

Enzyme Action in the Regulation of Plant Hormone Responses*

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Plants synthesize a chemically diverse range of hormones that regulate growth, development, and responses to environmental stresses. The major classes of plant hormones are specialized metabolites with exquisitely tailored perception and signaling systems, but equally important are the enzymes that control the dose and exposure to the bioactive forms of these molecules. Here, we review new insights into the role of enzyme families, including the SABATH methyltransferases, the methylesterases, the GH3 acyl acid-amido synthetases, and the hormone peptidyl hydrolases, in controlling the biosynthesis and modifications of plant hormones and how these enzymes contribute to the network of chemical signals responsible for plant growth, development, and environmental adaptation.

Plants produce an array of signaling molecules with essential roles in plant growth and development and control responses to environmental stresses, such as drought, herbivory, and pathogen attack. Many plant growth regulators, or phytohormones, were originally isolated as specialized metabolites with molecular structures that reflect their metabolic origins (Fig. 1A) (1). The chemical diversity of these molecules is linked to their biological functions and *in planta* effects through various signaling pathways. Over the past decade, efforts to understand the biosynthesis of plant hormones and their associated perception systems have revealed new biochemical pathways and identified the receptors and signaling events for the major classes of these molecules (Table 1) (2–43).

Although plant development, growth, and environmental responses are all determined by the complex integration of hormone-controlled signaling pathways, changes in cellular concentrations and the chemical structure of a hormone directly affect interaction with cognate receptors to control the duration of activation and potentiation of specific biological effects. The biosynthesis, degradation, and chemical modification of each class of plant hormone contribute to controlling those biological effects. For example, indole-3-acetic acid (IAA),⁴ the

major auxin, triggers hormone responses, whereas modified forms of IAA are used for storage, degradation, or inhibition of auxin signaling (Fig. 1B) (2–5). This minireview provides an overview of plant hormone synthesis; describes new insights into enzymatic modification for controlling plant growth regulators; and highlights how enzymes contribute to the intricate network required for plant growth, development, and environmental adaptations.

Overview of Plant Hormone Biosynthesis

Primary metabolism provides the building blocks of plant growth regulators (Table 1). From a chemical perspective, the molecules that control plant growth are specialized metabolites adapted for interaction with protein receptors to regulate a variety of biological outcomes. Pathways for the biosynthesis of the plant hormones are tightly regulated and integrated to control responses to a diverse array of developmental and environmental inputs.

Amino acid metabolism contributes to the synthesis of ethylene, auxin, and salicylic acid (SA). Production of ethylene, which stimulates fruit ripening and senescence of vegetative tissues, occurs by cyclization of *S*-adenosyl-L-methionine (AdoMet) into 1-aminocyclopropane-1-carboxylic acid (ACC) by ACC synthase and subsequent oxidation into ethylene by ACC oxidase. Regulation of ACC synthase controls ethylene production (10).

Aromatic amino acids are precursors of auxin and SA synthesis. Originally discovered through their effects on plant growth in light and gravity, auxins, especially the predominant hormone IAA, control a range of processes, such as maintenance of apical dominance, shoot elongation, and root initiation (5, 44). Initially reported as a plant growth regulator in 1926 (45), the biosynthesis of IAA from tryptophan has a long history involving multiple proposed pathways (5); however, new evidence suggests that tryptophan aminotransferase and the YUCCA flavin monooxygenase are the major route to auxin (46, 47). Maintenance of bioactive IAA levels requires a balance of synthesis, storage, degradation, transport, and modification (Fig. 1B) (2–5, 48).

SA plays a critical role in plant responses to biotrophic pathogens, which lead to increased SA levels at infection sites (19, 49). SA synthesis requires chorismate. Isotopic feeding studies suggest that conversion of phenylalanine into cinnamic acid and its subsequent metabolism to SA are one route (50–52). In contrast, genetic studies in *Arabidopsis thaliana* suggest that the main route for SA synthesis is conversion of chorismate to isochorismate, followed by breakdown to SA and pyruvate (53). Although bacteria metabolize isochorismate to SA and pyruvate using isochorismate-pyruvate lyase (54), a plant homolog of this enzyme remains unidentified.

Metabolites from the lipid and isoprenoid pathways support the synthesis of jasmonates, cytokinins, brassinosteroids, ab-

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⁴ The abbreviations used are: IAA, indole-3-acetic acid; SA, salicylic acid; AdoMet, *S*-adenosyl-L-methionine; ACC, 1-aminocyclopropane-1-carboxylic acid; ABA, abscisic acid; JA, jasmonic acid; JA-Ile, (+)-7-*iso*-jasmonoyl-

L-isoleucine; MeIAA, methyl-IAA; MES, methylesterase; MeSA, methylsalicylic acid; At, *A. thaliana*; Vv, *V. vinifera*.

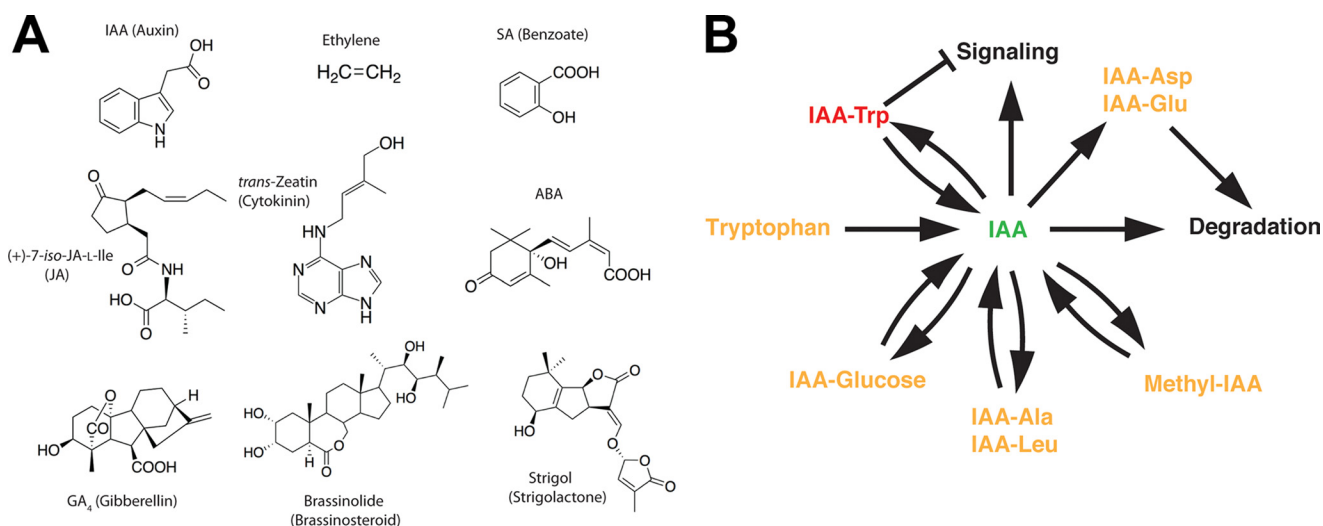


FIGURE 1. **Plant hormones and hormone homeostasis.** A, chemical structures of the major classes of plant hormones. B, representative chemical modifications of the auxin IAA. The bioactive (green), inactive (orange), and anti-auxin (red) forms of IAA are shown. These modifications are only some of the possible chemical changes that occur to this molecule.

TABLE 1

Summary of major plant hormones: biological effects, metabolic precursors, and receptor responses

ET, ethylene; GA, gibberellin; IPP, isopentenyl pyrophosphate; ARR, *Arabidopsis* response regulator.

Hormone	Active form	Effects	Precursor	Receptor
Auxin	IAA	Cell elongation; apical dominance; tropisms; branching; lateral roots	Tryptophan	AUX/IAA: repressor proteins; TIR1/AFBs: auxin F-box-binding proteins; auxin facilitates binding of AUX/IAA with SCF ^{TIR1/AFB} , resulting in degradation of AUX/IAA repressors and altered transcription of auxin-induced genes
ET	Ethylene	Flowering/fruit ripening; stress response; seed germination	Methionine	ETR1/ETR2/ERS1/ERS2/EIN4: histidine kinases (dimers); ET represses repressors of ET response, leading to activation of EIN3 and EREBP families of transcription factors
JA	JA-Ile	Plant defenses against insect herbivores; root growth inhibition; necrotrophic pathogen responses	α -Linolenic acid	COI1/JAZ co-receptor: COI1 F-box protein; component of the SCF ^{COI1} E3 ligase complex; targets JAZ proteins for degradation and leads to altered gene expression
Cytokinins	<i>trans</i> -Zeatin	Cell division; releases lateral buds from apical dominance; delays senescence; root growth	Adenine	CRE1: similar to histidine kinases; activates histidine kinase activity, initiating a phosphorelay that results in phosphorylation of type B ARRs to induce transcription of type A ARRs
Benzoates	SA	Systemic acquired resistance to pathogens; biotrophic pathogen responses	Chorismate	NPR1: transcriptional coactivator; NPR3/ NPR4: CUL3 E3 ligase adaptor proteins
ABA	ABA	Stomatal closure; seed maturation, germination, storage, desiccation tolerance; root/shoot growth; leaf senescence	IPP-derived tetraterpene (phytoene)	PYR/PYL/RCAR: START family of ligand-binding proteins; CHLH: cheletase; GTG1/2: G-proteins
GAs	GA ₁ , GA ₄	Stem elongation; root growth; seed germination; flowering; floral development; fruit growth; stresses	IPP-derived diterpene (geranylgeranyl diphosphate)	GID1: globular protein; GA-GID1 binds DELLA repressors, leading to targeting to the SCF complex and subsequent degradation by the 26 S proteasome
Brassinosteroids	Brassinolide	Cell division/elongation in stems/roots; photomorphogenesis; reproductive development; leaf senescence; stress responses	IPP-derived sesquiterpene (farnesyl diphosphate)	BRI1: dual-specificity kinase; dimerizes with BAK1, initiating phosphorelay prevention of phosphorylation of BES1/BZR1 to alter gene expression
Strigolactones	Strigol	Branching; leaf senescence; root development; plant-microbe interactions	Carotenoids	D14/DAD2: α/β -hydrolase; putative receptor/co-receptor; initiates SCF-mediated signaling via SCF ^{MAX2} complex

scisic acid (ABA), and gibberellins. The biosynthesis of these plant hormones generates a wide range of chemical diversity, much of which remains to be explored with respect to their biological effects.

First identified by their ability to inhibit plant growth, jasmonates affect seed germination, fertility, root growth, and responses to pathogens (1, 13, 55). Plants synthesize jasmonic acid (JA) from α -linoleic acid (13, 42, 43). JA synthesis begins in the plastid with conversion of α -linoleic acid to 12-oxo-phytodienoic acid. Following transport of 12-oxo-phytodienoic acid

to the peroxisome, a reduction reaction and multiple β -oxidation steps generate (+)-7-*iso*-JA (*i.e.* free JA). Conjugation of JA with isoleucine yields the bioactive hormone (+)-7-*iso*-jasmonoyl-L-isoleucine (JA-Ile) (13, 56, 57).

In plants, cytokinins promote cell division, and their synthesis occurs either *de novo* or through recycling of tRNAs containing a uridine at the first anti-codon position (58). In the *de novo* pathway, dephosphorylation of isopentenyl-adenosine-5'-phosphate followed by hydrolysis of the ribose yields *trans*-zeatin. Addition of chemical moieties at the adenosine amine

leads to diverse natural and synthetic cytokinins, such as kinetin (*N*⁶-furfuryladenine) and 6-benzylaminopurine (48).

Biosynthesis of brassinosteroids, ABA, strigolactones, and gibberellins extends from isoprenoid metabolism. Synthesis of brassinosteroids shares similar chemistry with mammalian steroid biosynthesis, as assembly of the core scaffold requires squalene synthase, steroid 5 α -reductase, and cytochrome P450 monooxygenases for formation of campesterol (31). Additional reactions lead to brassinosteroids with various functional groups (31, 55). Carotenoids are the basis for ABA and strigolactones. For ABA synthesis, epoxidation and processing of zeaxanthin to *trans*-violaxanthin occur in the plastid. Subsequent oxidative cleavage in the cytosol produces xanthoxin, which is further metabolized to ABA (22). Similarly, strigolactone synthesis involves breakdown of β -carotene; however, the exact pathway remains unclear (38). Gibberellins are a large group of diterpenoid carboxylic acids (27). Produced by plants, as well as fungi and bacteria, their synthesis begins with cyclization of geranylgeranyl diphosphate into *ent*-kaurene, followed by a series of reactions involving cytochrome P450 monooxygenases and 2-oxoglutarate-dependent dioxygenases (27).

Chemical Modifications

Although the biosynthesis of plant hormones builds from different primary metabolic pathways, these specialized compounds are all modified by common reactions. Modifications of plant hormones mark them for degradation, storage, or activation (Fig. 1*B*). For example, glycosylated forms of IAA, cytokinins, ABA, JA, and SA are common (19, 48, 55). Glycosylation usually leads to inactive storage forms of plant hormones that can be hydrolyzed for activation, but glycosylation of cytokinins tags these molecules for degradation (55). In addition, methylation, amino acid conjugation, hydroxylation, sulfonation, and carboxylation reactions lead to further tailoring of plant hormones (2, 48, 51, 55, 59).

Substantial progress toward understanding how different families of enzymes modify plant hormones provides a foundation for unraveling the biological consequences of these chemical changes. Because methylation and amino acid conjugation of plant hormones are well studied and the proteins involved in both formation and hydrolysis of these modifications have been structurally and biochemically characterized, the rest of this minireview is focused on the methyltransferases, acyl acid-amido synthetases, and the esterases/hydrolases associated with these modifications.

Plant Hormone Methylation

The methylation and demethylation of plant growth regulators rapidly switch chemical activity (Fig. 2*A*). Methyl derivatives of many plant growth regulators were isolated even before the parent compound's hormone activity was identified. For example, methyl jasmonate was isolated from *Jasminum grandiflorum* 2 decades before the role of JA in plant growth was discovered (13, 60). Methylated benzoates are also common across multiple plant species (61–64). Typically, methylated volatiles aid in long-distance communication, and the methylated forms are inactive as hormones and require removal of the modification for effect. Moreover, not all methylated plant

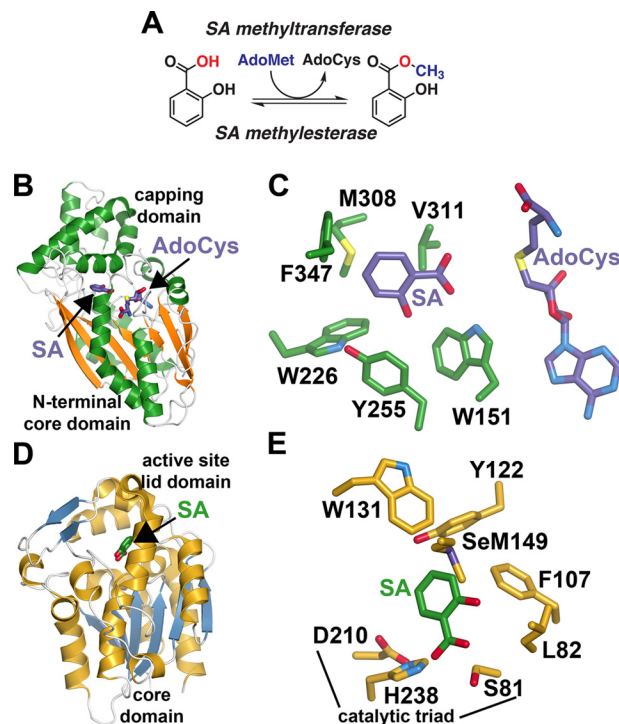


FIGURE 2. Structural overview of the SABATH and MES families. *A*, chemical reaction of SA methylation and demethylation. *AdoCys*, *S*-adenosylhomocysteine. *B*, overall structure of the SA methyltransferase from *C. breweri* (Protein Data Bank code 1M6E) (65) shown as a ribbon diagram. *C*, close-up view of the SA methyltransferase active site. *D*, overall structure of the *N. tabacum* MES SABP2 (Protein Data Bank code 1Y71) (71). *E*, active site view of *N. tabacum* SABP2 showing the positioning of SA and the catalytic triad.

hormones were discovered as volatiles. In *Arabidopsis*, overexpression of a methyltransferase (IAMT1) that converts IAA to methyl-IAA (MeIAA) leads to a curly leaf phenotype, which is typical of perturbed auxin homeostasis, even though MeIAA has not been identified *in vivo* (62). Similarly, *in planta* overexpression of methyltransferases active on gibberellins and JA *in vitro* results in phenotypes characteristic of low gibberellin and JA levels, respectively (63, 64).

In plants, the SABATH methyltransferases catalyze the AdoMet-dependent addition of methyl groups to a range of molecules (61, 65). Named after the first members of the family to be discovered, *i.e.* *SA* methyltransferase, *benzoic acid* methyltransferase, and *theobromine* synthase, these enzymes are encoded by large gene families in plants. For example, the genomes of *Arabidopsis* and rice (*Oryza sativa*) encode 24 and 41 SABATH methyltransferases, respectively. Typically, SABATH proteins function as *O*-methyltransferases that target carboxylic acids, but *N*- and *S*-methyltransferases have been reported (66, 67). Nearly all of the characterized SABATH methyltransferases display a high degree of substrate specificity, which suggests that members of this family evolved for specialized functions (61). An exception is a dual-specific enzyme in *Arabidopsis* that methylates benzoic acid and SA (68).

Crystallographic studies of multiple SABATH methyltransferases reveal the molecular basis of substrate specificity and catalysis (65). Following determination of the structure of SA methyltransferase from *Clarkia breweri* (Fig. 2*B*) (65), x-ray crystal structures of SABATH proteins involved in caffeine bio-

synthesis and IAA methylation were determined (66, 69). The structural fold of the SABATH family is an elongated, parallel, seven-stranded β -sheet, which is common to all AdoMet-dependent methyltransferases, with an α -helical domain capping the active site (Fig. 2B). Although the sequences of SABATH methyltransferases are highly divergent, the residues of the AdoMet-binding site are well conserved (65, 66, 69). In a typical SABATH methyltransferase, such as SA methyltransferase (Fig. 2, B and C), apolar residues interact with the adenine ring of AdoMet, and an invariant aspartate forms a bidentate interaction with the hydroxyl groups of the ribose group. Additional protein-ligand contacts with the carboxylate and amine groups of AdoMet anchor this molecule in the active site for catalysis.

Because SABATH proteins methylate chemically diverse molecules, the substrate-binding site is highly variable in sequence and structure. For example, the substrate-binding site of the *C. breweri* SA methyltransferase (Fig. 2C) consists of multiple apolar residues, with hydrogen bonds between the substrate carboxylic acid and the amine groups of Gln-25 and Trp-151 orienting it toward the AdoMet methyl group (65). In comparison, the substrate-binding site of IAAMT1 utilizes apolar residues for substrate binding (69). Although knowledge of binding residues has aided in the prediction of SABATH methyltransferase function (65, 69), many SABATH proteins with unknown physiological roles remain.

The SABATH methyltransferases catalyze an S_N2 reaction using substrate proximity and desolvation effects to drive the reaction (65). In the proposed reaction, the enzyme binds AdoMet and the methyl acceptor substrate to occlude solvent and orient the AdoMet methyl group toward the acceptor site. Thus, residues provided by the SABATH active site provide a three-dimensional scaffold for binding and orientation of the two substrates.

Plant Hormone Demethylation

Demethylation of plant growth regulators by methyltransferases (MESs) leads to activation of these molecules for their specific biological function (Fig. 2A) (70). MES proteins are part of the α/β -hydrolase enzyme superfamily and share a Ser-His-Asp triad (71, 72). In plants, the first MES discovered was an SA-binding protein from tobacco, which was proposed to function in methylsalicylic acid (MeSA) perception and signaling (73). Later work showed that this protein hydrolyzed MeSA to SA (71). As with the SABATH family, plants encode multiple MES-like genes. For example, *Arabidopsis* contains 20 MES-like genes, two of which (*A. thaliana* (*At*) *MES19* and *AtMES20*) are likely pseudogenes (72).

Crystallographic analysis of the tobacco (*Nicotiana tabacum*) MES SABP2 (salicylic acid-binding protein-2) provided the first structural insight into this protein family (71). The overall structure contains a core six-stranded parallel β -sheet surrounded by α -helices with an active site lid domain consisting of a three-stranded antiparallel β -sheet and three α -helices (Fig. 2D). The active site is at the interface of these two domains. The aromatic ring of SA binds to a hydrophobic pocket in the cap domain, and the main domain contains the catalytic triad responsible for hydrolysis of the methyl group (Fig. 2E). Although some of the MES proteins in *Arabidopsis* contain a

non-canonical catalytic triad in which the serine is replaced with an aspartate, this substitution does not compromise esterase activity (70, 72).

Biochemical screening of 15 *Arabidopsis* MES proteins identified enzymes that hydrolyzed MeIAA, MeSA, methyl jasmonate, and the artificial substrate *p*-nitrophenyl acetate (70). None of the *Arabidopsis* MESs were active with methyl gibberellins, and some showed no activity with any compound tested (70). Subsequent studies showed that AtMES16 demethylates chlorophyll catabolites and is important for chlorophyll degradation (74). Further work aimed at examining the expression patterns and quantifying substrate specificities of the plant MES family promises to develop a better understanding of the molecular and physiological roles of these proteins.

Plant Hormone-Amino Acid Conjugation

In plants, amino acid conjugates to IAA and JA dramatically alter the biological roles of these molecules (2–5, 13, 42, 43). Amino acid conjugation of auxins plays a central role in their homeostasis (2). For IAA, the free acid is the biologically active form of the hormone, with amino acid conjugation leading to inactivation (75). The metabolic fate of conjugated IAA depends on which amino acid is attached (Fig. 1B). Conjugation of either alanine or leucine to IAA leads to an inactive but readily hydrolyzed storage form (75). Conjugation of IAA with either aspartate or glutamate leads to hormone degradation (75). IAA-Trp acts as an anti-auxin to inhibit plant growth effects but does not compete with IAA for binding to the TIR1 auxin receptor (76). Typically, modification of IAA negates its plant growth effects. In contrast, amino acid conjugation of JA leads to formation of the biologically active hormone JA-Ile, which binds to the COI1 ubiquitin ligase/JAZ protein co-receptor to elicit its effects (56, 77, 78). Interestingly, JA-Trp also functions as an anti-auxin and suggests possible cross-talk that aids in balancing auxin and jasmonate signaling (77).

In plants, the enzymes that catalyze amino acid conjugation of plant hormones belong to the GH3 (Gretchen Hagen 3) family of acyl acid-amido synthetases (79). The first GH3 gene was identified in 1985 as an early auxin-responsive gene in *Glycine max* (soybean) (79). As with the SABATH and MES families, multiple genes encode GH3 proteins in each plant. For example, *A. thaliana* and rice encode 19 and 13 GH3 proteins, respectively (79–81). Genetic and physiological studies of various GH3 proteins indicate a diverse range of biological functions for these enzymes in jasmonate and auxin hormone signaling and for SA-related pathogen responses.

Staswick *et al.* (82) first characterized the *A. thaliana* GH3.11 (AtGH3.11/JAR1) protein after identifying the *jar1* (jasmonate resistant 1) mutant, which showed defective JA signaling, including reduced male fertility and resistance to exogenous JA treatment. Sequence analysis of AtGH3.11/JAR1 identified low homology with firefly luciferase and the ANL family of adenylating enzymes (83). Named after its founding members (acyl-CoA synthetases, non-ribosomal peptide synthetases, and luciferase), ANL enzymes catalyze a two-step reaction via an adenylated reaction intermediate (83). The GH3 acyl acid-amido synthetases use ATP to adenylate an acyl acid substrate in the first half-reaction. Nucleophilic attack by the

amine of an amino acid leads to release of AMP and formation of an amino acid conjugate (84, 85).

Subsequent studies in *Arabidopsis* and rice identified multiple GH3 proteins with roles in plant growth and development linked to auxin hormone signaling (86–91). Later biochemical analysis of these enzymes demonstrated conjugation of IAA to various amino acids (92). In *Arabidopsis*, knock-outs of the IAA-utilizing GH3 proteins show little effect on plant growth (86–91), which may result from functional redundancy. Over-expression lines for a few IAA-utilizing GH3 proteins in *Arabidopsis* all result in a dwarf phenotype along with other traits indicative of altered auxin signaling (86–91). In addition to jasmonate and auxin signaling, *Arabidopsis* mutants of the *AtGH3.12/PBS3* gene display increased disease susceptibility to virulent and avirulent forms of the pathogen *Pseudomonas syringae* (93). These mutants suggest that amino acid conjugates of SA and related benzoates may function as bioactive inducers of plant pathogen defense responses (94).

Although genetic studies link GH3 proteins to jasmonate, auxin, and SA responses, the biochemical understanding of these proteins was limited to a few examinations of reaction chemistry and substrate specificity (77, 81, 82). New insights into how these proteins modify plant hormones came from the crystal structures of two *Arabidopsis* GH3 proteins (*AtGH3.11/JAR1* and *AtGH3.12/PBS3*) and a grapevine (*Vitis vinifera* (Vv)) GH3 protein (*VvGH3.1*) (95, 96). As mentioned above, *AtGH3.11/JAR1* catalyzes formation of JA-Ile, the bioactive jasmonate hormone (77). Although mutants of *AtGH3.12/PBS3* display SA-related phenotypes in *Arabidopsis*, SA is not a substrate of this protein, and the physiologically relevant substrate is not known (94). The grapevine protein (*VvGH3.1*) conjugates IAA with aspartate (96). These structures define the overall GH3 fold as a large N-terminal domain with a β -barrel and two β -sheets flanked by α -helices and a smaller C-terminal domain consisting of a single four-stranded β -sheet flanked by two α -helices on each side (Fig. 3A). The active site is located at the interdomain interface, and the C-terminal domain is conformationally flexible, with its movement linked to each half-reaction (95).

The crystal structures reveal how the GH3 enzymes conjugate various amino acids to chemically diverse plant hormones. In each structure, residues of the N-terminal domain form the acyl acid/hormone-binding site. In *AtGH3.11/JAR1*, JA-Ile is bound in an elongated hydrophobic tunnel, with the oxylipin tail stacking across the surface of a tryptophan (95). The protein was crystallized with a racemic mixture of JA with the final structure including (–)-JA-Ile because the refinement statistics were modestly better with this enantiomer than with (+)-*iso*-JA-Ile; however, the quality of the ligand density allows modeling of either product. It is possible that the density represents a mixture of product enantiomers. The only polar interaction is a water-mediated hydrogen bond between the JA ketone group and a histidine side chain (95).

In comparison, the hormone-binding site of the IAA-utilizing *VvGH3.1* protein reveals extensive changes that provide a different hydrophobic site to accommodate the substrate indole ring and a hydrogen bond between an active site tyrosine hydroxyl group and the amine of the indole ring (Fig. 3B) (96).

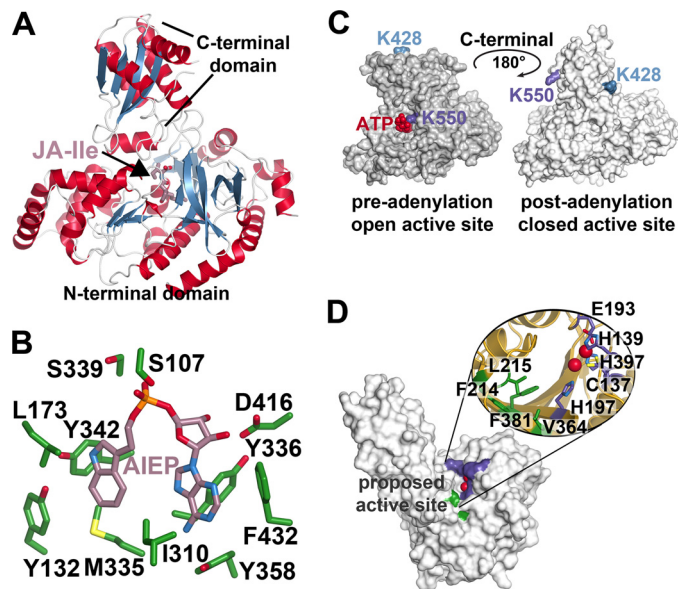


FIGURE 3. Structural overview of the GH3 acyl-amido synthetases and peptidyl hydrolase ILL. A, overall structure of *AtGH3.11/JAR1* (Protein Data Bank code 4EPL) (95). B, active site view of the *VvGH3.1* protein in complex with the non-hydrolyzable adenylation intermediate analog adenosine 5'-[2-(1*H*-indol-3-yl)ethyl]phosphate (Protein Data Bank code 4B2G) (96). C, surface representations of *AtGH3.12/PBS3* showing the rotation of the C-terminal domain during the pre-adenylation open active site conformation (Protein Data Bank code 4EWW) (95) and the post-adenylation closed active site conformation (code 4EPM) (95). D, surface view of the *AtILL2* peptidyl hydrolase (Protein Data Bank code 1XMB) (99). Residues corresponding to the apolar pocket proposed as the substrate-binding site are colored green, and the residues forming the canonical metal-binding site are colored purple.

Surprisingly, there are no interactions with the substrate carboxylate group, but the *VvGH3.1* structure was determined with adenosine 5'-[2-(1*H*-indol-3-yl)ethyl]phosphate, a non-hydrolyzable adenylated IAA analog lacking the IAA carbonyl moiety, so additional interactions may form with the authentic substrate (Fig. 3B). Previous analyses of the IAA-utilizing GH3 proteins demonstrate activity with a variety of different auxins, including phenylacetic acid and naphthaleneacetic acid, which differ in chemical structure from IAA, but the basis for binding these molecules remains unclear (85). Although the physiological substrate of *AtGH3.12/PBS3* is not known, the hormone site of this enzyme contains a number of polar residues, suggesting distinct substrate specificity compared with the JA- and IAA-conjugating enzymes.

Across the structures, the nucleotide site is identical to residues conserved in not only GH3 proteins but also the ANL superfamily (83, 95, 96). Three motifs define this site. The first motif is a serine-, threonine-, and glycine-rich P-loop that forms contacts with the α -phosphate; these interactions are essential for catalysis (95). Residues of the second motif (*i.e.* YGSSE) provide additional binding interactions. The tyrosine contacts the adenosine ring, the second serine hydrogen bonds with the α -phosphate, and the glutamate coordinates binding of the Mg^{2+} ion required for ATP binding. The third motif provides an aspartate for hydrogen bonding to the ribose hydroxyls of the bound nucleotide. Overall, these features orient ATP for attachment of the hormone carboxylic acid at the α -phosphate in the first half-reaction, leading to an adenylated intermediate (95).

Following adenylation of the acyl acid substrate and pyrophosphate release, rotation of the C-terminal domain by 180° yields an active site that covers the reaction intermediate (Fig. 3C) (95). This rotation prevents either the release or hydrolysis of the reaction intermediate before binding of the amino acid substrate. Rotation of the C-terminal domain also creates a tunnel for the amino acid to enter and orient for nucleophilic attack on the adenylated intermediate (95, 96). Although no crystal structure of a GH3 protein with an amino acid bound is currently available, the placement of the isoleucine group of JA-Ile in AtGH3.11/JAR1 and a bound malate in VvGH3.1 suggests a putative amino acid-binding site (95, 96).

Biochemically, the GH3 proteins catalyze the conjugation of amino acids to acyl acids via an adenylation reaction. Through this simple modification, these proteins regulate levels of active and inactive forms of jasmonates, auxins, and benzoates. The capacity of GH3 proteins to modify multiple related substrates may also allow for changes of metabolic pools across a biosynthetic pathway, in addition to modifying the final hormone product.

Plant Hormone Hydrolases

As with methylated hormones, amino acid-conjugated hormones can be hydrolyzed back to the free hormone and amino acid. The first hydrolase discovered with this activity was identified from a mutant screen of *Arabidopsis* looking for resistance to IAA-Leu treatment (97). The ILR (IAA-Leu-resistant) protein belongs to the M20 peptidase superfamily, of which there are six other homologs in *Arabidopsis* (97). Most enzymes in the M20 peptidase family are Zn²⁺-dependent, but the ILR proteins prefer Mn²⁺ (98). The crystal structure of an ILR protein from *Arabidopsis* (i.e. AtILL2) shows the bidomain fold representative of the superfamily (Fig. 3D) (99). The larger domain contains the active site and consists of eight β -strands with α -helical bundles on both sides. The smaller satellite domain adopts an α/β -sandwich topology. In other M20 peptidases, this domain serves as a dimerization interface; however, AtILL2 functions as a monomer (99). Although AtILL2 was crystallized in the absence of metal ions, it contains a canonical histidine-rich metal-binding site (Fig. 3D). Because the crystal structure of AtILL2 is an apoenzyme form, computational ligand docking experiments identified a putative IAA-binding site and suggested that the difference between a leucine and a tyrosine in the site may control access of substrates with smaller versus larger amino acid conjugates, respectively (99). Crystal structures with non-hydrolyzable analogs of the substrates or reactions products along with biochemical analysis of each family member are needed to understand the mechanism of substrate specificity in the plant hormone hydrolases.

Summary

The integration of multiple signaling pathways and hormone responses determines how plants develop and grow; however, the perception of small molecule signals by receptors is only one piece of this biological puzzle. The range of responses controlled by plant hormones requires enzyme action in biosynthetic, storage, degradation, and mobilization pathways and through recognition by cognate receptors (many of which are

also enzymes in plants). These processes control the dose and duration of exposure to plant growth regulators.

Recent efforts summarized here provide new molecular insights into how large enzyme families catalyze similar types of modifications on chemically diverse plant growth regulators to alter their biological functions. In addition to the growing biochemical and structural understanding of enzymatic regulation of plant hormones, efforts to elucidate the temporal and spatial expression patterns of hormone-modifying enzymes will provide a physiological context for these reactions. The specialized chemicals that plants use to control their growth are a rich pallet for coloring plant responses to internal and external stimuli.

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