and miR172a/b under High Osmotic Stress

To identify new miRNA-mRNA pairs related to high osmotic conditions, we used a deep sequencing technique to profile small RNA populations in leaf and root tissues of plants under high osmotic stress and control conditions. We recovered around 20 million reads from each library, which corresponded to at least 18 million reads after adapter trimming, and more than 60% of these perfectly matched *Arabidopsis* genome sequences. This resulted in at least 11 million total reads for each library, which were used in subsequent analyses (see Supplemental Table 1 online). First, we searched for global changes in the population of small RNAs under high osmotic stress (see Supplemental Figure 1 online). Apart from minor changes observed among 21- and 24-nucleotide small RNAs, there were no significant differences in small RNA size distribution between the osmotic stress and control sample (see

Supplemental Figures 1A and 1B online). To eliminate the noise generated by high-copy-number small RNA species, we asked whether the same pattern would be observed if small RNA species were only counted once, without any regard to copy number. Here, again, we did not detect any significant differences between the osmotic stress and control small RNA population (see Supplemental Figures 1C and 1D online). Since there were no major global changes in the small RNA population, we searched for specific small RNAs whose accumulation was affected by stress. All annotated miRNAs and *trans*-acting small interfering RNAs were queried in our sequencing data sets, and most of them did not change under stress. However, the accumulation of miR164a/b, miR167a/b, and miR172a/b was reduced under high osmotic stress (Table 1).

RNA gel blot analyses confirmed miR167a/b accumulation was reduced under high osmotic stress compared with control in leaf samples, whereas miR391 remained at similar levels in both tissues (Figure 1A). Using a specific probe, we found that the accumulation of pri-miR167a, which is the most highly expressed miR167 precursor, was also reduced under high osmotic stress (Figure 1B). Therefore, the reduced accumulation of miR167a/b under stress was mainly due to changes at the pre-miR167 level and not due to regulation of precursor processing.

A miR167a/b mRNA Target, *IAR3*, Is Upregulated in High Osmotic Stress

Like animal small RNAs, the inhibitory effects of plant miRNAs on target gene expression may involve translation inhibition, but plant miRNAs also mediate mRNA cleavage at the complementary site (Brodersen et al., 2008; Lin et al., 2008; Pant et al., 2008). We focused on the mRNA cleavage function, which could be monitored by changes in mRNA levels.

To find new target mRNAs, we investigated expression levels of putative target mRNAs in microarrays using samples identical to those used in small RNA deep sequencing experiments. Our computer analysis predicted nine genes that were potential miR167 targets, and seven of them were represented in the ATH1 array (see Supplemental Table 2 online). Only *IAR3* showed increased mRNA levels of at least twofold under water stress (Figures 1C and 1D; see Supplemental Data Set 1 online). *IAR3* encodes an indole-3-acetic acid (IAA)-Ala hydrolase, which releases bioactive auxin (IAA) from inactive auxin storage (IAA-Ala;

Table 1. Downregulated miRNAs with mRNA Targets in High Osmotic Stress

miRNA Annotation	Normalized Read Nos.				Fold Change		Target Genes	
	Leaf		Root		Leaf	Root	Previous Studies	This Study
	Control	Osmo	Control	Osmo				
miR172a/b	1,074	164	377	257	0.01	0.35	<i>AP2</i>	
miR164a/b	1,547	811	2,404	1,771	0.27	0.38	<i>NAC1, CUC1, NAC2/ORE1</i>	
miR167a/b	39,610	19,051	8,787	7,252	0.23	0.44	<i>ARF6/8</i>	<i>IAR3</i>

Normalized read numbers (reads per million) of miR164a/b, miR167a/b, and miR172a/b in leaf and root tissues of control and high osmotic stress (Osmo) plants, as well as fold changes. Thirty-day-old plants were subjected to high osmotic stress (300 mM mannitol, osmotic stress) or 0 mM mannitol (control) for 3 h and sampled. Previously identified and putative target mRNAs are shown. This study focused on *IAR3*.

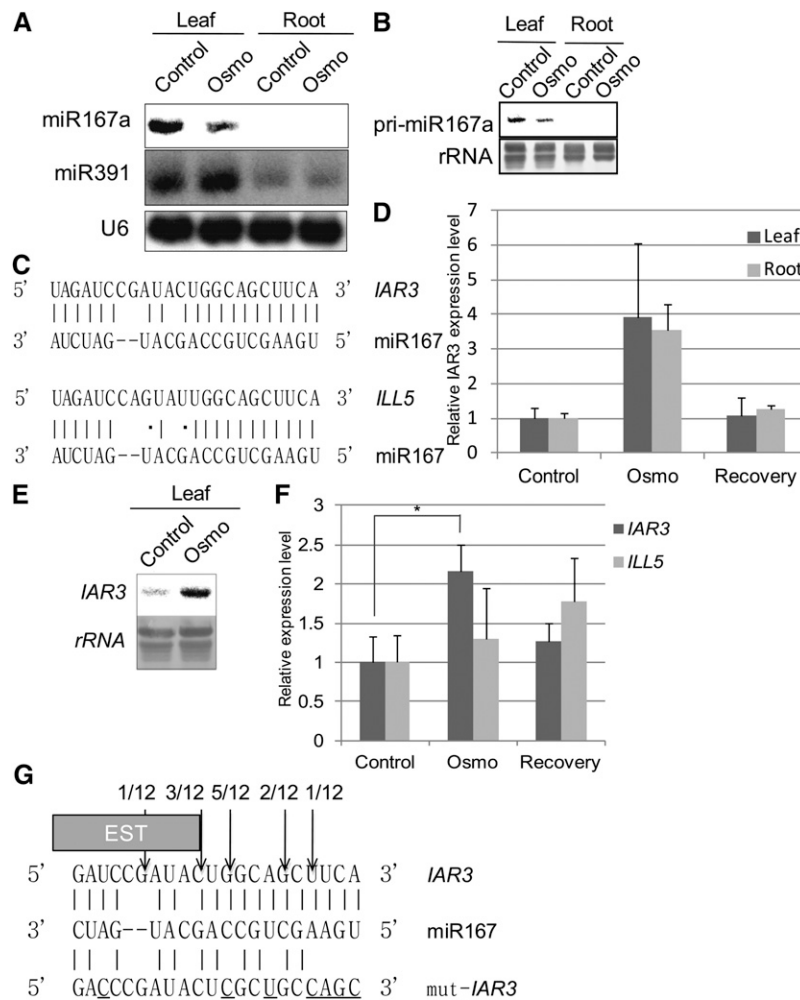


Figure 1. The *IAR3* mRNA Accumulation Pattern Is Inversely Correlated with That of miR167a/b.

(A) RNA gel blot analysis of miR167a/b and miR391 expressions in high osmotic stress (Osmo) and control leaf and root tissues. Conditions were identical to those in Table 1. U6 accumulation is shown as a loading control and each lane contained 20 μ g total RNA.

(B) RNA gel blot analysis of pre-miR167a expression in control and high osmotic stress (Osmo) leaf and root tissues. 25S rRNA was used as a loading control and each lane contained 10 μ g total RNA.

(C) Sequence complementarity of miR167a/b with *IAR3* and *ILL5* mRNAs. Hydrogen bond is shown by a vertical line, G·U wobble is shown by a dot, and gap is shown by a dash.

(D) Relative expression levels of *IAR3* mRNA in high osmotic stress samples (Osmo) compared with control from ATH1 microarray analyses. Microarray analyses were performed using identical samples as deep sequence analyses. In addition, materials that were sampled 24 h after stress conditions were analyzed to observe transient changes. These samples were designated "Recovery." Bars show SE ($n = 3$).

(E) RNA gel blot analysis of the *IAR3* mRNA expression pattern in leaf tissue. 25S rRNA was used as a loading control and each lane contained 10 μ g RNA.

(F) Relative expression patterns of *IAR3* and *ILL5* mRNAs using quantitative RT-PCR with gene-specific primers. Bars show SE ($n = 3$). Asterisks indicate a significant difference between Osmo and mock treatment, based on *t* test ($P < 0.05$).

(G) The 5' end of the cleaved product determined by sequencing is indicated by an arrow in the miRNA:mRNA base-pairing diagram, along with the number of clones analyzed. The box shows the end of a truncated *IAR3* EST clone (accession number EBENXNS01CL7RN). The bottom sequence shows a mutation strategy to generate the miR167-resistant form of *IAR3* without changing its amino acid sequence. Substituted nucleotides are underlined.

Davies et al., 1999). To evaluate a possible transient mode of mRNA accumulation, we harvested recovery tissues from plants that were returned to the normal medium for 24 h after stress treatment. In this sample, *IAR3* expression returned to control levels, strengthening our hypothesis that *IAR3* mRNA was transiently upregulated in high osmotic stress. Therefore, our

subsequent work focused on this putative miR167a/b-*IAR3* pair. To validate this experiment, we verified that genes mediating ABA signaling, such as *ABA INSENSITIVE1/2*, and downstream marker genes, such as *KIN1*, *RESPONSIVE TO DESSICATION29A/B*, and *COLD-REGULATED15A/B*, were upregulated (see Supplemental Figure 2 online).

RNA gel blot analyses confirmed increased accumulation of the putative target *IAR3* mRNA (Figure 1E). The *Arabidopsis* genome, however, also encodes *IAA-LEUCINE RESISTANT-LIKE GENE5 (ILL5)*, which is located next to *IAR3* on chromosome 1 and is 85% identical to *IAR3* at the nucleotide sequence level (Davies et al., 1999; LeClere et al., 2002). Sequence analysis of the miR167 complementary site uncovered a three-nucleotide difference between *IAR3* and *ILL5*, one occurring in the mismatched nucleotide between miR167 and *IAR3* and the other two resulting in the hydrogen bond to G·U wobble changes (Figure 1C). Our initial computer prediction did not identify *ILL5* as a potential target of miR167a/b because the total mismatch number exceeded the specified threshold.

To distinguish between *IAR3* and *ILL5* mRNAs, we designed discriminating gene primers and analyzed mRNA levels using quantitative RT-PCR. We reproduced microarray and RNA gel blot results for *IAR3* but did not observe meaningful changes in *ILL5* in response to water stress (Figure 1F). Although we cannot completely rule out the possibility that *ILL5* is also a target, our results indicated that *IAR3* was the preferred target of miR167a/b.

We noted that annealing between miR167 and *IAR3* at the target site would generate an exceptional two-nucleotide asymmetric bulge. To determine if there were alternative forms of miR167, or miR167-like small RNAs that could better align with *IAR3* mRNAs, we aligned all small RNAs in our osmotic stress and control data set to *IAR3*. We found eight miR167a-like small RNAs, which were one to three nucleotides longer than miR167a (see Supplemental Table 3 online). These small RNAs had a similar alignment to *IAR3* as miR167a. However, their read numbers were too low to be functionally significant (see Supplemental Table 4 online). Thus, miR167a seemed to be the main mediator of the *IAR3* cleavage.

***IAR3* mRNA Is Cleaved at the miR167a/b Complementary Site**

We used 5' rapid amplification of cDNA ends (RACE) to determine the cleavage site of *IAR3* mRNA. Sequence analysis of 12 independent clones placed the 5' end of the cleaved fragment in the middle of the miR167//*IAR3* mRNA complementary site (Figure 1G). We also found an EST clone (accession number EBENXNS01CL7RN), whose 3' end is located at the second highest cleavage position (Figure 1G). Together, these results demonstrate cleavage of *IAR3* mRNA at the center of the miR167a/b complementary site.

***iar3* Mutants Are Insensitive to High Osmotic Stress**

The upregulation of *IAR3* mRNA in osmotic stress indicated that *IAR3* was a potential new positive regulator in the high osmotic stress pathway. The *iar3* mutant was originally isolated by Davies et al. (1999) in their effort to identify a gene product that can convert IAA-Ala to IAA using the inhibitory effect of IAA-Ala on *Arabidopsis* root growth. Whereas primary root growth was inhibited in the wild type, *iar3* mutant primary roots continued to grow on IAA-Ala-supplemented media; hence, the name *IAA-Ala Resistant3 (IAR3)*. The root growth inhibition in both IAA-Ala-supplemented media and high osmotic conditions led us to

hypothesize that, under high osmotic stress, the upregulated *IAR3* expression might contribute to primary root growth inhibition by releasing biologically active IAA. To test this hypothesis, we obtained two knockout lines harboring independent T-DNA insertions in the *IAR3* locus (Alonso et al., 2003). These lines were designated as *iar3-5* and *iar3-6* in sequence after *iar3-1* to *iar3-4* mutants that were previously characterized by Davies et al. (1999). The two mutants, *iar3-5* and *iar3-6*, contain T-DNA insertions in the 5' untranslated region and the promoter of *IAR3*, respectively, and *IAR3* mRNA expression levels were significantly reduced in both mutants compared with the wild type (see Supplemental Figure 3 online).

Because *IAR3* is a hydrolase that releases free IAA from an inactive form (IAA-Ala), we analyzed free IAA accumulation levels in the wild type and *iar3-5* mutants. Figure 2A shows that under control conditions, *iar3-5* accumulated less IAA than the wild type, as previously reported (Rampey et al., 2004). Under high osmotic conditions, the wild type accumulated increased amounts of IAA compared with control conditions, whereas *iar3-5* accumulated reduced amounts compared with the wild type in both control and high osmotic conditions. These results suggest that upregulated *IAR3* expression levels under high osmotic conditions may contribute to increased amounts of free IAA in *Arabidopsis*.

Since IAA is a major factor regulating primary root growth and lateral root development, we compared root morphologies of *iar3* mutant and wild-type plants. Under high osmotic conditions, wild-type plants showed reduced primary root growth but increased lateral root development. By contrast, *iar3* mutants were insensitive to the effect of high osmotic stress on root development (Figure 2B). We observed reduced miR167 expression levels under stress conditions primarily in leaves, but not as clearly in roots (Figures 1A and 1B), whereas *IAR3* expression was enhanced in both leaf and root samples (Figures 1D to 1F). It could be, in osmotic stressed root samples, the reduction of miR167 levels was tissue specific and, therefore, not detectable by RNA gel blot analyses. Another possibility is that as the leaf miR167 level decreased under stress conditions, the resulting increase in *IAR3* mRNA and protein levels led to an increased amount of IAA, which was transported to the root system. Quantitative analyses showed no obvious differences in primary root growth between the wild type and *iar3* mutants under control conditions. By contrast, under high osmotic stress, both *iar3* mutant alleles displayed marked insensitivity, whereas wild-type primary root growth was severely inhibited (Figure 2C).

With respect to lateral root formation, *iar3-6* produced slightly fewer lateral roots compared with the wild type only on day 5 of control conditions; however, we did not observe any consistent and significant difference between *iar3* mutant alleles and the wild type at other time points under the same conditions (Figure 2D). Upon transfer to high osmotic conditions, the wild type exhibited robust lateral root formation, whereas *iar3* mutants were insensitive to the stress and formed fewer lateral roots (Figure 2D). In addition, *iar3-6* mutants produced fewer lateral roots than *iar3-5* mutants under stress conditions. This was probably due to the tendency of *iar3-6* to form fewer lateral roots compared with *iar3-5* under control conditions. Therefore,

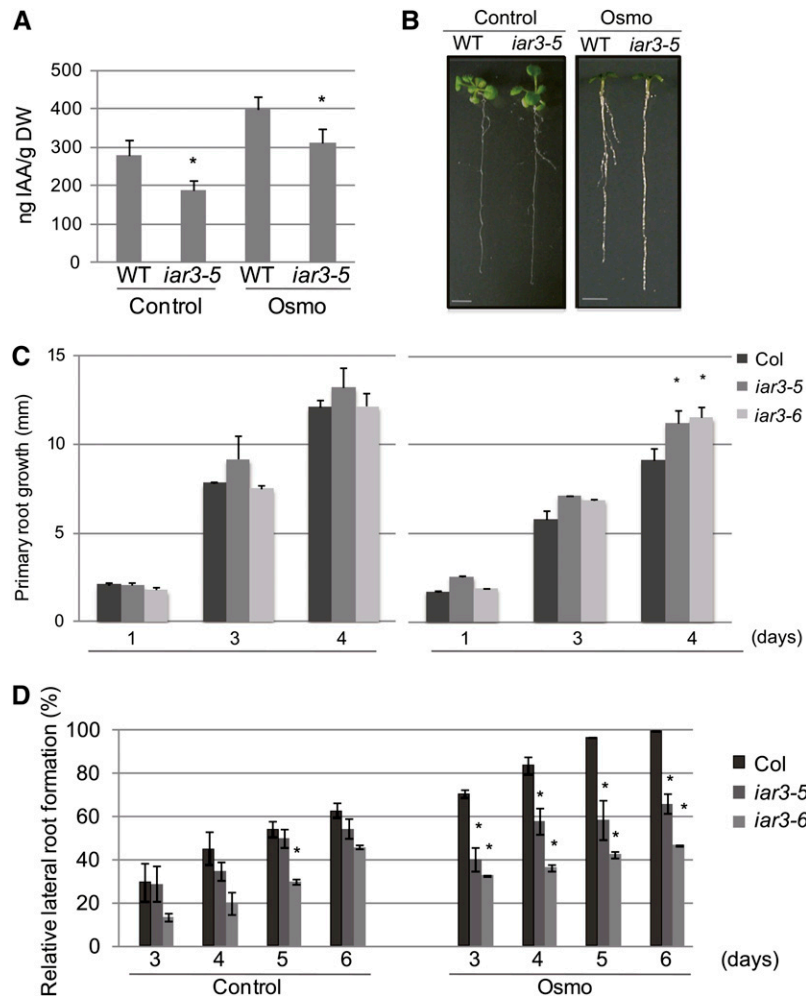


Figure 2. *iar3* Mutants Are Insensitive to Osmotic Stress.

(A) Measurement of endogenous free IAA levels by liquid chromatography–electrospray ionization–tandem mass spectrometry. Increase in dry weight (DW) due to mannitol uptake was adjusted by dividing the dry weight by the average increase in dry weight under mannitol treatment. Bars show SE ($n = 23$). Asterisks indicate a significant difference between the wild type (WT) and the mutant, based on t test ($P < 0.05$).

(B) Morphology of the wild type (Col) and *iar3-5* under high osmotic stress (right) and control (left). Wild-type and *iar3-5* plants were grown in control media for 8 d and transferred to the Osmo plate (MS containing 0.25 M mannitol) or control plate (MS alone). Pictures were taken after 1 week (Osmo) or 10 days (Control). Bar = 5 mm.

(C) Time-course experiments of the wild type, *iar3-5*, and *iar3-6*, quantifying primary root growth (mm) after seedlings were transferred to Osmo or control. Root lengths were analyzed using ImageJ (National Institutes of Health). Bars show SE ($n \geq 12$). Asterisks indicate a significant difference between the wild type and the mutant, based on t test ($P < 0.05$).

(D) Time-course experiment quantifying lateral root numbers in the wild type, *iar3-5*, and *iar3-6*. Experiments were done as in **(B)**. Lateral roots were counted under a stereomicroscope (Leica). Lateral root numbers are represented as percentage of wild-type lateral root numbers in Osmo media at day 6. Bars show SE ($n \geq 20$). Asterisks indicate a significant difference between the wild type and mutant, based on t test ($P < 0.01$).

although the two alleles seemed to produce slightly different numbers of lateral roots, their responses to stress were very similar. These results suggest that *IAR3* is a new positive regulator of high osmotic stress signaling and *iar3* mutants are compromised in their response to this stress. The simplest explanation for the underlying mechanism is that bioactive IAA released by *IAR3* contributes to the developmental changes in roots. Only one of the two alleles tested here showed a slight decrease in lateral root formation under control condition.

This is consistent with the notion that auxin is also involved in the normal course of lateral root development. Under this condition, *IAR3* is expressed at low levels but presumably plays a functional role. On the other hand, there is a greater demand for auxin under high osmotic stress conditions when plants need to develop more lateral roots. The contribution of *IAR3* to plant development becomes much greater under stress, which explains its stress-induced expression (Figures 1D to 1F).

Direct Involvement of miR167 and IAR3

To obtain evidence that miR167 directly mediates *IAR3* mRNA cleavage, we introduced seven nucleotide mismatches into the miR167 complementary site in the *IAR3* mRNA sequence without changing the encoded amino acids, and the mutant gene version was designated *mut-IAR3* (Figure 1G).

We transformed *iar3-5* mutants with either *IAR3:mut-IAR3* or *IAR3:WT-IAR3*. mRNA analyses of two independent T1 lines for each construct showed that *mut-IAR3/iar3-5* lines expressed much higher *IAR3* levels than *WT-IAR3/iar3-5* lines (Figure 3A). For this experiment, we used control conditions to allow higher miR167 accumulation levels, thus providing a more sensitive condition than that of high osmotic stress. To determine the

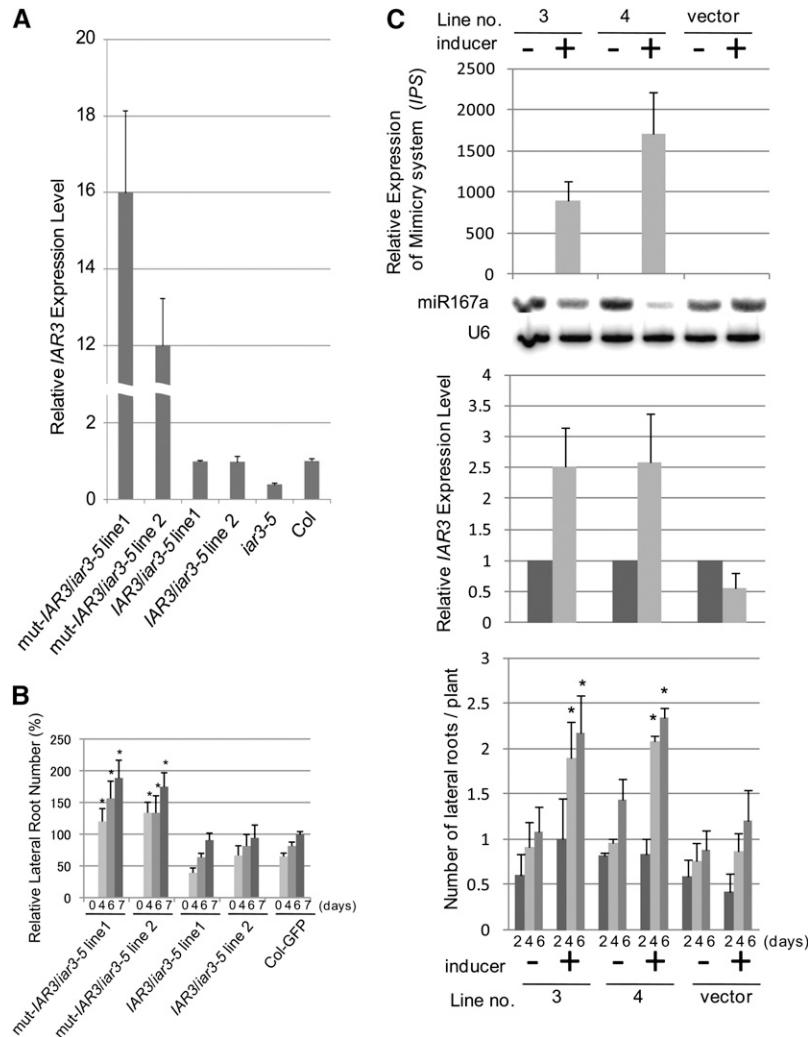


Figure 3. Relationships between miR167, *IAR3*, and Lateral Root Numbers.

(A) Relative expression levels of *IAR3* mRNAs using quantitative RT-PCR in *IAR3:IAR3/iar3-5*, *IAR3:mut-IAR3/iar3-5*, the wild type, and *iar3-5*. Bars show SE ($n = 3$). Asterisks indicate a significant difference between the wild type and *iar3-5* or transgenic lines based on *t* test ($P < 0.05$).

(B) Lateral root numbers were analyzed in *IAR3:IAR3/iar3-5*, *IAR3:mut-IAR3/iar3-5*, and 35S:*GFP* (for green fluorescent protein) control plants. Average lateral root numbers of T2 population derived from two independent T1 lines are shown. Antibiotic-resistant plants were transferred to a vertical plate after 1 week and lateral root numbers scored. At day 0 (d0) there were no lateral roots. Bars show SE ($n \geq 12$). Asterisks indicate a significant difference between *IAR3* promoter:*mut-IAR3* and 35S:*GFP*, based on *t* test ($P < 0.05$).

(C) Transcript levels of target mimicry against miR167 (*MIM167*, top), miR167a (RNA gel blot), *IAR3* (middle panel), and lateral root numbers (bottom) were assessed in two independent *MIM167* transgenic lines. Vector control or 2-week-old seedlings were grown in control media for 2 weeks and transferred to media containing 0 or 50 μM β -estradiol. Seedlings were harvested after 5 d for RNA gel blot and quantitative PCR analyses. Bars indicate SE ($n = 3$). Lateral root numbers were counted under a stereomicroscope at 2, 4, and 6 d after seedlings were transferred to media containing 0 or 50 μM β -estradiol. Bars indicate SE ($n \geq 12$). Asterisks indicate a significant difference between *MIM167* and vector control transgenic lines based on *t* test ($P < 0.05$).

physiological consequence of the accumulated *IAR3* mRNA levels, we examined lateral root formation in these lines under the same conditions as in Figure 3A. Plants expressing *mut-IAR3* developed more lateral roots than wild-type *IAR3* transgenic plants (Figure 3B). Under this condition, we did not observe consistent and significant difference between the wild type and *iar3-5* (Figure 2D). Similar results were obtained when the native *IAR3* promoter was replaced by a 35S promoter (see Supplemental Figure 4 online). Together, these results indicate that the higher *IAR3* mRNA level in *mut-IAR3* transgenic plants compared with wild-type *IAR3* transgenic lines was due to increased stabilization through mutations at the miR167a complementary site.

To confirm further that *IAR3* mRNA accumulation is directly regulated by miR167, we introduced an inducible target mimicry construct against miR167a (*MIM167a*), so that we could sequester and downregulate miR167a activity in an inducer-dependent manner (Zuo et al., 2000; Franco-Zorrilla et al., 2007). Consistent with our expectations, two independent lines with high levels of *MIM167a* expression downregulated miR167 levels in an inducer-dependent manner. In the presence of inducer, these lines showed significantly increased accumulation of endogenous *IAR3* mRNA compared with untreated controls (Figure 3C, top and middle panels). We also observed that these lines exhibited increased numbers of lateral roots in an inducer-dependent manner (Figure 3C, bottom panel). Thus *MIM167a* transgenic plants phenocopied *IAR3:mut-IAR3* transgenic lines, which is in good agreement with our hypothesis. As a control, we monitored *ILL5* mRNA levels and found no clear mRNA upregulation in the presence of inducer (see Supplemental Figure 5A online, left panel). This is consistent with our data that *ILL5* mRNA levels did not change under stress (Figure 1F).

Using these lines, we also examined *AUXIN RESPONSE FACTOR6* (*ARF6*) and *ARF8* mRNA levels as these mRNAs harbor sequence complementarity to miR167 and have been shown to be targets of miR167 (Wu et al., 2006). We found that *ARF6* and *ARF8* mRNA levels were also elevated in an inducer-dependent manner (see Supplemental Figure 5A online, middle and right panels). This result prompted us to examine *ARF6/8* involvement in osmotic stress. First, we confirmed by an independent analysis that *ARF6/8* mRNA expression patterns did not change under osmotic stress (see Supplemental Figure 5B online). Second, we examined single and double mutant phenotypes under stress. Single mutants *arf6-2* and *arf8-3* showed no clear difference compared with the wild type in control and stress conditions with respect to lateral root development and primary root growth (see Supplemental Figures 5C and 5D online). However, to rule out functional redundancies, we also examined phenotypes of the *arf6-2arf8-3* double mutant. The double mutant showed only a marginal phenotype under osmotic stress conditions in terms of primary root growth and lateral root development compared with the *arf6-2* single mutant, which segregated from the same population as the *arf6-2arf8-3* double mutant is infertile (see Supplemental Figures 5E and 5F online). Mutant plants of *arf8-3* developed more lateral roots than wild-type and *arf6-2* plants under stress at day 6. This was probably due to a faster growth rate of *arf8-3*, as this mutant also produced more lateral roots at day 3 under control

conditions. These results suggest that although *ARF6/8* mRNA levels responded to reduction of miR167 levels in artificial systems like the *MIM167* lines, under osmotic stress, miR167 was preferentially targeted to *IAR3* over *ARF6/8* by an unknown mechanism.

The miR167-*IAR3* Relationship Is Evolutionarily Conserved in Vascular Plants

Since *IAR3* is a positive regulator of high osmotic stress responses, the miR167-*IAR3* interaction might have an important role in this process. Thus, we hypothesized that this interaction had evolved under selective pressure to conserve a responsive mechanism in high osmotic conditions, such as long drought.

We verified that the sequence of miR167a has been almost completely conserved among vascular plant species (see Supplemental Table 5 online). In all nine analyzed species, the only difference in miR167a was either an additional 22nd nucleotide or a substitution at the last nucleotide (miRBase).

The miR167 target site on *IAR3* mRNA is also conserved at both the amino acid and nucleotide levels (Campanella et al., 2003). Since the miR167 complementary site is located in an exon, this conservation is most likely due to the selective pressure on the amino acid sequence and not its relationship with miR167a/b. To observe sequence conservation at the miR167 complementary site in a manner that is affected by miR167 targeting, we took advantage of synonymous substitutions, which often occurred in the 3rd nucleotide of triplet codons. Figure 4 shows *IAR3* sequences from different vascular plant species. Although there is no clear conservation of the 3rd nucleotide of amino acids located in 3' and 5' ends of the target site, those at the center of the complementary site, which is thought to be crucial for the cleavage, are almost completely conserved. We quantified the conservation rate by calculating actual and theoretical nucleotide conservation ratios, which confirmed that the actual conservation ratios were consistently higher than the theoretical values at the center (Figure 4, bottom). In fact, the 3rd nucleotide of Leu was most conserved and this position was exactly where the most frequent cleavage occurred in our 5' RACE analyses (Figure 1G). The first nucleotide of Leu did not seem to be conserved since there were synonymous substitutions. However, the corresponding nucleotide of the miR167a/b was G, which meant that both C and T were complementary nucleotides. Thus, this nucleotide was almost completely conserved as well.

Given that *IAR3* but not *ILL5* responded to miR167, we predicted that there was no similar sequence conservation of the miR167 target site between *IAR3* and other members of the *IAR3* family. Indeed, there was no significant sequence conservation at the synonymous substitution sites (see Supplemental Table 6 online). Taken together, the miR167a-*IAR3* relationship appears to be evolutionarily conserved.

IAR3 Is a Positive Regulator of Drought Stress Tolerance

That *IAR3*-dependent increase in lateral root development occurred during high osmotic stress and that *IAR3* was an evolutionarily conserved target of miR167a suggested a possible role

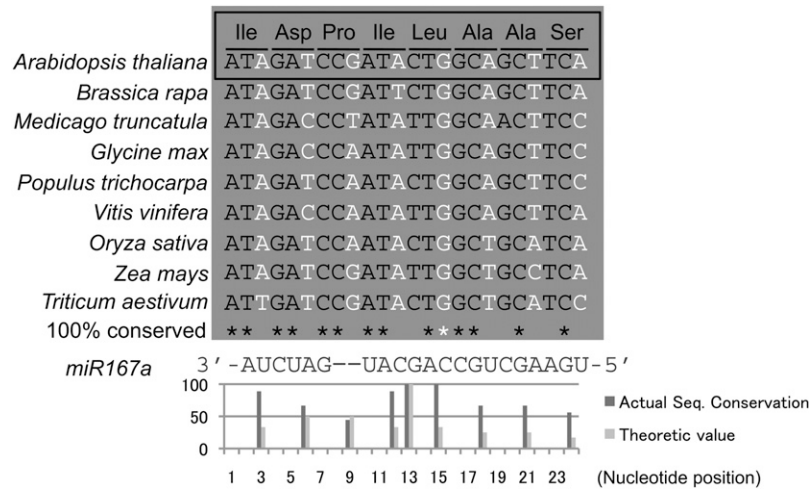


Figure 4. *IAR3* Sequence of the miR167 Target Site Is Evolutionarily Conserved.

IAR3 sequences of the miR167 target site from *Arabidopsis*, *Brassica rapa*, *M. truncatula*, soybean, *Populus trichocarpa*, grape (*Vitis vinifera*), rice, maize, and wheat (*Triticum aestivum*) are shown. The amino acid sequence is shown on top. Third nucleotide of triplet codon is indicated in white; 100% conserved nucleotide in first or second nucleotide of triplet codon is indicated by a black asterisk; that of a 3rd nucleotide is indicated by a white asterisk. miR167a/b sequence is shown at the bottom. The graph shows the percentage of nucleotide sequence conservation for the third nucleotide of each amino acid, calculated from the species shown in this figure.

of *IAR3* in drought stress tolerance. To explore this possibility, we analyzed the drought stress tolerance of *iar3* mutants and the wild type by measuring their capacity to survive for 2 weeks after withholding water. We found that the *iar3* mutants were significantly more sensitive to drought stress than the wild type (Figure 5). Thus, the function of *IAR3* was indispensable in drought tolerance.

DISCUSSION

By a combination of deep sequencing of small RNAs and microarray profiling of mRNAs, we showed that plants responded to high osmotic stress by downregulating miR167 expression, which in turn, upregulated *IAR3* mRNA abundance (Figure 6). We identified a new and functional miRNA-mRNA pair by showing (1) an inverse correlation in expression pattern between the two partners; (2) *IAR3* mRNA cleavage at the miR167 complementary site; (3) transgenic plants expressing a miR167-resistant form of *IAR3* without changing its encoded amino acid sequence exhibited a constitutive stress response, and these plants showed high levels of *IAR3* mRNA accumulation and increased lateral root development compared with plants expressing the wild type form of *IAR3*; and (4) miR167a downregulation by target mimicry technique resulted in the upregulation of *IAR3* transcripts as well as lateral root development.

We also showed that *IAR3*, which releases bioactive IAA from an inactive precursor, is a previously unknown positive factor in root architecture changes in this process (Figure 6). This is consistent with the rich genetic and experimental evidence that auxin is the morphogenic trigger for lateral root formation (Benková et al., 2009) and that ABA and auxin interact in seedling stage (Belin et al., 2009). Consistent with its function in

root architecture changes under high osmotic stress, *IAR3* is also important for drought tolerance. Our observation is also consistent with the result of Liu et al. (1992) using excised segments of *Vigna* hypocotyl that showed that IAA applied simultaneously with osmotic stress enhanced the adaptive recovery of elongation growth by enhancing wall extensibility.

Taken together, our results suggest that the miR167-*IAR3* interaction plays a role in root architecture changes under high osmotic stress as well as in drought stress tolerance. In agreement with this notion, the miR167-*IAR3* interaction appears to be evolutionarily conserved.

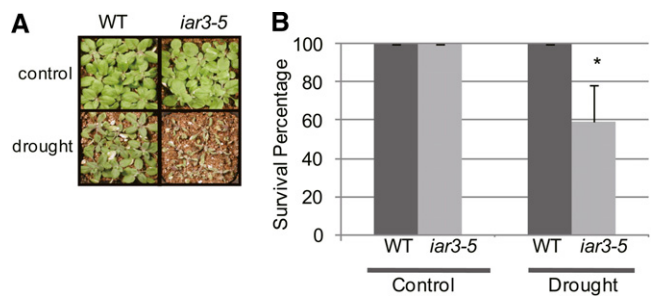


Figure 5. *IAR3* Is a Positive Regulator of Drought Tolerance.

(A) *iar3-5* and wild-type (WT; Col) plants were grown for 2 weeks after soil transfer, with half of the population subjected to drought stress by withholding the water supply and the other half watered normally. Pictures were taken 2.5 weeks after withholding water.

(B) Col and *iar3-5* plants that survived the drought stress were scored after resuming the water supply. Bars show SE ($n = 27$). Asterisks indicate a significant difference between the wild type and *iar3-5*, based on *t* test ($P < 0.05$).

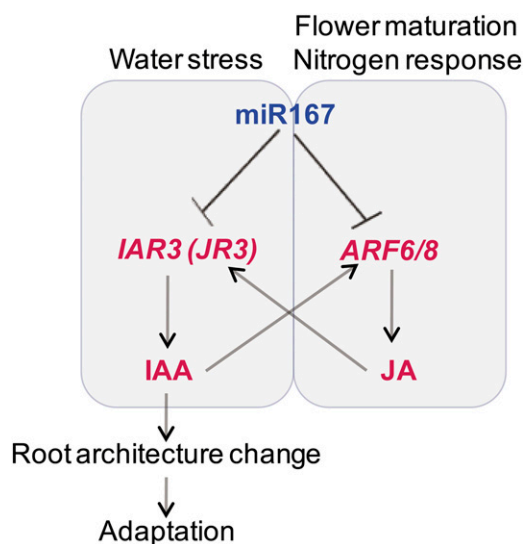


Figure 6. A Working Model of an Integrated View of the First and Secondary Circuits of miR167 Pathways.

Blue letters represent lower expression levels, and red letters represent increased mRNA levels. Dotted arrows show possible secondary effects, in which jasmonic acid produced from the ARF6/8 pathway stimulates accumulation of *IAR3* mRNA, which is also called *JR3*. Likewise, auxin released from *IAR3* stimulates *ARF6/8*.

Localization of the miR167-*IAR3* Circuit

In addition to *IAR3* mRNA, *ARF6/8* mRNAs are also targets of miR167 (Wu et al., 2006). Gifford et al. (2008) showed that the miR167-*ARF6/8* circuit regulates nitrogen-responsive root development by fine-tuning the ratio of initiating/emerging lateral roots. In support of this role, both pre-miR167a and *ARF8* were expressed in the pericycle, where lateral roots emerge (Dubrovsky et al., 2000; Hardtke, 2006; De Smet et al., 2006; Parizot et al., 2008; Petricka and Benfey, 2008). Since these authors performed global cellular expression profiling (Gifford et al., 2008), we searched their database to see if *IAR3* was also expressed in the pericycle. Indeed, we found that among different root tissues, the highest *IAR3* expression level was detected in the pericycle (see Supplemental Figure 6 online). This is consistent with our finding that miR167 and *IAR3* interact directly and that *IAR3* enhances lateral root development (Figure 6).

However, miR167 does not seem to target *ARF6/8* under high osmotic stress (see Supplemental Figures 5B to 5E online). In the future, it would be interesting to investigate how plants use a common regulator (miR167) to perceive different inputs (osmotic stress or nitrogen application) but have distinct outputs as a result.

Although miR167-*ARF6/8* and miR167-*IAR3* circuits do not interact directly, we note that *ARF6/8* promote jasmonic acid production (Nagpal et al., 2005), and jasmonic acid is a strong inducer of *IAR3*, which is identical to *JASMONATE RESPONSIVE3* (*JR3*). On the other hand, bioactive auxin that is released by *IAR3* could easily stimulate the expression of *ARF6/8*, which are auxin-responsive factors. Thus, although these two pathways are not linked in the initial response, they may interact in secondary responses (Figure 6).

Evolutionary Conservation of miR167-*IAR3*

The miR167-*IAR3* relationship appears to be evolutionarily conserved because sequences of synonymous substitution sites are conserved in evolutionarily distant plant species. This is consistent with the recent findings of Wang et al. (2011) that the functions of miR156 and *SPL* mRNAs encoding transcription factors in juvenile-to-adult transition are conserved between *Arabidopsis*, maize (*Zea mays*), and trees (Wu et al., 2009; Wang et al., 2009). The observed versus actual conservation rate revealed that the 5' end of miR167 is more conserved than the 3' end, in agreement with the result of Lin et al. (2009), who reported that the 5' half of artificial miRNA is crucial for its recognition. Furthermore, the complete conservation of the guanine residue at the cleavage site is also consistent with the result of artificial miRNA-targeted virus sequence evolution studies (Lafforgue et al., 2011). Analyses of virus sequence mutations in the lineages that broke artificial miRNA suppression showed that the nucleotide sequence at the cleavage site was most frequently mutated, suggesting also that this cleavage site is crucial in mRNA cleavage by miRNA.

What makes the miR167-*IAR3* relationship advantageous during the course of evolution is not known for certain. Perhaps, developmental flexibility—short and highly branched root architecture—in water-limiting conditions contributes to plant survival in long droughts.

While most of the angiosperm species evolved one to four orthologs of *IAR3*, legume species such as *Medicago truncatula* and soybean (*Glycine max*) evolved seven orthologs (www.phytozome.net). Perhaps these *IAR3* orthologs contribute to better fitness through optimizing root architecture for maximum nitrogen assimilation. Another point is that *M. truncatula* is much more drought tolerant than other leguminous species (Gonzalez et al., 1995; Gálvez et al., 2005). It would be interesting to explore the functional divergence of *M. truncatula IAR3s* in terms of legume-rhizobium symbiosis and drought tolerance.

PIN3 Possibly Functions Downstream of *IAR3*

Our microarray analyses showed that auxin-related genes, such as *PIN-FORMED3* (*PIN3*), *INDOLE-3-ACETIC ACID INDUCIBLE30* (*IAA30*), *MYB DOMAIN PROTEIN14*, and *HOMEBOX PROTEIN40*, were upregulated in high osmotic conditions (see Supplemental Figure 7 online). These genes could be responding to auxin released by *IAR3*. *PIN3*, which controls the direction and rate of cellular auxin efflux, is of particular interest as it is a key factor in the establishment of auxin gradient in adaptation responses such as gravitropism, phototropism, and shade avoidance (Friml et al., 2002; Ding and Friml, 2010; Keuskamp et al., 2010).

In roots, *PIN3* has been reported to mediate the early phase of root gravity response and this is inhibited by cold stress (Friml et al., 2002; Shibasaki et al., 2009). This auxin transporter is localized in pericycle and columella cells (Friml et al., 2002; Bllilou et al., 2005), and the former cell type is where miR167 and *IAR3* are expressed (see above). *PIN3* proteins in columella cells contribute to the reestablishment of the auxin gradient in gravitropism (Friml et al., 2002), and in pericycle cells, *PIN3* proteins are involved in the lateral root development (Benková et al.,

2003). Under normal growth conditions, *pin3* mutants produce fewer lateral root initials, and the double mutant *pin1 pin3* shows defects in lateral root primordium development (Benková et al., 2003). Expressed in the inner cells of lateral root primordia, PIN1 subcellular localization changes polarity from transverse to lateral membranes toward the primordium tip. This correlates with the establishment of an auxin gradient with its maxima at primordium tip (Benková et al., 2003). Interestingly, PIN3 is expressed at the base of lateral root primordia (Benková et al., 2003).

It is possible that, under high osmotic stress, free auxin released by IAR3 stimulates lateral root development through PIN3 where the latter contributes to auxin gradient establishment in the lateral root primordium by an unknown mechanism. Future experiments should test if the PIN3 subcellular localization changes under high osmotic conditions, particularly in relation to lateral root primordia.

METHODS

Plant Materials and Growth Conditions

Arabidopsis thaliana ecotype Columbia-0 (Col-0) was used as the wild type. *iar3-5* (SALK_042101) and *iar3-6* (SALK_090805) were obtained from the ABRC (Alonso et al., 2003). For hydroponic culture, wild-type seeds after a 3-d cold treatment at 4°C were sowed on rock wool (Grodan) presoaked in media containing 1.75 mM sodium phosphate buffer (pH5.8), 1.5 mM MgSO₄, 2.0 mM Ca(NO₃)₂, 3.0 mM KNO₃, 67 μM Na₂EDTA, 8.6 μM FeSO₄, 10.3 μM MnSO₄, 30 μM H₃BO₃, 1.0 μM ZnSO₄, 24 nM (NH₄)₆Mo₇O₂₄, 130 nM CoCl₂, and 1 μM CuSO₄ (Fujiwara et al., 1992; Taniguchi et al., 1998). In short, plants were grown at 22°C with a 150 μmol m⁻² s⁻¹ fluorescence rate and under 12-h/12-h light/dark cycle in MGRL hydroponic medium. At this time, three separate populations of seedlings were defined. Two populations of 30-d-old plants were then subjected to high osmotic stress by adding 0.3 M mannitol to the basal MGRL medium for 3 h in the light (designated: Osmo treatment). The remaining population was treated without 0.3 M mannitol (designated: Mock). Recovery plants were returned to the MGRL medium for 1 d after the stress treatment and sampled (designated: Recovery). This procedure was used to treat three independent biological replicates. Leaf and root samples were collected separately from the same set of plants. There were a total of 18 samples (two tissues × three samples × three replicates).

RNA Isolation and sRNA Library Construction

For small RNA library construction, three biological replicates were pooled and total RNA was extracted using Trizol reagent (Invitrogen) according to the manufacturer's protocol. Small RNAs (18 to 28 nucleotides) from 40 μg total RNA were separated and then fractionated using 15% polyacrylamide urea gel. Subsequent steps were performed by the protocol of Digital Gene Expression for Small RNA Sample Preparation (Illumina). Qualities and quantities of individual sRNA libraries were confirmed using a bioanalyzer. Solexa sequencing was performed at the Genomics Resource Center in the Rockefeller University following the manufacturer's instructions (Illumina).

Small RNA Gel Blot Hybridization and 5' RACE

RNA gel blot and 5' RACE were performed essentially according to Reyes and Chua (2007). Briefly, total RNA was separated on 15% polyacrylamide urea gels and then transferred onto Amersham Hybond-N⁺ (GE Healthcare). The membrane was prehybridized with Ultrahyb-oligo

buffer (Ambion) at 42°C for at least 1 h after UV cross-linking. For probe labeling, 15 pmol oligo DNA complementary to a mature miRNA sequence was radioactively labeled by T7 polynucleotide kinase reaction (NEB). [³²γ] ATP (40 μCi) was used for the phosphorylation reaction at 37°C for 1 h. Radioactive ATP unincorporated was removed by a mini quick spin oligo columns (Roche), and the purified probe was used for hybridization at 42°C overnight. The membrane was washed with 2× SSC and 0.1% SDS. The First Choice 5' RLM-RACE Kit (Ambion) was used for 5' RACE according to the manufacturer's instructions.

Phenotypic Assays

For root growth assays, plants were grown for 1 week (Murashige and Skoog [MS], 0 or 0.5% Suc, and 0.8% agar) and transferred to vertical plates containing MS medium (half-strength MS, 0.5% Suc, and 0.6% agar) supplemented with 0.25 M mannitol for high osmotic stress treatments. Growth rates of primary roots were measured using ImageJ software (National Institutes of Health). Drought assays were done according to Catala et al. (2007). All experiments were performed with at least three biological replicates.

Small RNA Sequencing Data

Adapter sequence was clipped by the Perl program. All trimmed reads were mapped to the *Arabidopsis* genome (TAIR9) using local C program. Only reads with a perfect hit(s) were selected for further analysis.

Prediction and Identification of *Arabidopsis* miRNAs and Their mRNA Targets

The method of Wang et al. (2004) was used.

GeneChip Arrays

In short, targets for *Arabidopsis* GeneChip ATH1 were synthesized from 1 μg total RNA extracted by RNA easy extraction kit (Qiagen). In total, there were 18 individual GeneChip targets comprising three biological replicates for CL (Col leaf), CR (Col root), OL (osmotic-stressed leaf), OR (osmotic-stressed root), RL (recovery leaf), and RR (recovery root). Target synthesis, hybridization, and scanning of all arrays were performed at the Genomics Resource Center, Rockefeller University, following the manufacturer's instructions (Affymetrix). Background correction was performed as described by Irizarry et al. (2003). Quantile normalization was performed as described by Bolstad et al. (2003). Finally, targets were summarized into single-gene expression (probe set) values on a per chip basis using the median polish algorithm (Tukey, 1977). Differential gene expression was determined for each of the following comparisons: OR/CR, OR/RR, OL/CL, and OL/RL. Significance was determined at P values below 0.05, adjusted for multiplicity using the Bonferroni correction method (22,811 probe sets). All comparisons were made using the *ebayes* function implemented under the *limma* package in the R statistical environment (Clayton and Kaldor, 1987; Smyth, 2005). Adjusted P values for all the genes represented in the ATH1 array can be found in Supplemental Data Set 1 online.

Plasmid Construction, Plant Transformation, and RNA Isolation

Plant transformation, plasmid construction, and RNA gel blot analyses were done as described by Kinoshita et al. (2010). Promoter fusions were constructed by QuikChange Lightning Site-Directed Mutagenesis (Stratagene) and In-Fusion Cloning Kit (Clontech) according to the manufacturers' instructions using oligonucleotides in Supplemental Table 7 online. pBA002a was used as a vector without promoter with *EcoRV* and *MluI* as restriction enzymes (Møller et al., 2001). Gateway (Invitrogen)-compatible inducible

vector and pBA002 vector were used for inducible *MIM167* cloning and 35S:*IAR3* cloning, respectively (Zuo et al., 2000).

Endogenous Auxin Measurement

Free IAA was measured in three biological replicates according to Sugawara et al. (2009).

Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL databases under the following accession numbers: *IAR3* (AT1G51760), *ILL5* (AT1g51780), *ARF6* (AT1G30330), *ARF8* (AT5G37020), miR167a (AT3G22886) and *PIN3* (AT1G70940). Raw data are available from the Gene Expression Omnibus under accession numbers GSE36560 (small RNA) and GSE36789 (microarray).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Size Distribution of Small RNAs from Leaves and Roots in Control and High Osmotic Stress.

Supplemental Figure 2. Relative Expression Levels of High Osmotic Stress-Responsive Genes.

Supplemental Figure 3. *IAR3* mRNA Levels in *iar3-5* and *iar3-6*.

Supplemental Figure 4. Relationships between miR167, *IAR3*, and Lateral Root Numbers in Constitutive Promoter System.

Supplemental Figure 5. Involvement of *ILL5*, *ARF6*, and *ARF8* in the MIM167 System.

Supplemental Figure 6. Tissue-Specific Expression of *IAR3* mRNA in Root Tissue.

Supplemental Figure 7. Relative Expression Patterns of Auxin-Responsive Genes.

Supplemental Table 1. Summary of Deep Sequencing Results.

Supplemental Table 2. List of miR167 Putative Targets.

Supplemental Table 3. miR167-Like Small RNAs That Might Target *IAR3* mRNA.

Supplemental Table 4. Normalized Expression Levels of miR167 and miR167-Like RNAs in Stress.

Supplemental Table 5. Sequence Conservation of miR167 among Vascular Plants.

Supplemental Table 6. miR167a Target Site Conservation among *IAR3* Family Members.

Supplemental Table 7. List of Oligonucleotide Sequences.

Supplemental Data Set 1. Significance Tests for All Comparisons.

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AUTHOR CONTRIBUTIONS

N.K., H.W., and N.-H.C. designed research. N.K. performed most of the experiments. H.K. and Y.K. measured endogenous auxin. H.W., C.M., and J.L. carried out bioinformatics analyses. Results were discussed with all authors. The article was written by N.K. and N.-H.C. and read by all the authors.

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