

Cloning and Characterization of *IAR1*, a Gene Required for Auxin Conjugate Sensitivity in Arabidopsis

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Most indole-3-acetic acid (IAA) in higher plants is conjugated to amino acids, sugars, or peptides, and these conjugates are implicated in regulating the concentration of the free hormone. We identified *iar1* as an Arabidopsis mutant that is resistant to the inhibitory effects of several IAA–amino acid conjugates but remains sensitive to free IAA. *iar1* partially suppresses phenotypes of a mutant that overproduces IAA, suggesting that *IAR1* participates in auxin metabolism or response. We used positional information to clone *IAR1*, which encodes a novel protein with seven predicted transmembrane domains and several His-rich regions. *IAR1* has homologs in other multicellular organisms, including *Drosophila*, nematodes, and mammals; in addition, the mouse homolog KE4 can functionally substitute for *IAR1* in vivo. *IAR1* also structurally resembles and has detectable sequence similarity to a family of metal transporters. We discuss several possible roles for *IAR1* in auxin homeostasis.

INTRODUCTION

Indole-3-acetic acid (IAA) is the major endogenous auxin and participates in many plant developmental processes, including cell enlargement and division, differentiation of vascular tissue, initiation of lateral roots, apical dominance, and responses to environmental stimuli such as gravity and light (reviewed by Estelle and Klee, 1994; Bennett et al., 1998). Plants contain little free IAA; most IAA is found conjugated to amino acids, peptides, sugars, or high molecular weight glycans. These conjugates have been implicated in such processes as storage, transport, and protection from oxidative degradation (reviewed in Cohen and Bandurski, 1982; Bandurski et al., 1995). Plants apparently also permanently inactivate excess IAA by conjugation (reviewed in Normanly, 1997). For example, many plants form IAA-Asp as an intermediate in IAA catabolism (Tsurumi and Wada, 1986; Monteiro et al., 1988; Tuominen et al., 1994; Östin et al., 1998).

Conjugation and hydrolysis of conjugates are probable mechanisms used to regulate the concentrations of free IAA (reviewed in Normanly and Bartel, 1999), and characterization of the genes involved in these processes is a prerequisite to understanding this regulation. Although diverse land plants conjugate IAA to glucose and other molecules (Sztejn

et al., 1995, 1999), the only plant gene so involved that has been identified is the maize *iaglu* gene, which encodes an enzyme esterifying IAA to glucose (Szczeszen et al., 1994). Various plants hydrolyze IAA conjugates, and IAA-glucose hydrolases have been identified in maize, potato, oat, and bean (Kowalczyk and Bandurski, 1990; Jakubowska et al., 1993). IAA-Ala hydrolases have been partially purified from bean and carrot (Cohen et al., 1988; Kuleck and Cohen, 1993), and Chinese cabbage extracts contain isoenzymes that hydrolyze IAA-Ala, IAA-Asp, and IAA-Phe (Ludwig-Müller et al., 1996).

The ability of several IAA conjugates to mimic the effects of free IAA on plant growth (reviewed by Bartel, 1997) has been exploited to identify Arabidopsis mutants that respond abnormally to IAA conjugates. The *ilr1* (IAA-Leu-resistant) mutant elongates roots at concentrations of IAA-Leu that inhibit wild-type root growth. The gene defective in *ilr1* encodes an amidohydrolase with high affinity for IAA-Leu and IAA-Phe and lesser affinity for IAA-Ala, IAA-Gly, and IAA-Val (Bartel and Fink, 1995). Four *ILR1*-like genes, *ILL1*, *ILL2* (Bartel and Fink, 1995), *ILL3*, and *ILL5* (Davies et al., 1999), have been identified on the basis of their similarity to *ILR1*. A sixth member of this gene family is defective in the *iar3* (IAA-Ala-resistant) mutant and encodes an amidohydrolase specific for IAA-Ala (Davies et al., 1999). Two other mutants, *icr1* and *icr2*, are resistant to growth inhibition by IAA-Phe, IAA-Ala, and IAA-Gly (Campanella et al., 1996), but the genes defective in these mutants remain unidentified.

We have been identifying and characterizing Arabidopsis mutants with altered responses to IAA–amino acid conjugates to elucidate the role of conjugate metabolism in plant

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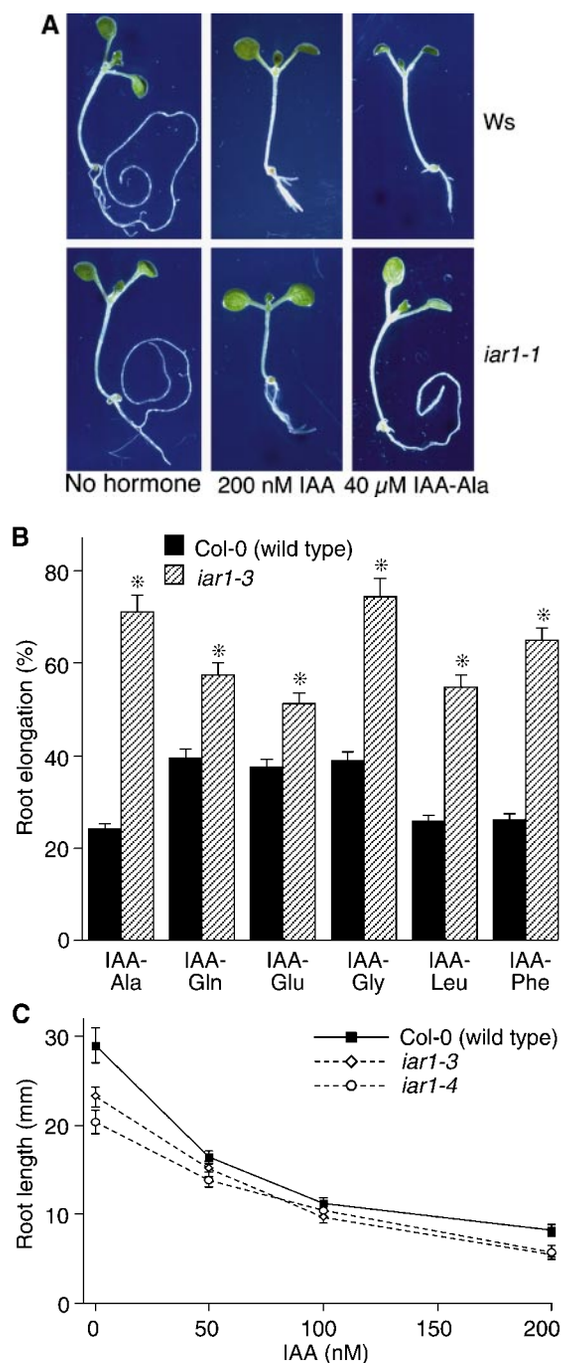


Figure 1. *iar1* Roots Have Decreased Sensitivity to IAA–Amino Acid Conjugates and Normal Sensitivity to IAA.

(A) Eight-day-old *iar1-1* mutant and Wassilewskija (Ws) wild-type seedlings were removed from the agar and photographed after growth on medium containing no hormone, 200 nM IAA, or 40 μ M IAA-Ala.

(B) Wild-type Columbia (Col-0) and *iar1* mutant seed were plated on medium containing 20 μ M of the indicated IAA–amino acid conjugates. After 8 days, the primary root of each seedling was measured

development. Here, we report the isolation of the *iar1* auxin conjugate-resistant mutant and the positional cloning of *IAR1*, which encodes a polytopic membrane protein with several His-rich regions. *IAR1* represents a novel component of the auxin conjugate metabolic machinery in plants, and several possible models for its function are discussed.

RESULTS

Isolation and Characterization of *iar1* Mutants

Certain IAA–amino acid conjugates mimic free IAA in bioassays (reviewed in Bartel, 1997), and IAA conjugates that inhibit Arabidopsis root elongation have been used in mutant screens to identify genes involved in hydrolysis of IAA conjugates (Bartel and Fink, 1995; Davies et al., 1999). We initially identified the *iar1* mutant because its roots elongate at concentrations of IAA-Ala that are normally inhibitory, as shown in Figure 1. Table 1 lists the seven *iar1* alleles, four of which were isolated in screens for IAA-Ala resistance. Three additional *iar1* alleles were identified as enhancers of the *ilr1* mutant, which is defective in an IAA conjugate hydrolase (Bartel and Fink, 1995). These alleles were isolated based on the increased resistance of *ilr1 iar1* double mutants to IAA-Leu or IAA-Phe. The recessive nature of the *iar1* mutations (data not shown) suggests that they are loss-of-function alleles.

We quantitated the IAA conjugate resistance of the *iar1* alleles by comparing root lengths of seedlings grown on medium containing various conjugates. All *iar1* alleles are resistant to IAA-Ala, *iar1-4* and *iar1-5* being the strongest and weakest alleles, respectively (data not shown). Because *ilr1 iar1* double mutants are highly resistant to IAA-Leu and IAA-Phe (data not shown), we tested the response of *iar1* single mutants to these conjugates in root elongation assays. *iar1-3* is resistant to IAA-Phe at all concentrations tested (10 to 40 μ M) but is resistant to IAA-Leu only at lower concentrations (10 to 20 μ M; Figure 1B and data not shown). *iar1* roots are also resistant to elongation inhibition by IAA-Gly, IAA-Gln, and IAA-Glu (Figure 1B), indicating that *IAR1* is

and normalized against growth on unsupplemented medium. Error bars indicate standard errors of the means ($n = 13$), and asterisks indicate significant differences from the wild type (Student's *t* test, $P < 0.0001$).

(C) Wild-type (Col-0) and *iar1* mutant seed were plated on medium containing the indicated concentrations of IAA. After 8 days, the primary root of each seedling was measured. Error bars indicate standard errors of the means ($n = 10$).

Table 1. *iar1* Mutant Alleles

Allele	Accession	Screen	Mutagen	Codon Change ^a	Amino Acid Change	Allele Detection
<i>iar1-1</i>	Ws	IAA-Ala ^R	EMS	<u>GGA</u> → <u>AGA</u>	Gly ⁴⁰⁰ →Arg	Destroys HpaII site
<i>iar1-2</i>	Ws	IAA-Ala ^R	EMS	<u>GGA</u> → <u>AGA</u>	Gly ⁴²³ →Arg	Destroys AciI site
<i>iar1-3</i>	Col-0	IAA-Ala ^R	Fast neutron	<u>GGAATG</u> → <u>GGGCCTG</u>	Frameshift at codon 458	Creates HaeIII site
<i>iar1-4</i>	Col-0	IAA-Ala ^R	Fast neutron	65 to 80-kb deletion including entire <i>IAR1</i> coding region		Late-flowering due to <i>FKF1</i> deletion ^b
<i>iar1-5</i>	Ws	<i>ilr1</i> enhancer IAA-Leu ^R	EMS	<u>AGA</u> → <u>AAA</u>	Arg ³⁵⁷ →Lys	Sequencing
<i>iar1-6</i>	Ws	<i>ilr1</i> enhancer IAA-Leu ^R	EMS	<u>CAA</u> → <u>TAA</u>	Gln ¹⁶² →stop	Destroys HphI site
<i>iar1-7</i>	Ws	<i>ilr1</i> enhancer IAA-Phe ^R	EMS	<u>tagCT</u> → <u>taaCT</u> (3' splice site of first intron) ^c	Predicted to disrupt splicing	Creates AluI site

^aNucleotide changes are underlined.

^bNelson et al., 2000.

^cIntron bases are in lowercase letters; exon bases are in uppercase letters.

quired for IAA conjugate sensitivity in general rather than being conjugate specific. In contrast, *iar1* plants respond normally to a range of concentrations of free IAA (Figures 1A and 1C), indicating that mutations in *IAR1* alter conjugate sensitivity rather than general auxin responsiveness.

As shown in Figure 2, exogenous IAA or IAA conjugates can inhibit Arabidopsis hypocotyl elongation in the light; accordingly, we examined the effects of IAA conjugates on *iar1* hypocotyl elongation. *iar1* mutant hypocotyls respond normally to free IAA but are resistant to the inhibitory effects of IAA-Ala on elongation (Figure 2). These results suggest that IAR1 functions in the hypocotyls as well as the roots of Arabidopsis seedlings.

The *iar1* Mutation Partially Suppresses *alf1* Phenotypes

The Arabidopsis *aberrant lateral root formation* (*alf1*) mutant (Celenza et al., 1995), also isolated as *rooty* (King et al., 1995), *superroot1* (Boerjan et al., 1995), and *hookless3* (Lehman et al., 1996), overproduces both free and conjugated IAA (Boerjan et al., 1995; King et al., 1995; Lehman et al., 1996). *alf1* plants have short primary roots, display increased numbers of adventitious and lateral roots, fail to make an apical hook when germinated in the dark, and are completely sterile under normal growth conditions (Celenza et al., 1995). These phenotypes can be copied by applying auxin to wild-type plants (Boerjan et al., 1995; Celenza et al., 1995; King et al., 1995; Lehman et al., 1996). To investigate whether IAR1 influences IAA responses in vivo, we generated homozygous *iar1/iar1 alf1/alf1* plants (see Methods). *iar1* partially rescues the fertility defect of *alf1*, in that double mutants yield some seed. In addition, *iar1 alf1* plants are able to form a partial apical hook when germinated in the dark (data not shown). We quantified the degree to which *iar1* suppresses *alf1* by monitoring root elongation on unsupplemented medium. As shown in Figure 3, the *iar1* muta-

tion suppresses the *alf1* root elongation defect. The observation that loss of *IAR1* function can partially alleviate defects in the *alf1* auxin-overproducing mutant suggests that IAR1 normally acts to increase the concentrations of free IAA or the sensitivity to IAA.

Suppression of the *iar1* Phenotype by Manganese

The Arabidopsis amidohydrolases characterized to date require a metal cofactor. Mn²⁺ and Co²⁺ are the most effective

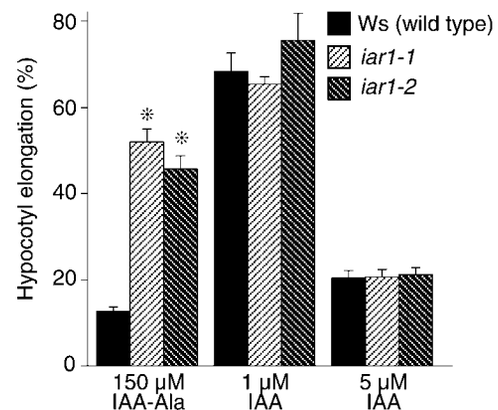


Figure 2. *iar1* Hypocotyls Have Decreased Sensitivity to IAA-Ala.

Wild-type and *iar1* seed were plated on medium containing the indicated concentrations of IAA or IAA-Ala. After 8 days in yellow-filtered light at 22°C, hypocotyls were measured and normalized against growth on unsupplemented medium. Error bars indicate standard errors of the means ($n = 20$ for Ws and *iar1-1*, $n = 9$ for *iar1-2*), and asterisks indicate significant differences from the wild type (Student's *t* test, $P < 0.0001$).

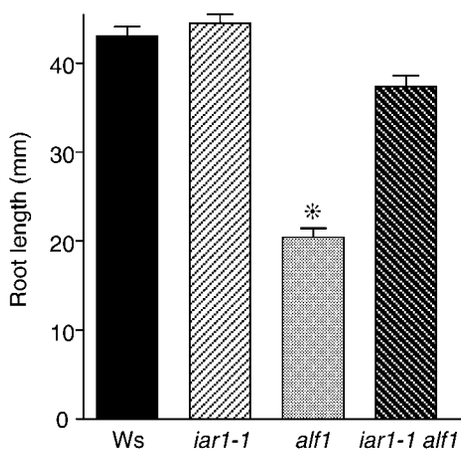


Figure 3. *iar1* Suppresses the *alf1* Root Elongation Defect.

Seed of Ws (wild type), *iar1-1* and *alf1* single mutants, and *iar1-1 alf1* double mutants were plated on unsupplemented medium. After 10 days under yellow-filtered light at 22°C, the primary root of each seedling was measured. Homozygous *alf1* mutants were identified among the progeny of an *alf1/ALF1* parent based on the epinastic cotyledon phenotype. Error bars indicate standard errors of the means ($n \geq 18$), and the asterisk indicates a significant difference from the wild type (Student's *t* test, $P < 0.0001$).

cofactors identified for heterologously expressed Arabidopsis IAA-amino acid conjugate hydrolases purified from *Escherichia coli* (Bartel and Fink, 1995; Davies et al., 1999; R. Tellez and B. Bartel, unpublished data). Because mutations in these amidohydrolases can cause IAA-amino acid conjugate resistance (Bartel and Fink, 1995; Davies et al., 1999), mutations that prevent the hydrolases from assembling with or retaining the proper cofactor also might cause this phenotype. We examined the response of *iar1* to Mn^{2+} alone or to IAA-Ala and Mn^{2+} . As shown in Figure 4A, Mn^{2+} alone did not dramatically inhibit root growth at the concentrations tested. However, high Mn^{2+} concentrations partially suppressed the IAA-Ala resistance of *iar1* (Figure 4B). This finding suggests that IAR1 might be necessary to provide the correct metal cofactor to the IAA-amino acid conjugate hydrolases.

IAR1 Cloning

To elucidate the role of IAR1 in conjugate metabolism, we cloned *IAR1* by using a map-based positional approach. A recombination mapping population was prepared by outcrossing *iar1* plants in the Wassilewskija (Ws) accession to wild-type Columbia (Col-0) plants. DNA isolated from IAA-Ala-resistant F_2 progeny was analyzed with polymerase chain reaction (PCR)-based polymorphic markers (Konieczny and Ausubel, 1993; Bell and Ecker, 1994). The *iar1* mutation

was localized to an unsequenced (at that time) region at the bottom of chromosome 1, between simple sequence length polymorphism markers nga280 and nga111 (Bell and Ecker, 1994). An 800-kb yeast artificial chromosome (YAC) contig had been described in this interval (Vijayraghavan et al., 1995), and we used subclones from these and other nearby YACs to develop additional PCR-based markers for mapping (see Methods). We localized the *iar1* mutation between the markers 9H12LE and 13A11RE, as shown in Figure 5. Using PCR and DNA hybridization analysis, we determined that this interval is completely contained on the 116-kb bacterial artificial chromosome (BAC) T7E4 (Figure 5A).

We constructed a complementation library from BAC T7E4 (see Methods) and used *Agrobacterium* to transform (Clough and Bent, 1998) these subclones into the *iar1* mutant. T_2 progeny of *iar1-3* plants transformed with the various constructs were tested for conjugate resistance. Approximately 75% of the T_2 progeny of C37 (Figure 5B) transformants had wild-type IAA-Ala sensitivity, as expected for a rescuing construct. We sequenced the 12-kb insert of C37 and identified several potential coding regions. One of these genes was altered in all seven *iar1* alleles. Three alleles (*iar1-1*, *iar1-2*, and *iar1-5*) had mutations that caused amino acid substitutions; *iar1-6* had a nonsense mutation; the mutation in *iar1-7* probably disrupted splicing because it changed the final G of the first intron to an A; *iar1-3* had a frameshift mutation predicted to replace the last 11 amino acids of IAR1 with an alternative 14-amino acid stretch (Table 1; Figure 5C); and *iar1-4* had an ~65- to 80-kb deletion that included the entire C37 region (Nelson et al., 2000). Finally, the 6.5-kb genomic clone pBIN/IAR1g (see Methods), which contains the *IAR1* open reading frame with 2.1 kb of DNA upstream of the initiator ATG, completely restored IAA-Ala sensitivity to the *iar1* mutant (Figure 5D), confirming that we had identified *IAR1*.

IAR1 Encodes a Membrane Protein with Multiple His-Rich Regions

We isolated a full-length *IAR1* cDNA (GenBank accession number AF216524) by hybridization to a cDNA library (see Methods). Comparing this cDNA with various genomic sequences revealed that the *IAR1* coding region has 10 introns (Figure 5C) and potentially encodes a 469-amino acid protein with seven or eight transmembrane domains, as shown in Figures 6A and 7. IAR1 is 28% identical to the *Drosophila* Catsup (catecholamines up) protein (GenBank accession number AAF37226) and 26% identical to the predicted mouse KE4 and human HKE4 proteins (Abe et al., 1988; St.-Jaques et al., 1990; Janatipour et al., 1992; Ando et al., 1996). Uncharacterized IAR1 homologs are found in other multicellular organisms, including *Drosophila* (GenBank accession numbers AAF49687, AAF50401, and AAF56960), *Caenorhabditis elegans* (GenBank accession numbers CAB17070, CAB05297.1, and CAB02806.1), and zebrafish (GenBank ac-

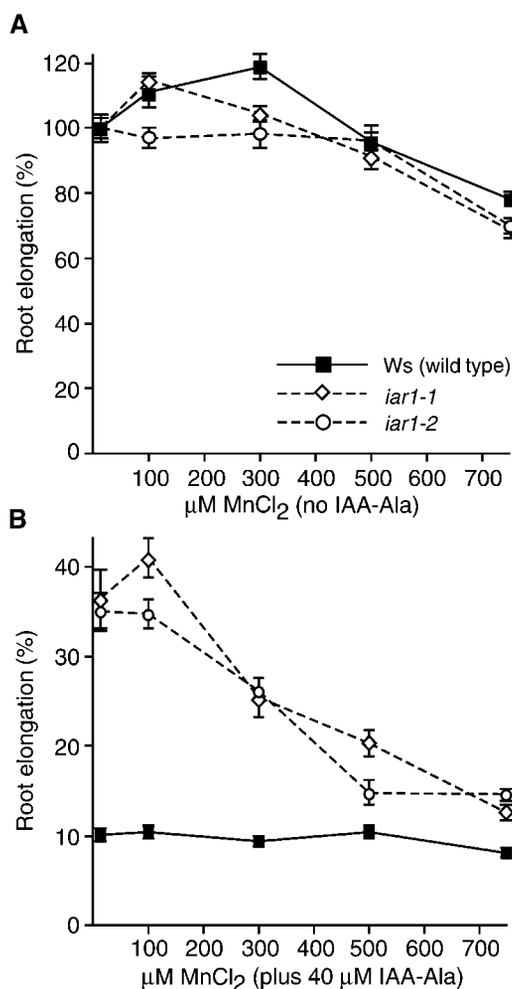


Figure 4. Manganese Suppresses *iar1* IAA-Ala Resistance.

Wild-type (Ws) and *iar1* mutant seed were plated on medium containing 14 to 750 μM manganese either with or without 40 μM IAA-Ala. After 8 days under yellow-filtered light at 22°C, the primary root of each seedling was measured and normalized to growth on unsupplemented medium (which contained 14 μM manganese). Error bars indicate standard errors of the means ($n = 12$).

(A) Without IAA-Ala.

(B) With 40 μM IAA-Ala.

cession number AAF05821). Figure 6A shows an alignment of IAR1 with similar proteins from *Drosophila* and mammals.

The closest IAR1 homologs of known function are members of the ZIP (ZRT, IRT-like protein) family of metal transporters (Eng et al., 1998; Guerinot and Eide, 1999; Guerinot, 2000), which are ~ 10 to 13% identical to IAR1 (Figure 6B). These widespread transporters have been characterized from yeast (Zhao and Eide, 1996a, 1996b; MacDiarmid et al., 2000), *Arabidopsis* (Grotz et al., 1998), and mammals (Lioumi et al., 1999; Gaither and Eide, 2000).

The Mouse *KE4* Gene Can Functionally Substitute for *IAR1* in *Arabidopsis*

The *IAR1* homologs *KE4* and *HKE4* are mouse and human genes of unknown function (Abe et al., 1988; St.-Jaques et al., 1990; Janatipour et al., 1992; Ando et al., 1996). In addition to 26% amino acid identity (Figure 6A), *IAR1* and *KE4* share very similar hydropathy plots, suggesting comparable membrane-spanning topologies (Figure 7). To determine whether the *Arabidopsis* *IAR1* and mammalian *KE4* proteins function analogously, we transformed *iar1-3* mutant plants with a construct expressing the *KE4* cDNA from the constitutive cauliflower mosaic virus 35S promoter (35S-*MmKE4*). As shown in Figure 8, the 35S-*MmKE4* construct substantially restores IAA-Ala sensitivity to the *iar1* mutant, indicating that the mouse *KE4* gene can substitute for *IAR1* in vivo. This finding suggests that *IAR1* and *KE4* have similar functions in *Arabidopsis* and mammals.

Several *Arabidopsis* *ZIP* genes have been isolated (Grotz et al., 1998) that restore growth on low-zinc medium to a *S. cerevisiae* *zrt1 zrt2* double mutant that is defective in high- and low-affinity plasma membrane-localized zinc transporters (Zhao and Eide, 1996a, 1996b). We tested whether *IAR1* could rescue this mutant because these transporters are apparently related to *IAR1* (Figure 6B) and are structurally similar (Figure 7). In contrast to the expression of several *Arabidopsis* *ZIP* genes (Grotz et al., 1998), *IAR1* expression does not rescue the *zrt1 zrt2* strain in the presence of limiting zinc (data not shown), suggesting that *IAR1* does not encode a plasma membrane-localized zinc transporter. However, perhaps *IAR1* does not fold or localize properly in yeast, because the human hZIP2 zinc transporter also fails to rescue *zrt1 zrt2* mutants (Gaither and Eide, 2000).

DISCUSSION

Requirement of *IAR1* for IAA-Amino Acid Conjugate Sensitivity

Phenotypic analysis of *iar1* mutants indicates that *IAR1* participates in IAA-amino acid conjugate metabolism or sensing. *iar1* mutant roots and hypocotyls respond normally to free IAA but are less sensitive than wild-type roots and hypocotyls to several IAA-amino acid conjugates (Figures 1 and 2). In contrast, two other IAA-amino acid-resistant mutants, *ilar1* and *iar3*, are defective in IAA conjugate hydrolases that have distinct substrate specificities and display correspondingly restricted conjugate resistance profiles (Bartel and Fink, 1995; Davies et al., 1999). The broad *iar1* conjugate resistance profile suggests a general role for *IAR1* in IAA-amino acid conjugate metabolism rather than an IAA-Ala-specific role.

Mutation of *IAR1* partially alleviates several auxin overproduction phenotypes of the *alf1* mutant, indicating that *IAR1*

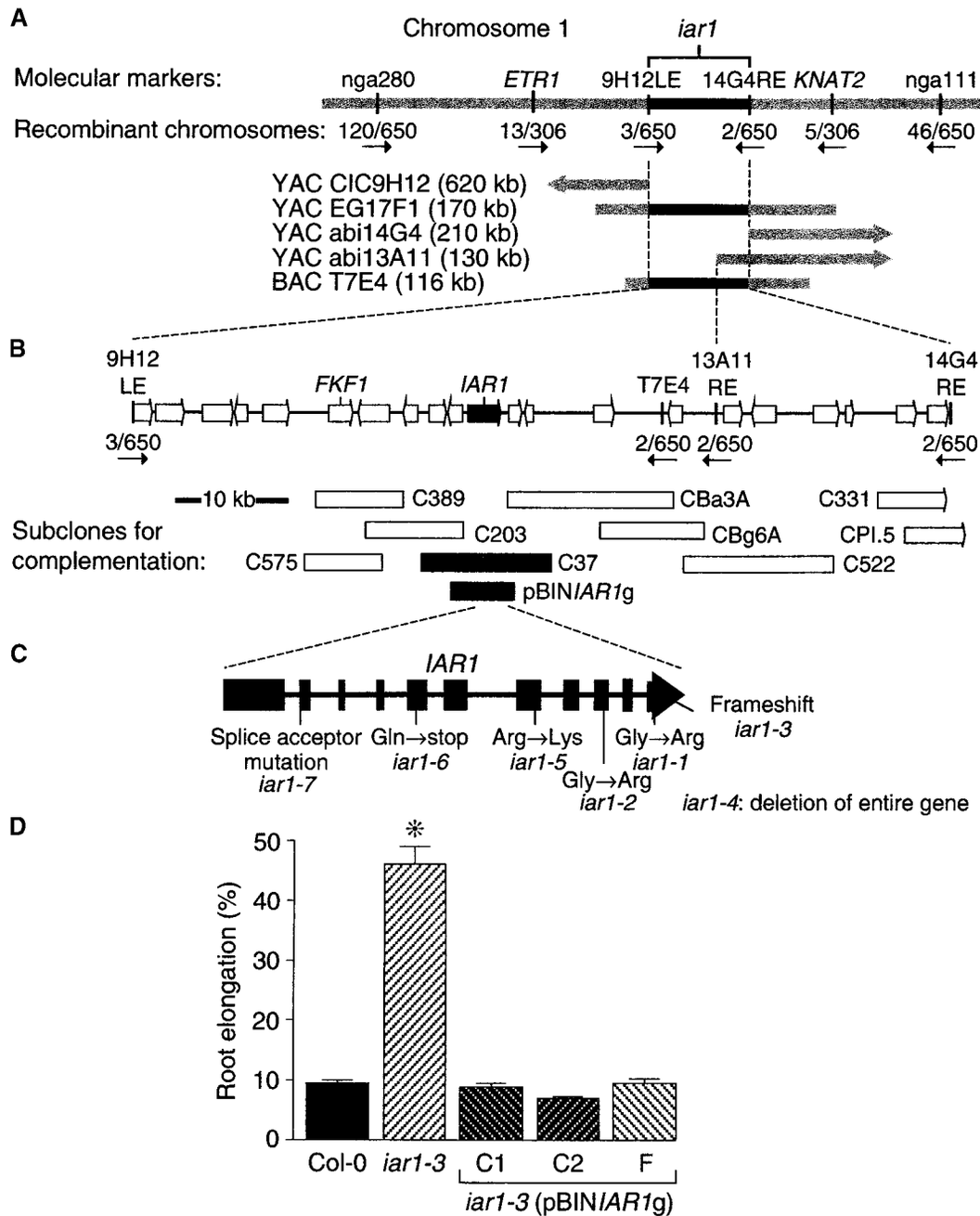


Figure 5. Positional Cloning of *IAR1*.

(A) Recombination mapping with PCR-based markers nga111 and nga280 (Bell and Ecker, 1994) localized *iar1* to the bottom of chromosome 1. This position was refined to between *ETR1* (Chang et al., 1993) and *KNAT2* (Lincoln et al., 1994). Additional markers from nearby YAC clones included 9H12LE and 14G4RE (Nelson et al., 2000) and a marker made from the right end of YAC abi13A11 (see Methods). These three markers hybridized to BAC T7E4 (Choi et al., 1995).

(B) A complementation library was constructed (see Methods) from BAC T7E4, and subclones that were tested for *iar1* complementation are indicated by rectangles below the predicted genes (arrows) in the region. Subclones shown in black rescued the *iar1* IAA-Ala resistance, and those shown in white did not.

(C) Positions of the seven *iar1* mutations are shown below a scheme of the *IAR1* coding region. Exons are indicated by closed rectangles, and introns are represented by lines.

(D) A genomic construct containing the predicted *IAR1* open reading frame controlled by its own promoter rescued the *iar1* mutant phenotype.

normally increases free IAA or sensitivity to IAA. The *alf1* mutant has shortened primary roots, has more adventitious and lateral roots, and is completely sterile (Celenza et al., 1995), probably because of increased concentrations of free IAA (Boerjan et al., 1995; King et al., 1995; Lehman et al., 1996). The *iar1* mutation partially suppresses the *alf1* phenotype, restoring normal root elongation (Figure 3) and partially restoring fertility and the ability to form an apical hook when germinated in the dark (data not shown).

IAR1 Homologs in Other Multicellular Organisms

The *IAR1* gene potentially encodes a 469–amino acid protein that is 26 to 28% identical to the *Drosophila* Catsup protein and to mammalian KE4 proteins encoded in the major histocompatibility complex (MHC) class II locus (Abe et al., 1988; St.-Jaques et al., 1990; Janatipour et al., 1992; Ando et al., 1996). Interestingly, these *IAR1*-like genes (Figure 6A) probably play similar or identical roles in plants and animals, because expression of the mouse *KE4* cDNA restores normal IAA-Ala sensitivity to *iar1* plants (Figure 8). These proteins all have predicted cleavable N-terminal signal sequences, several regions rich in His residues, and six or seven transmembrane domains (Figure 6A). Their almost identical hydropathy plots (Figure 7) suggest common membrane topologies. *IAR1* is most similar to its homologs in and near potential transmembrane domains, and two of the *iar1* point mutations have Arg residues substituted for conserved Gly in predicted transmembrane domains (Figure 6A), an indication that these regions are functionally important.

IAR1 Similarity to the ZIP Family of Metal Transporters

Intriguingly, members of the ZIP family of metal transporters (reviewed by Guerinot and Eide, 1999; Guerinot, 2000) also have N-terminal signal sequences, seven or eight transmembrane domains (Figure 7), and His-rich regions. Although most ZIP family members were isolated as zinc transporters (Zhao and Eide, 1996a, 1996b; Grotz et al., 1998; Gaither and Eide, 2000), results of both direct uptake assays (Eide et al., 1996; Korshunova et al., 1999; Pence et al., 2000) and competition experiments (Grotz et al., 1998; Gaither and Eide, 2000) suggest that ZIPs can have varied

substrate specificities. *IAR1*, *KE4*, and *Catsup* have weak identity to the predicted transmembrane domains of ZIP family members (Figure 6C and data not shown), including most of the consensus residues noted previously (Eng et al., 1998). The conserved His residues in transmembrane domains IV and V of the ZIPs that are essential for metal transport in *Arabidopsis* *IRT1* (Rogers et al., 2000) are conserved in *IAR1*, *Catsup*, and *KE4* (Figure 6C). However, the *IAR1* region corresponding to ZIP transmembrane domains IV and V contains only one transmembrane domain, as predicted by the SMART (Schultz et al., 2000) and PSORT (Nakai and Kanehisa, 1992) motif prediction programs. Although these predicted helices are amphipathic in the ZIPs (Eng et al., 1998), this region in *IAR1*, *Catsup*, and *KE4* contains even more polar residues. Moreover, *IAR1* family members share a conserved Pro residue in the region corresponding to transmembrane domain V of the ZIPs (Figure 6C). This region of *IAR1* is probably important because the conservative substitution of a Lys for an Arg at position 357 results in the IAA–amino acid conjugate resistance of the *iar1-5* allele and because this region is very similar in *IAR1*, *KE4*, and *Catsup* (Figure 6A).

A striking feature of *IAR1* is the presence of several extremely His-rich regions. His represents 17% of the 88 residues between the predicted signal sequence and the first predicted transmembrane domain and 32% of the 40 residues between the second and third transmembrane domains. In addition, the loop between the third and fourth predicted transmembrane domains has seven consecutive His residues (Figure 6A). The *Drosophila* and mammalian *IAR1* homologs are also His-rich in these areas (Figure 6A), suggesting that these residues are important for *IAR1* function. Although the membrane topologies of *IAR1* and its homologs are unknown, two of these areas flank the third predicted transmembrane domain, indicating that both faces of the membrane should have His-rich regions. Five of these regions in *IAR1* are potential metal-binding motifs with the sequence (HX)₃₋₆ (Eng et al., 1998; Guerinot and Eide, 1999). Most ZIP family members contain these motifs between their third and fourth predicted transmembrane domains (Eng et al., 1998), and one of the *IAR1* repeats is in this position (Figure 6A). Although *IAR1* and the ZIPs share only ~10% amino acid identity, this similarity between *IAR1* family members and ZIP family members suggests a common evolutionary ancestor (Figure 6B) and also that *IAR1* may transport or bind metals.

Figure 5. (continued).

Seed from the wild type (Col-0), the *iar1-3* mutant, and three *iar1-3* transgenic lines homozygous for the pBIN*IAR1g* construct, shown in (B), were plated on medium containing 40 μ M IAA-Ala. After 8 days of growth under yellow-filtered light at 22°C, the primary root of each seedling was measured and standardized against growth on unsupplemented medium. Error bars indicate standard errors of the means ($n \geq 17$), and the asterisk indicates a significant difference from the wild type (Student's *t* test, $P < 0.0001$).

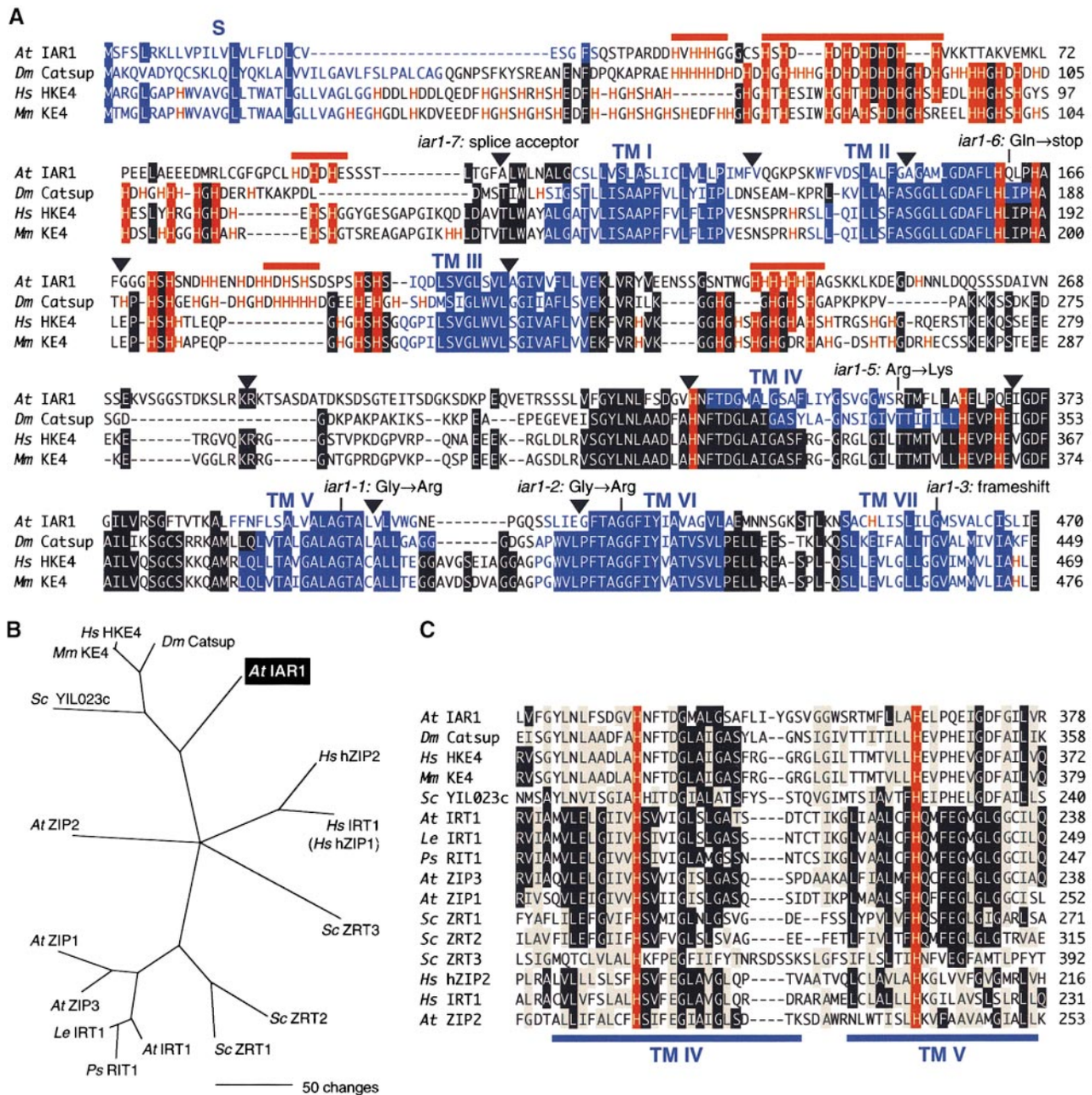


Figure 6. Alignment of IAR1 and Similar Proteins from Other Organisms.

(A) IAR1, *Drosophila* (*Dm*) Catsup, human (*Hs*) HKE4 (Janatipour et al., 1992; Ando et al., 1996), and mouse (*Mm*) KE4 (Abe et al., 1988; St-Jaques et al., 1990) predicted proteins were aligned with the Megalign program (DNASar, Madison, WI) by using the Clustal method (Higgins and Sharp, 1989) with PAM250 residue weights. Residues conserved in at least three proteins are shaded. Triangles indicate the positions of introns in the *IAR1* coding sequence. Positions of *iar1* mutations are shown above the IAR1 sequence, except for *iar1-4*, which is a deletion of the entire *IAR1* gene. The His residues are highlighted in red, and IAR1 potential metal-binding sequences of the type (HX)₃₋₆ are indicated by red bars above the sequence. Regions predicted by the SMART program (Schultz et al., 2000) to be transmembrane domains (TM I to VII) are indicated in blue.

(B) Phylogenetic tree of IAR1 and its relatives. The tree reconstructs the evolutionary relationship between IAR1 family members from **(A)** and ZIP proteins, including Arabidopsis (*At*; Korshunova et al., 1999) and tomato (*Le*) IRT1 (GenBank accession number AF136579); pea (*Ps*) RIT1 (GenBank accession number AF065444); Arabidopsis ZIP1, ZIP2, and ZIP3 (Grotz et al., 1998); yeast (*Sc*) ZRT1 (Zhao and Eide, 1996a), ZRT2 (Zhao and Eide, 1996b), ZRT3 (MacDiarmid et al., 2000), and YIL023c (GenBank accession number P40544); and human (*Hs*) IRT1 (hZIP1) and

Possible Roles for IAR1 in IAA Conjugate Metabolism

IAR1 encodes a putative polytopic membrane protein weakly similar to metal transporters. This single-copy gene lacks close Arabidopsis relatives, at least in the sequenced portion of the genome (~98% complete in September 2000; <http://arabidopsis.org/agi.html>). However, even *iar1* null alleles lack striking morphological phenotypes in the absence of exogenous auxin conjugates, indicating that the plant can compensate for *IAR1* deficiency in laboratory growth conditions. Figure 9 presents two models that could explain how the loss of *IAR1* might result in IAA conjugate resistance.

Arabidopsis encodes a family of amidohydrolases, some of which hydrolyze IAA-amino acid conjugates and contain sequence motifs that suggest an endoplasmic reticulum (ER) lumen localization (Bartel and Fink, 1995; Davies et al., 1999). If *IAR1* encodes an ER-localized IAA conjugate transporter, then loss of *IAR1* function would prevent conjugates from reaching the hydrolases and would thus cause conjugate resistance. However, we did not detect conjugate transport in microsomes prepared from yeast expressing *IAR1* (data not shown). Additionally, *iar1* rescue by the mouse *KE4* cDNA (Figure 8) indicates that *IAR1* and *KE4* probably function similarly in plants and mammals. The lack of an obvious need for such a transporter in mammals argues against a direct role for *IAR1* in IAA-amino acid conjugate transport.

The shared structural features with the ZIP family of transporters (Figures 6 and 7) suggest that *IAR1* might transport a metal. The characterized Arabidopsis amidohydrolases require a metal cofactor, and mutations in these genes can result in IAA-amino acid conjugate resistance (Bartel and Fink, 1995; Davies et al., 1999). Therefore, it is possible that mutations that prevent hydrolases from assembling with or retaining the proper metal might also cause this phenotype. Supplying more of the required cofactor might alleviate IAA-amino acid resistance resulting from such a mutation. This model has precedent in the copper requirement of Arabidopsis ethylene receptors (Rodríguez et al., 1999). Certain copper transporter mutants respond to an ethylene receptor antagonist as if it were ethylene, a phenotype that can be suppressed by providing exogenous copper (Hirayama et

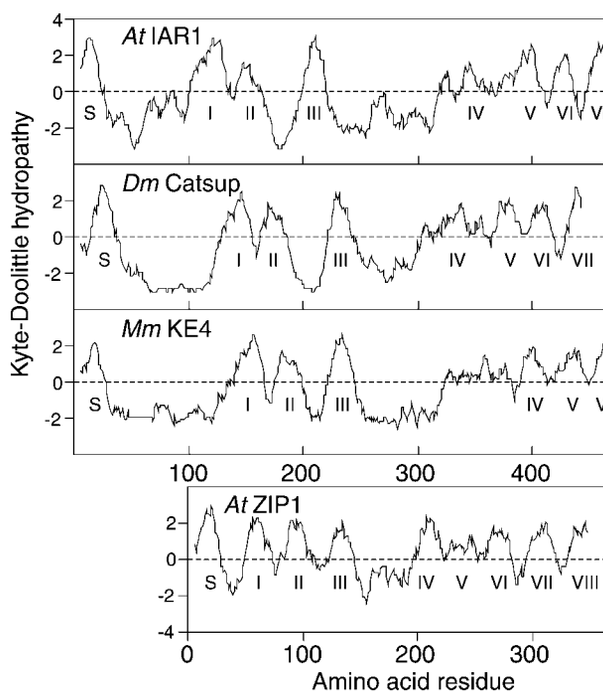


Figure 7. *IAR1* Encodes a Potential Transmembrane Protein.

Hydropathy was calculated for *IAR1*, *Drosophila (Dm) Catsup*, mouse (*Mm*) *KE4* (Abe et al., 1988; St.-Jaques et al., 1990), and Arabidopsis (*At*) *ZIP1* (Grotz et al., 1998) with a window size of 13 and a linear weight variation model (Kyte and Doolittle, 1982). Positive and negative values indicate hydrophobic and hydrophilic regions, respectively. Predicted signal sequences are indicated by S, and the transmembrane domains predicted by the SMART program (Schultz et al., 2000) are numbered.

al., 1999). Manganese is an effective amidohydrolase cofactor, and the partial suppression of the IAA-Ala resistance of *iar1* by exogenous manganese (Figure 4B) suggests that *IAR1* might be an ER-localized manganese transporter. However, *IAR1* expression does not rescue the manganese hypersensitivity of the yeast *pmr1* mutant (data not shown),

Figure 6. (continued).

hZIP2 (Lioumi et al., 1999; Gaither and Eide, 2000). Sequences (from predicted transmembrane domain IV to the end of the proteins) were aligned as described in (A), and the phylogenetic tree was generated by using PAUP 4.0b3a (Swofford, 2000). The bootstrap method was performed for 100 replicates with a maximum parsimony optimality criterion. All characters were weighted equally. Starting trees for the heuristic search were obtained by random stepwise addition, and tree-bisection-reconnection was the branch-swapping algorithm.

(C) Alignment of predicted transmembrane domains IV and V from the ZIP family members described in (B) with the similar region from *IAR1* family members. Identical residues and conservative changes seen in at least seven proteins are shown in black and gray boxes, respectively. Transmembrane domains in the ZIP family described by MacDiarmid (2000) are indicated by blue bars below the alignment. The His residues conserved with those in Arabidopsis *IRT1* that are required for zinc transport (Rogers et al., 2000) are indicated in red.

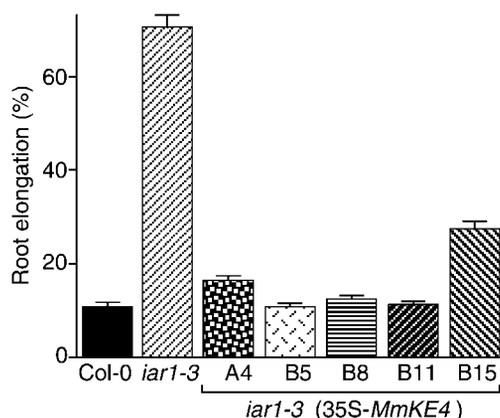


Figure 8. The Mouse *KE4* Gene Can Functionally Substitute for *IAR1*.

Wild-type (Col-0), *iar1-3*, and homozygous T_3 progeny of five independent lines of *iar1-3* plants transformed with 35S-*MmKE4*, a construct containing the mouse *KE4* cDNA (Abe et al., 1988; St.-Jaques et al. 1990) under the control of the cauliflower mosaic virus 35S promoter, were grown for 8 days under yellow-filtered light at 22°C on medium containing 40 μ M IAA-Ala. The primary root of each seedling was measured and standardized against growth on unsupplemented medium. Error bars indicate standard errors of the means ($n \geq 20$).

which is defective in an ER-localized calcium and manganese transporter (Rudolph et al., 1989; Dürr et al., 1998). In addition, Arabidopsis ECA1 transports manganese into the ER and rescues the *pmr1* mutant (Liang et al., 1997). Finally, this model would require that the topology of *IAR1* in the membrane be reversed compared with that of the ZIPs, which transport metals into the cytoplasm (Eng et al., 1998; Guerinot and Eide, 1999; MacDiarmid et al., 2000). Therefore, we think it unlikely that *IAR1* transports manganese into the ER.

A related possibility is that *IAR1* exports an inhibitory metal out of the ER. Zinc and copper inhibit the *IAR3* amidohydrolase in vitro, even when stoichiometric manganese is present (Lasswell, 2000). Yeast ZRT1 and human hZIP2 are plasma membrane localized (Gitan et al., 1998; Gaither and Eide, 2000), whereas yeast ZRT3 is in the vacuolar membrane, where it appears to mobilize stored zinc in zinc-limited cells (MacDiarmid et al., 2000). Therefore, one could speculate that *IAR1* transports zinc, copper, or another inhibitory metal out of the ER and away from the hydrolases (Figure 9A). This model is consistent with the suppression of the *iar1* mutant phenotype by manganese (Figure 4B); excess manganese may outcompete the inhibitory metal from the amidohydrolase and restore activity. It will be interesting to identify the membrane to which *IAR1* localizes, because an ER localization would be consistent with this model.

A recent report on the *Drosophila* Catsup protein (Stathakis et al., 1999), which is ~28% identical to *IAR1*

(Figure 6A), suggests an alternative role for *IAR1* in IAA conjugate responses. The rate-limiting step in catecholamine biosynthesis is the tyrosine hydroxylase (TH)-mediated conversion of L-tyrosine to 3,4-dihydroxy-L-phenylalanine (Kumer and Vrana, 1996). TH is transcriptionally, translationally, and post-translationally regulated (Kumer and Vrana, 1996). *Catsup* mutant flies have normal amounts of TH protein but increased TH activity, which leads to a lethal increase in catecholamine synthesis and suggests that *Catsup* is a post-transcriptional negative regulator of TH (Stathakis et al., 1999) (Figure 9B). Interestingly, Arabidopsis can hydroxylate IAA conjugates (Östin et al., 1998), perhaps as a mechanism to detoxify excess auxin. Thus, if *IAR1* regulates an auxin conjugate hydroxylase the way *Catsup* regulates TH, increased conjugate hydroxylase activity could explain the IAA-amino acid conjugate resistance of the *iar1* mutant (Figure 9B). A preliminary report suggests that *Catsup* interacts directly with TH (Burton et al., 2000), so it will be interesting to isolate proteins that physically or geneti-

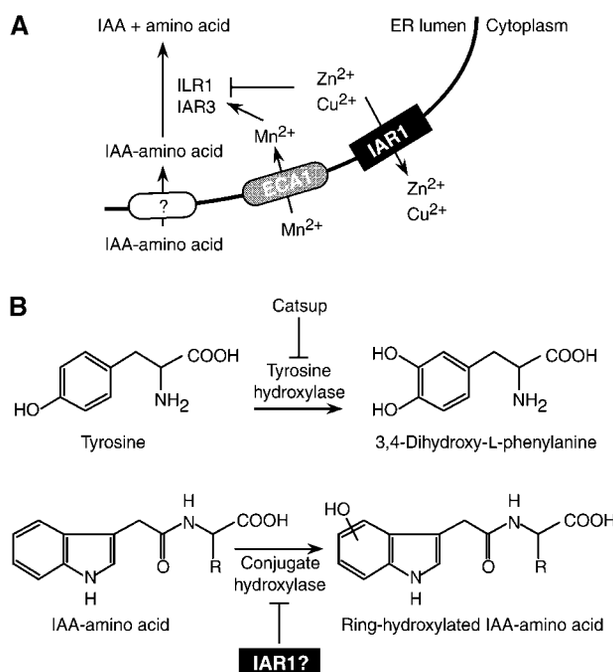


Figure 9. Possible Models for *IAR1* Function.

(A) *IAR1* may transport inhibitory metals out of the ER. This model is suggested by the weak similarity of *IAR1* to ZIP family metal transporters, the suppression of the *iar1* phenotype by the conjugate hydrolase cofactor Mn²⁺, and the inhibition of conjugate hydrolases by Cu²⁺ or Zn²⁺.

(B) *IAR1* may inhibit an auxin conjugate hydroxylase. This model is suggested by the similarity of *IAR1* to the *Drosophila* tyrosine hydroxylase inhibitor *Catsup* (Stathakis et al., 1999) and is consistent with the presence of ring-hydroxylated IAA conjugates in plants. Flat-tipped arrows indicate inhibitory interactions. See text for details.

cally interact with IAR1; these may include the targets of IAR1 action in conjugate metabolism.

IAR1 is a novel membrane protein that is involved directly or indirectly in auxin homeostasis. The identification of the *IAR1* gene demonstrates the capacity of genetic screens to uncover unanticipated players in biochemical processes, and the conservation of IAR1 homologs in metazoans suggests that similar proteins play important roles beyond the plant kingdom.

METHODS

Plant Materials and Growth Conditions

Arabidopsis thaliana seed were surface-sterilized (Last and Fink, 1988) and grown on plant nutrient medium (PNS) containing 0.5% sucrose (Haughn and Somerville, 1986) solidified with 0.6% agar. PNS either contained no supplement or was supplemented with 50 nM to 5 μ M indole-3-acetic acid (IAA; from a 10 mM stock in ethanol), 20 to 150 μ M IAA-amino acid conjugates (from 100 mM stocks in ethanol), 15 μ g/mL kanamycin (Kan; from a 25 mg/mL stock in water), 7.5 μ g/mL glufosinate-ammonium (BASTA; from a 15 mg/mL stock in 25% ethanol), or 100 to 750 μ M MnCl₂ (from a 1 M stock in water). IAA and IAA conjugates were from Aldrich (Milwaukee, WI) or were synthesized (S. LeClere and S.P.T. Matsuda, unpublished data). Plates were wrapped in gas-permeable Leukopor surgical tape (Beierdorf Inc., Norwalk, CT) and grown at 22°C in 24-hr illumination under yellow long-pass filters (25 to 45 μ E m⁻² sec⁻¹) to prevent breakdown of indolic compounds (Stasinopoulos and Hangarter, 1990). Plants transferred to soil (Metromix 200; Scotts, Marysville, OH) were grown at 22 to 25°C under continuous illumination (~200 μ E m⁻² sec⁻¹) with Cool White fluorescent bulbs (Sylvania, Versailles, KY).

Mutant Isolation

The *iar1-1* and *iar1-2* mutants were isolated from pools of Wassilewskija (Ws) seed mutagenized with ethyl methanesulfonate (EMS) as described previously (Normanly et al., 1997). Approximately 24,000 surface-sterilized M₂ seed were grown on PNS supplemented with 50 μ M IAA-Ala. Two-week-old putative mutants with long roots were transferred to soil and allowed to set seed. The resulting M₃ seed were screened separately for resistance to 50 μ M IAA-Ala and for sensitivity to 1 μ M IAA. *iar1-3* and *iar1-4* were isolated similarly from M₂ progeny of fast neutron-mutagenized Columbia (Col-0) *gl1-1* seed (purchased from Lehle Seeds, Round Rock, TX). The other alleles were identified from M₂ pools of EMS-mutagenized *ilr1-1* seed on PNS supplemented with 100 μ M IAA-Leu (*iar1-5* and *iar1-6*) or 70 μ M IAA-Phe (*iar1-7*), as described previously (Davies et al., 1999). The independence of the seven alleles was confirmed by sequencing polymerase chain reaction (PCR) amplification products of genomic DNA. *iar1* mutants were backcrossed at least three times to the parental accession (Col-0 or Ws) before phenotypic analysis.

To obtain the *iar1-1 alf1* homozygous double mutant, an *alf1/ALF1* heterozygous plant (accession Ws) was crossed to the *iar1-1* mutant, and F₂ plants with the phenotypes of both single mutants were transferred to soil and allowed to set seed. Some of these plants were

partially fertile (unlike *alf1/alf1 IAR1/IAR1* plants). These plants yielded 100% IAA-Ala-resistant, *alf1*-like progeny, indicating that they were homozygous for both mutations. Root lengths were measured in progeny from these *iar1-1/iar1-1 alf1/alf1* plants after 10 days at 22°C under continuous yellow-filtered light.

Genetic Mapping of *iar1*

To map the *iar1* mutation, we used segregating populations from crosses between *IAR1/IAR1* (accession Col-0) and *iar1/iar1* (accession Ws) plants. Genomic DNA prepared (Celenza et al., 1995) from 325 IAA-Ala-resistant F₂ plants was scored by using published (Konieczny and Ausubel, 1993; Bell and Ecker, 1994; Nelson et al., 2000) and new PCR-based polymorphic markers. New markers developed for mapping include 13A11RE (amplification with the oligonucleotides 5'-GGTCTTCTGTATTTCACATGGC-3' and 5'-GTGACA-ATAAGATAATTAGATCCGG-3' yielded a 144-bp product with one Tsp509I site in Ws and none in Col-0), T7E4 (amplification with the oligonucleotides 5'-CGGTTTCGAGAGTTGCCTCGT-3' and 5'-GCG-AACTGAATAAAAACCTCCG-3' yielded a 260-bp product with one MseI site in Ws and none in Col-0), *ETR1* (amplification with the oligonucleotides 5'-ATCTCCGTAACCGCTCTTGTCACC-3' and 5'-ACC-ACCACCATCTTGTTCTCTAT-3' yielded a 1200-bp product with one NcoI site in Ws and none in Col-0), and *KNAT2* (amplification with the oligonucleotides 5'-GGTAGCCATATCAGTTCATTG-3' and 5'-ATTTAGTTGACACATCAGCTATC-3' yielded a 660-bp product with three AluI sites in Ws and two in Col-0).

Complementation Library Construction

DNA (~50 μ g) from the T7E4 bacterial artificial chromosome (BAC) (Choi et al., 1995) was partially digested with Sau3AI and separated on 0.8% agarose gel containing 1 mM guanosine to prevent UV light damage to the DNA (Gründemann and Schömig, 1996). Fragments of ~10 kb were purified by using the Qiaex II gel extraction kit (Qiagen, Valenica, CA), ligated into a BamHI-digested pBIN19 plant transformation vector (Bevan, 1984), and transformed into *Escherichia coli* strain DH5 α . A contig of overlapping clones covering the region in which *iar1* mapped was assembled by using a combination of DNA blot hybridization and sequencing. Selected clones were introduced into *Agrobacterium tumefaciens* GV3101 (Koncz and Schell, 1986) by electroporation (Ausubel et al., 1995). *iar1-1* and *iar1-3* mutant plants were transformed with these constructs by the floral dip method (Clough and Bent, 1998), and Kan-resistant transformants were transferred to soil ~14 days after the T₁ seed had been plated on PNS supplemented with 15 μ g/mL Kan. T₂ plants were tested for rescue of the *iar1* IAA-Ala-resistant phenotype by plating on 40 μ M IAA-Ala and examining root elongation after 8 days.

pBIN*IAR1g* was made by cloning a 6.5-kb EcoRI fragment from C37 into the EcoRI site of pBIN19. This construct was transformed into *iar1-3* as described above, and homozygous lines were identified by examining the pattern of Kan resistance in the T₃ generation.

To determine the sequences of the various *iar1* mutations, we amplified genomic DNA by PCR with the following pairs of oligonucleotides: 5'-GAGATGCAAAGAGCAACACTC-3' with 5'-CACTGATGG-AATGGCATTAGG-3', and 5'-CACTGATGGAATGGCATTAGG-3' with 5'-CTGCAAGAACTCCAGCAACAGCTA-3' (40 cycles of 15 sec at 94°C, 15 sec at 55°C, and 30 sec at 72°C); and 5'-GAACCAGGA-CAATCATCGTTG-3' with 5'-GGTACTGCTATCCAATCCGG-3', and

5'-CGTGACGATCACGTGCATCATC-3' with 5'-GCGAATCCGAAT-GAGAATGATC-3' (40 cycles of 30 sec at 95°C, 30 sec at 56°C, and 3 min at 72°C). Products were sequenced directly with the primers used for amplification after sequential precipitations with ethanol, polyethylene glycol, and ethanol (Ausubel et al., 1995). The base changes in most *iar1* alleles alter restriction enzyme recognition sites (Table 1).

cDNA Isolation

A full-length *IAR1* cDNA was isolated by hybridizing an Arabidopsis accession Landsberg *erecta* cDNA library (Minet et al., 1992) with a 1.2-kb HindIII/PstI genomic fragment subcloned from the rescuing C37 construct (Figure 5B) containing the last four exons of the *IAR1* gene. The NotI insert of the cDNA was subcloned into the NotI site of pBluescript KS+ (Stratagene, La Jolla, CA) to create KS/*IAR1c*, which was sequenced by using vector-derived and internal primers (GenBank accession number AF216524).

Transgenic Lines Expressing the Mouse *KE4* cDNA

A full-length cDNA encoding the mouse *IAR1* homolog *KE4* in the vector pME18S-FL3 was purchased from Research Genetics (Huntsville, AL). The ~1.6-kb cDNA insert was excised with StuI and XmnI and ligated into the SmaI site of 35SpBARN (S. LeClere and B. Bartel, unpublished data) to create 35S-*MmKE4*, such that the mouse *KE4* cDNA was under the control of the 35S promoter. This construct was transformed into *iar1-3* plants as described above, transformants were selected on PNS supplemented with 7.5 µg/mL BASTA, and homozygous lines were identified by examining the pattern of BASTA resistance in the T₃ generation.

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Cloning and Characterization of *IARI*, a Gene Required for Auxin Conjugate Sensitivity in *Arabidopsis*

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