

# Significance of Inducible Defense-related Proteins in Infected Plants

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## Key Words

antimicrobial activity, defense signaling, developmental regulation, pathogenesis-related proteins, resistance, stress alleviation

## Abstract

Inducible defense-related proteins have been described in many plant species upon infection with oomycetes, fungi, bacteria, or viruses, or insect attack. Several types of proteins are common and have been classified into 17 families of pathogenesis-related proteins (PRs). Others have so far been found to occur more specifically in some plant species. Most PRs and related proteins are induced through the action of the signaling compounds salicylic acid, jasmonic acid, or ethylene, and possess antimicrobial activities in vitro through hydrolytic activities on cell walls, contact toxicity, and perhaps an involvement in defense signaling. However, when expressed in transgenic plants, they reduce only a limited number of diseases, depending on the nature of the protein, plant species, and pathogen involved. As exemplified by the PR-1 proteins in *Arabidopsis* and rice, many homologous proteins belonging to the same family are regulated developmentally and may serve different functions in specific organs or tissues. Several defense-related proteins are induced during senescence, wounding or cold stress, and some possess antifreeze activity. Many defense-related proteins are present constitutively in floral tissues and a substantial number of PR-like proteins in pollen, fruits, and vegetables can provoke allergy in humans. The evolutionary conservation of similar defense-related proteins in monocots and dicots, but also their divergent occurrence in other conditions, suggest that these proteins serve essential functions in plant life, whether in defense or not.

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**SA:** salicylic acid  
**JA:** jasmonic acid  
**ET:** ethylene  
**R:** resistance gene  
**PR:**  
pathogenesis-related  
**SAR:** systemic  
acquired resistance

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## INTRODUCTION

Plants possess both preformed and inducible mechanisms to resist pathogen invasion. Extant morphological barriers, secondary metabolites (phytoanticipins), and antimicrobial proteins must be avoided or overcome for pathogens to be able to invade a plant. Once contact has been established, elicitors produced and released by the pathogen induce further defenses, comprising the reinforcement of cell walls, the production of phytoalexins, and the synthesis of defense-related proteins (133). In the past few years, plant microarray data have been collected showing that in both compatible and incompatible plant-pathogen interactions, hundreds of genes are up- and downregulated. In many cases, differences between susceptibility and resistance are associated with differences in the timing and magnitude of these changes rather than with the expression of different sets of genes (141). As evident from the occurrence of mainly *Arabidopsis* mutants with enhanced disease susceptibility (*eds*) (115), plants possess a basal resistance against their pathogens, which is overcome, manipulated, or suppressed by these pathogens to allow successful infection and tissue colonization (25, 98, 127, 182). *Arabidopsis* mutants affected in the production or action of the signaling compounds salicylic acid (SA), jasmonic acid (JA), or ethylene (ET) likewise show an enhanced disease-susceptibility phenotype upon infection by specific pathogens, indicating that these regulators play a role in the basal resistance against these pathogens (77, 143). The same regulators have also been implicated in certain types of nonhost resistance and in *R*-gene-mediated resistance, suggesting that expression of these different types of resistance involves activation of partly similar defensive mechanisms (54, 144). Whether or not a plant turns out to be susceptible or resistant is likely determined by the speed and magnitude with which these mechanisms are activated and expressed and by their effectiveness against

individual pathogens with different modes of attack.

Because the availability of microarrays is largely limited to *Arabidopsis*, current views are based mostly on a few selected interactions, e.g., those of *Arabidopsis* with the oomycete *Hyaloperonospora parasitica* (87), the fungus *Alternaria brassicicola* (156), and the bacterium *Pseudomonas syringae* pv. *maculicola* (46). Data from other plant species infected by other pathogens or attacked by insects are usually interpreted with reference to the former. Although different plant species react to infection by activation of similar defensive mechanisms, their regulation may differ in important details. For instance, basal resistance against the fungus *Botrytis cinerea* is regulated by SA in tomato but by JA and ET in tobacco (1, 45). Systemic JA-mediated induced resistance against insect herbivory in tomato involves the mobile signaling peptide systemin, but in other plant species no obvious counterpart of this transportable signal is evident (61). Such differences may be at the basis of the specificity in plant-pathogen interactions, as only a small number of potential pathogens is able to infect any given plant species.

Among the genes that in *Arabidopsis* are activated in response to infections, many appear to be involved in transcriptional regulation, signal transduction, various metabolic activities, and defense (32). The current status of transcription factors and signaling pathways involved in plant reactions to microorganisms has been the subject of several recent reviews (19, 38) and is not discussed here; neither are changes in metabolic pathways that occur during disease development, or involved in the readjustment of plant functioning, in response to an infecting pathogen. Instead, this review concentrates on inducible proteins that have been implicated in active defense and could play a role in restricting pathogen development and spread in the plant. Most of these defense-related proteins correspond to pathogenesis-related proteins (PRs) or the products of so-called SAR genes, which were identified several years ago as being associated

with resistance reactions of plants to various pathogens (12, 149). The study of these proteins has now shifted almost entirely to the analysis of the expression of the corresponding genes, but the relationships between the proteins and their encoding genes are not always obvious. Defense-related proteins commonly occur as families of closely related homologues whose mRNAs may cross-hybridize to greater or lesser extents. Corresponding mRNAs in newly studied plant species are commonly identified on the basis of cross-hybridization with heterologous probes. At the gene level, sequences are annotated on the basis of homology to an arbitrary member of the family without knowledge of whether and where the gene(s) are expressed. Of several proteins that are grouped within the same family on the basis of their serological relatedness and/or biological activity, the amino acid sequences are only partly known, if at all. Under these circumstances, in many cases the correspondence between genes/mRNAs and proteins is far from clear. For example, single, chemically induced PR-1, PR-2, and PR-5 proteins from *Arabidopsis* intercellular washing fluid were purified and characterized, and genes corresponding to these proteins were cloned and sequenced (148). However, it recently became clear that the gene annotated as *PR-1* on the Affymetrix *Arabidopsis* ATH1 GeneChip (At2g19990) is actually a homologue of the *PR-1* gene (At2g14610) that corresponds to the characterized protein (78). With the exception of a number of nucleotide sequences in tobacco and tomato—for which the PR-proteins have been fully classified—it is generally unclear which gene/mRNA within a family corresponds to which protein, or whether specific genes are expressed at all.

The resulting confusion is exacerbated by the use by different authors of various designations for the same gene or protein [e.g., Table 2 in (155)]. Although a unifying nomenclature for PR-proteins was described as early as 1994 (153), classification of PRs beyond tobacco and tomato has been hampered by the paucity of data on the properties of the

## PR-PROTEINS IN WINE

Grape berries accumulate PRs during ripening and as a result of induction by biotic or abiotic stress. Because PRs are resistant to proteolytic attack and low pH, they survive vinification and can adversely affect the clarity and stability of the wine. For instance, haziness develops as a result of the presence of mainly chitinases and thaumatin-like proteins. To remove the proteins, wines are cleared by adding an adsorptive compound, followed by the settling or precipitation of partially soluble components from the wine during layering. This process may also remove important wine aroma and flavour compounds and reduce wine quality.

It is assumed that the presence of PRs in the berries contributes to basal resistance against pathogens. Thus, a benefit to the plant is a nuisance to the winemaker (40).

proteins in other plant species and the restriction of data to the mRNA (cDNA) level. Originally, PRs were classified on the basis of their characteristics as plant proteins induced in pathological or related situations. Related proteins occurring in the absence of pathogen infection were to be referred to as “PR-like” proteins (PRLs). The term PRL was not adopted by the scientific community, in part because the distinction between PRs and PRLs became blurred when it was found that specific PRs were sometimes present in healthy tissues and the levels of certain pre-existing PRLs were increased after pathogen infection. The term “pathogenesis-related proteins” then became a collective term for all microbe-induced proteins and their homologues to the extent that enzymes such as phenylalanine ammonia-lyase (PAL), peroxidase, and polyphenoloxidase, which are generally present constitutively and only increased during most infections, are often also referred to as PRs. The latter was never intended to be the case (153), as there are numerous enzyme activities that are increased in response to pathogen attack and which may also play a role in defense (53). For this reason, in this review the general term “inducible defense-related proteins” is used to indicate

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**PRL:** PR-like protein

**PAL:** phenylalanine ammonia-lyase

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**Table 1** Recognized families of pathogenesis-related proteins

Family	Type member	Properties	Gene symbols
PR-1	Tobacco PR-1a	Unknown	<i>Ypr1</i>
PR-2	Tobacco PR-2	$\beta$ -1,3-glucanase	<i>Ypr2</i> , [ <i>Gns2</i> (' <i>Glb</i> ')]
PR-3	Tobacco P, Q	Chitinase type I, II, IV, V, VI, VII	<i>Ypr3</i> , <i>Chia</i>
PR-4	Tobacco 'R'	Chitinase type I, II	<i>Ypr4</i> , <i>Chid</i>
PR-5	Tobacco S	Thaumatocin-like	<i>Ypr5</i>
PR-6	Tomato Inhibitor I	Proteinase-inhibitor	<i>Ypr6</i> , <i>Pis</i> (' <i>Pin</i> ')
PR-7	Tomato P <sub>69</sub>	Endoproteinase	<i>Ypr7</i>
PR-8	Cucumber chitinase	Chitinase type III	<i>Ypr8</i> , <i>Chib</i>
PR-9	Tobacco "lignin-forming peroxidase"	Peroxidase	<i>Ypr9</i> , <i>Prx</i>
PR-10	Parsley "PR1"	Ribonuclease-like	<i>Ypr10</i>
PR-11	Tobacco "class V" chitinase	Chitinase, type I	<i>Ypr11</i> , <i>Chic</i>
PR-12	Radish Rs-AFP3	Defensin	<i>Ypr12</i>
PR-13	Arabidopsis THI2.1	Thionin	<i>Ypr13</i> , <i>Thi</i>
PR-14	Barley LTP4	Lipid-transfer protein	<i>Ypr14</i> , <i>Ltp</i>
PR-15	Barley OxOa (germin)	Oxalate oxidase	<i>Ypr15</i>
PR-16	Barley OxOLP	Oxalate-oxidase-like	<i>Ypr16</i>
PR-17	Tobacco PRp27	Unknown	<i>Ypr17</i>

Further details can be found at <http://www.bio.uu.nl/~fytopath/PR-families.htm>.

**Chitinases:**

enzymes that cleave poly- $\beta$ -1,4-N-acetylglucosamine (chitin)

**Lysozyme:**

an enzyme that cleaves bacterial peptidoglycans

**Thaumatocin:**

a sweet protein from the fruit of the African shrub *Tbaumatococcus daniellii*

those proteins that are mostly nondetectable in healthy tissues and for which induction at the protein level has been demonstrated after infection by one or more pathogens. Inducible defense-related proteins encompass both the known PR-protein families and nonclassified proteins meeting the criteria above. The term "defense-related" refers to the fact that these proteins are induced in association with resistance responses but does not by itself imply a functional role in defense. However, because some of these proteins have at least potential antimicrobial activities, a role in resistance to pathogens appears plausible and is discussed below.

**FAMILIES OF DEFENSE-RELATED PROTEINS AND THEIR OCCURRENCE**

Inducible defense-related proteins were first discovered in tobacco reacting hypersensitively to *Tobacco mosaic virus* (TMV) and later shown to occur in plant species from at least 13 families upon infection by oomycetes, fungi, bacteria, viruses, and viroids, as well as ne-

matode or insect attack (149). The recognized PRs have been extensively reviewed (12, 29, 72, 75, 179) and currently comprise 17 families of induced proteins (**Table 1**). The families are numbered in the order in which they were discovered. A type member, usually the first or most prominent one, was chosen and families were defined further on the basis of their common biochemical and biological properties. A role of several families in limiting pathogen activity, growth, and spread fits with the identification of the PR-2 family as  $\beta$ -1,3-endoglucanases and the PR-3, -4, -8, and -11 as endochitinases, which could act against fungi. The chitinases, as well as the proteinase inhibitors (PR-6), could also target nematodes and herbivorous insects. Members of the PR-8 family also possess lysozyme activity and may be directed against bacteria, whereas defensins (PR-12) (80, 142) and thionins (PR-13) (9, 36) both have broad antibacterial and antifungal activities. Some lipid transfer proteins (PR-14) have antifungal and antibacterial activities (44) and members of the PR-1, and the thaumatocin-like PR-5 families have been associated with

activity against oomycetes. Notably, the prominent PR-1 proteins are often used as markers of the enhanced defensive state conferred by pathogen-induced systemic acquired resistance (SAR), but their biological activity has remained elusive (155). PR-7 is an endoproteinase that is the most conspicuous PR in tomato (66). It might aid in microbial cell wall dissolution. PR-9 is a specific type of peroxidase that could act in cell wall reinforcement by catalyzing lignification (104) and enhance resistance against multiple pathogens. PR-10 shows homology to ribonucleases, and some members do have weak ribonuclease activity (16). There are no other families of PRs that are directed specifically against viruses, and it has sometimes been assumed that the ribonuclease activity of PR-10-type proteins points to a role in defense against these pathogens (102). However, recently an antifungal PR-4-type protein from wheat was shown to also possess ribonuclease activity (18). The families PR-15, -16, and -17 have been added recently. PR-15 and -16 are typical of monocots and comprise families of germin-like oxalate oxidases and oxalate oxidase-like proteins with superoxide dismutase activity (8), respectively. These proteins generate hydrogen peroxide that can be toxic to different types of attackers or could directly or indirectly stimulate plant-defense responses (e.g., 34, 62). PR-17 proteins have been found as an additional family of PRs in infected tobacco, wheat, and barley and contain sequences resembling the active site of zinc-metalloproteinases (21), but have remained uncharacterized so far. A putative novel family (PR-18) comprises fungus- and SA-inducible carbohydrate oxidases, as exemplified by proteins with hydrogen peroxide-generating and antimicrobial properties from sunflower (27). Not all families seem to be represented in all plant species, and occurrence and properties of different members within a family may differ strongly.

The described SAR genes, whose coordinate induction correlates with the onset of SAR, encompass most PR genes, as well as

a protein designated SAR 8.2, whose levels are strongly increased in tobacco in response to TMV infection (169). Among other nonclassified proteins resembling PRs in their induction by pathogens are an amylase in tobacco (55), the DRR206 protein in pea (26), cell wall hydroxyproline-rich glycoproteins (37), glycine-rich proteins (119), polygalacturonase-inhibiting proteins (31), lipoxygenases (41), and lipase-like gene products (64). Important groups of antimicrobial proteins that are present in various plant organs but have not been reported to be induced by pathogen attack—and hence are not PRs—are ribosome-inactivating proteins, lectins, and various types of cysteine-rich peptides (12).

In leaves, PRs appear to be present both in epidermal and mesophyll cells, as well as in the vascular bundles. As an example, in response to infection by *Phytophthora infestans*, potato accumulates PR-1b in the vicinity of the successfully colonized leaf area and of the epidermal cell layer in particular. Additional locations within infected leaves were stomatal guard cells, glandular trichomes, crystal idioblasts and the vascular bundles (57). Many defense-related proteins are synthesized with an N-terminal signal peptide determining translocation into the ER, followed by secretion into the apoplast. These proteins accumulate extracellularly and can be collected easily in intercellular washing fluid. PR-type proteins have been collected from xylem fluid of tomato, broccoli, rape, pumpkin, and cucumber (17, 67, 111, 112) and from guttation fluid of barley leaves (51), suggesting that secretion into the veins entails uptake and transport in the transpiration stream. Other proteins have additional extensions specifying deposition into the vacuole. PR-10-type proteins are the only family of which all members seem to be cytoplasmic.

After their characterization as proteins induced as a result of pathogen or insect attack, many of the same or closely related proteins/mRNAs have been found to be expressed in a developmentally controlled,

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#### Systemic acquired resistance (SAR):

the phenomenon that plants acquire an enhanced defensive capacity against subsequent pathogen attack as a result of a primary, limited infection

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### Somatic embryogenesis:

embryo formation as a result of dedifferentiation of diploid cells

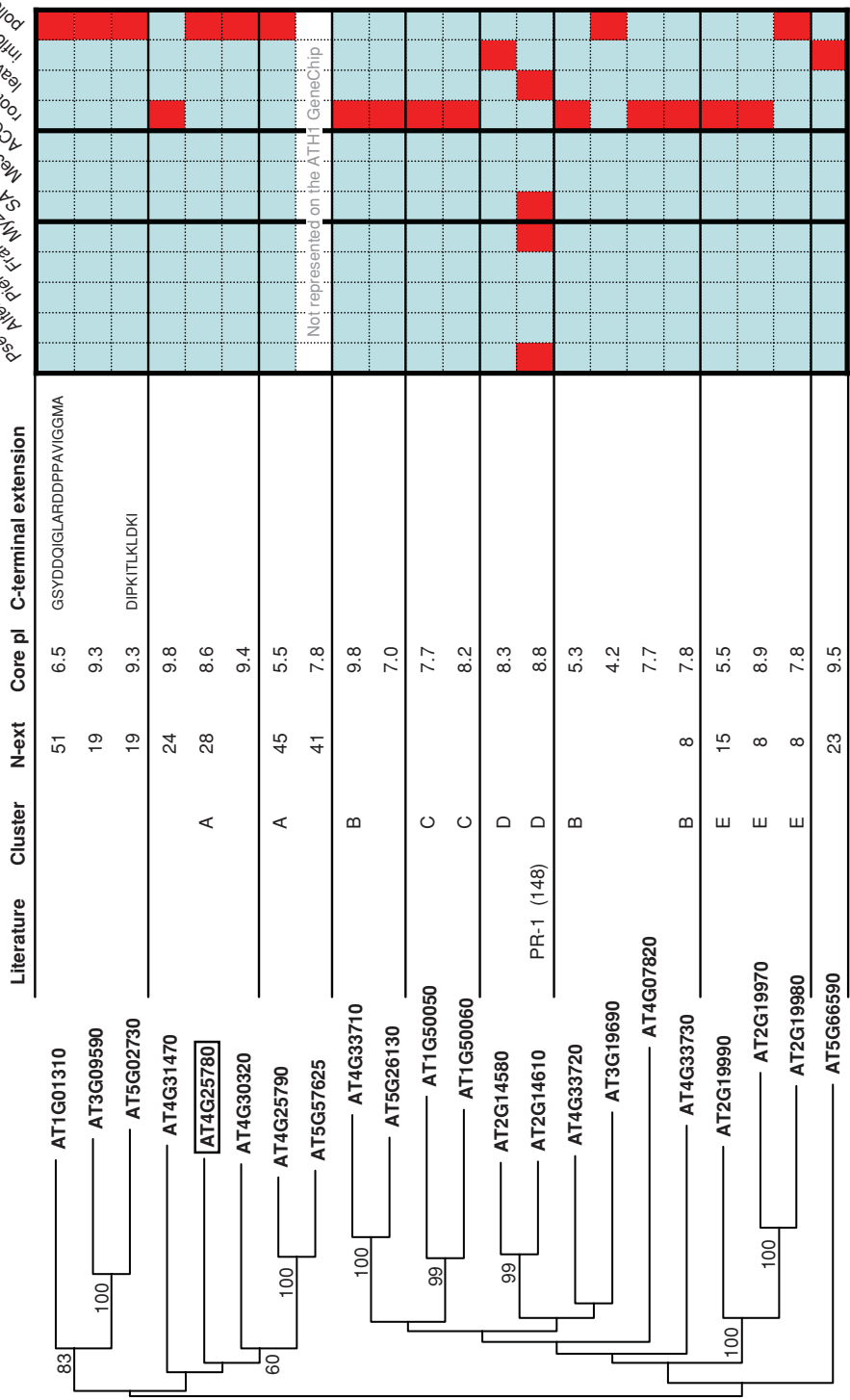
organ-specific manner in healthy plants (12, 162). Indeed, the same proteins can both appear during specific developmental stages and be induced in response to infection in the same organs (see **Figure 1**). For instance, pathogen-inducible basic PR-2 glucanase and PR-3 chitinase in tobacco are not detectable in young leaves of noninfected plants, but accumulate over their lifetime and are particularly abundant in roots. These and many other PRs are also present in floral organs of various plant species. PR-10-type proteins are widespread and have been found in pollen from various species. A PR-10-type protein from mung bean was reported to specifically bind cytokinin (43), whereas similar proteins from *Thalictrum flavum* and *Hypericum perforatum* function as metabolic enzymes in plant secondary metabolism (120). In addition, a general plant steroid carrier function, including brassinosteroid binding, has been suggested for PR-10-type proteins (88). A PR-2-type protein has been shown to be necessary for normal pollen development in tobacco and rice (170). Moreover, basic PR-2- and PR-

3-type proteins in tobacco, tomato, and pea seeds play a role in germination by degrading the cell walls of the seed coat and allowing the emerging radicle to protrude, or protect the exposed inner tissues of the seed against microbial entry (83, 93, 172). In carrot, PR-3- and PR-4-type chitinases are required for embryogenesis to proceed beyond the globular stage (76). A common occurrence of chitinases in embryogenic tissues has been associated with enzymatic activity on arabinogalactan proteins (e.g., 33, 105). These findings indicate that PR-type proteins can have a developmental role and, through their enzymatic activities, may generate signal molecules that could act as endogenous elicitors in morphogenesis. Such elicitors could also play a role in activating other types of defensive responses. Several PRs, such as PR-1-, -2-, -3-, and -4-type proteins and proteinase inhibitors, have been shown to be induced in abscission zones (114) and might be involved in cell wall loosening or in defense of the scarified tissue to invasion by bacterial and fungal pathogens. Many pathogen-inducible proteins in leaves

### Figure 1

Phylogenetic relationships and structural characteristics of predicted PR-1 proteins of *Arabidopsis thaliana*. All 22 proteins with a predicted PR-1 domain (Pfam code PF00188) encoded by the *Arabidopsis* genome were aligned with the PR-1-type protein of the fungus *Fusarium graminearum* (FG02744; not shown) as an outgroup for phylogenetic analysis. The gene model of AT1G50050 was adjusted (second intron removed) to improve alignment of the translation product to the other proteins. Additions to the conserved PR-1 domain at N termini and C termini were trimmed and gap-rich regions that could not be aligned unambiguously were removed. This alignment was used to construct a phylogenetic tree using neighbor-joining. Bootstrap percentages are provided only for branches receiving 60% or more support. Branch length reflects the extent of sequence divergence. The lengths of N-terminal extensions (N-ext) to the core sequence are the distance between the predicted signal peptide cleavage site (SignalP) and the first residue of the conserved core (a range is given if there is more than one potential signal peptide cleavage site). C-terminal extensions to the core are given beyond the conserved sequence "P[F/Y]". Isoelectric point (pI) is given for the conserved core sequences as calculated at [http://www.expasy.org/tools/pi\\_tool.html](http://www.expasy.org/tools/pi_tool.html). Gene clusters (i.e. homologous genes in close proximity on the same chromosome) are indicated with arbitrary letters. Boxed: the only protein with an ortholog (Os02g54560) in rice. On the right, an overview is given of the expression of the *Arabidopsis* PR-1 gene family from Affymetrix ATH1 GeneChip data. PR-1 gene expression was analyzed upon infection by the pathogenic bacterium *Pseudomonas syringae* pv. *tomato* and fungus *Alternaria brassicicola*, infestation by the herbivorous insects *Pieris rapae* (Cabbage white butterfly), *Frankliniella occidentalis* (Western flower thrips), or *Myzus persicae* (green peach aphid), exogenous application of salicylic acid (SA: 1 mM) or methyl jasmonate (MeJA: 100  $\mu$ M), or in different plant organs (roots, leaves, inflorescence, or pollen). Red squares indicate significant up-regulation. AT5G57625 is not represented on the ATH1 GeneChip. Gene expression data are derived from (32), and the *Arabidopsis* Microarray Database and Analysis Toolbox GENEVESTIGATOR (183).

*Pseudomonas syringae*  
*Alternaria brassicicola*  
*Plasmodium parasitica*  
*Myzus persicae*  
 SA  
 MeJA  
 AOC  
 roots  
 leaves  
 inflorescence  
 pollen



Affymetrix ATH1 GeneChip data

are constitutively present in storage tissues, such as fruits, seeds, and tubers (11, 131). This holds particularly for proteinase and amylase inhibitors, various types of lectins, defensins, thionins, and some lipid transfer proteins (12). Besides conferring protection against predation and disease (85), these proteins constitute a storage form of nitrogen (106) and might contribute to the survival of the organs during environmentally unfavorable periods.

### THE RELATIONSHIP BETWEEN PR-1-TYPE PROTEINS AND PLANT DEFENSE

The PR-1 family is strongly conserved and appears to be represented in every plant species investigated to date. Homologues have been found in fungi, insects, and vertebrates, including human, but of all PR-families, its function is the least understood (155). PR-1-type proteins are very similar in structure with 35% sequence identity among all PR-1 proteins and pair-wise sequence identities between 56% and 97%. All PR-1 proteins are structurally similar in having four  $\alpha$ -helices and four  $\beta$ -strands and share a number of strictly conserved residues, including six cysteines. These findings suggest that the unique molecular structure of PR-1 defines a protein module that has been retained during evolution and must serve one or more important functions. In tobacco (*Nicotiana tabacum* cv. Samsun NN) at least 16 PR-1-type genes appear to be present (24). Three acidic (1a, 1b, and 1c) and one basic (1g) protein with different biological properties were identified as being induced upon TMV infection. Additional or different homologues were found in other tobacco cultivars or in related *Nicotiana* species. In tomato, closely related homologues of the major extracellular, acidic tobacco PR-1a, -b, and -c are basic proteins (153). These findings indicate that PRs that are classified in the same family on the basis of sequence homology can have different properties and, hence, may differ substantially in biological activity.

In vitro, tomato PR-1c reduced germination of sporangia and germ-tube length of *P. infestans* and, in vivo, its application reduced the surface area of leaf discs infected with this oomycete. Basic tobacco PR-1g was similarly active against this pathogen, whereas tobacco PR-1a and -1b were only marginally so (96). However, neither in tomato nor in tobacco have transgenic plants engineered to constitutively express these proteins been described to possess enhanced resistance against *P. infestans*. This does not imply that these proteins could not contribute to basal resistance against this pathogen in vivo. However, other factors may already be sufficient to achieve basal resistance, or the pathogen is less vulnerable to these defenses in vivo. Extracellular defense-related proteins are perfectly located to contact invading attackers before tissue penetration has taken place, and have been considered a possible first line of defense. However, it takes time before the proteins start to accumulate. Thus, an effective pathogen is likely to have passed into further tissues before the induced proteins become sufficiently active. Consequently, their function may be directed more against following invaders, or constitute part of the biochemical barrier raised by SAR against subsequent infections.

Developmentally regulated defense-related proteins that are located in the vacuole could act as an effective second line of defense when the pathogen causes tissue damage. When the vacuole is disrupted, the lysosome-like contents are released and could engulf the pathogen. Many pathogens, including *P. infestans*, have a hemibiotrophic lifestyle in which, at first, they avoid damaging infected cells. Only later, when they have already colonized further tissues, the earlier infected cells collapse and massive liberation of hydrolytic enzymes occurs. Again, the plant would react too late to effectively stall the pathogen. Nevertheless, in the subtle but dynamic interplay between the pathogen and the plant, the balance may be shifted depending on the speed and magnitude of attack and defense.

When pathogen-inducible PR-1 genes in tobacco were expressed individually under the control of the constitutive 35S promoter, the transformed plants were slightly more resistant to blue mold, caused by *Peronospora tabacina*, and black shank, due to *Phytophthora parasitica* f.sp. *nicotianae*, but not to diseases caused by the fungus *Cercospora nicotianae*, the bacterium *Pseudomonas syringae* pv. *tabaci*, or several viruses (12). Overexpression of a TMV-inducible basic PR-1-encoding gene from pepper enhanced tolerance to *P. parasitica* var. *nicotianae*, *Ralstonia solanacearum*, and *P. syringae* pv. *tabaci* (121). The apparent association between PR-1 proteins and enhanced resistance against oomycetes has been noted, but too few data on nonoomycete pathogens have been reported to conclude that PR-1 is directed specifically against oomycete attack.

Such functional analysis would be greatly aided by mutant or knock-out lines in which all members of the PR-1 family were lacking or nonfunctional. Transient silencing of *PR-1* expression by double-stranded RNA interference in barley allowed the mildew fungus *Blumeria graminis* f.sp. *bordei* to penetrate the cell wall more frequently (126). However, the effect was small. Given the number of PR-1 proteins in tobacco and tomato and the lack of knowledge about the number of possible genes, this has not been attempted in other species. Although known PR-1 proteins contain a widely conserved sequence encompassing the fourth  $\alpha$ -helix and first part of the third  $\beta$ -strand, CGHYTQVVW[R/K]X[S/T][V/T][R/S]XGC (155), there is insufficient homology at the nucleotide level to allow down-regulation of all genes through RNA interference or virus-induced gene silencing.

In the fully sequenced genomes of *Arabidopsis* and rice, 22 and 39 PR-1-type genes are present, respectively (**Figures 1, 2**). Most rice PR-1 genes are present in clusters (**Figure 2**). A special case is a cluster on chromosome 7 with 14 PR-1 genes. This cluster contains one full and one partial duplication of a  $\sim$ 32-kb genomic region containing three PR-1 genes, as well as many transposons. As

a result, these three PR-1 genes have two or four exact copies in this cluster. Most of the remaining PR-1 genes in rice and about half of those in *Arabidopsis* are present in clusters of two to four genes, with at most four intervening genes. In several cases, but not always, clusters contain genes that are more related to each other than to the other paralogs in the genome, indicative of local duplication events.

In phylogenetic trees containing homologous proteins from both *Arabidopsis* and rice, only one branch was found with proteins from both species. These two, At4g25780 and Os02g54560, are likely to be orthologs. Next to relatively high similarity of their core sequences, additional evidence for orthology is that both are predicted to contain N-terminal extensions of similar length (29 and 36 residues, respectively) that contain cysteines, a feature that is not observed for any other PR-1 protein of either plant species. Paucity of mixed branches of PR-1 proteins indicates that the current complement of PR-1 proteins of *Arabidopsis* and rice has developed mostly independently, through loss of orthologs and gene duplications after divergence of the species. How far this can have a bearing on the antimicrobial activities of these genes is unclear. In *Arabidopsis* only a single PR-1 gene (At2g14610) is activated by infection, insect attack, or chemical treatment, whereas ten and eight different PR-1-type genes are constitutively expressed in roots and pollen, respectively (**Figure 1**). In rice, JA-inducible cDNAs corresponding to one acidic and one basic PR-1 protein have been characterized (2), but induction of the proteins themselves upon infection has not been investigated.

The retention of the high numbers of homologues can only be explained by an important function in plant life. Recently, a 28-kDa PR-1 family member, Tex 31, from the venom duct of the cone snail *Conus textile* was shown to have serine protease inhibitor-sensitive proteolytic activity, with the likely catalytic residues falling within the structurally conserved PR-1 domain (92). Similarly,

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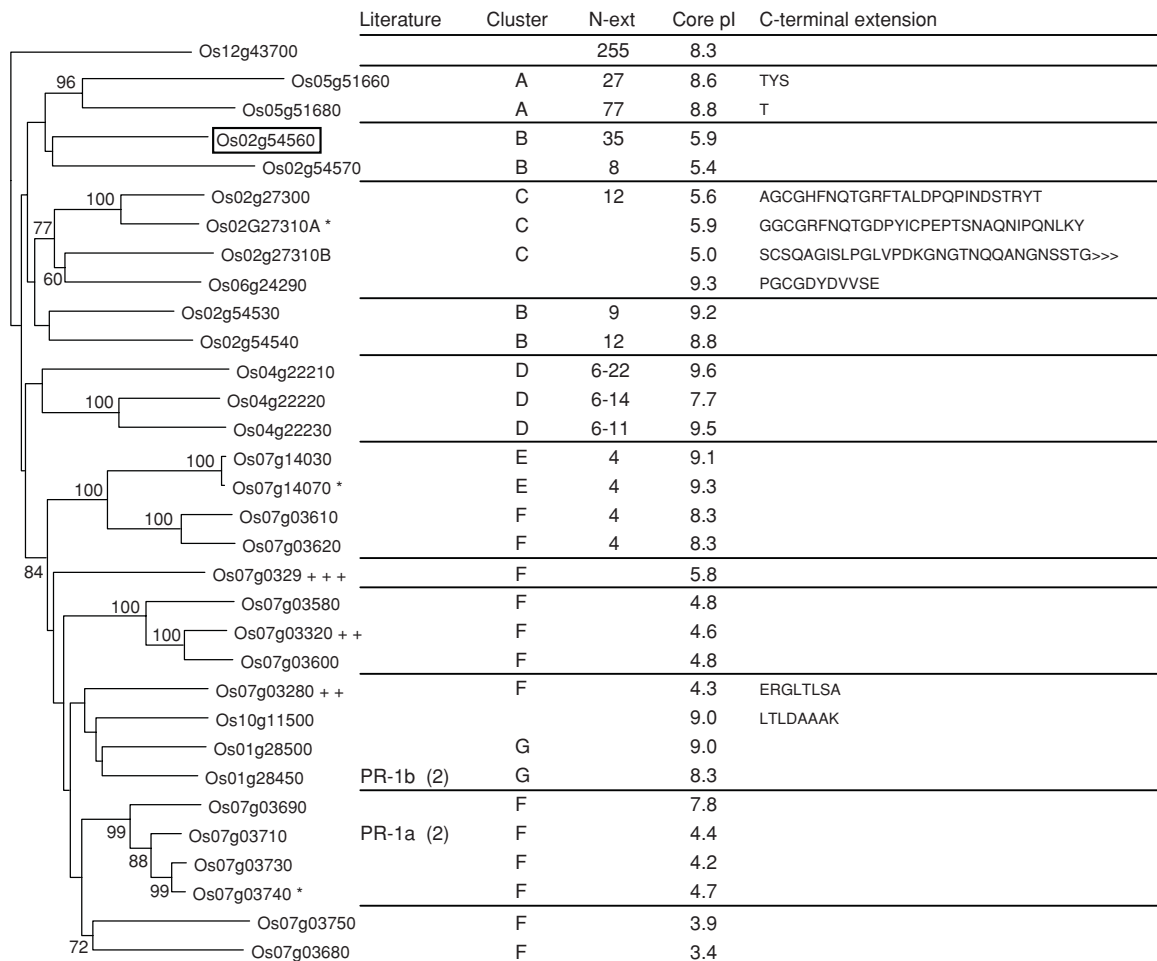
**Paralogs:**

homologous genes within a species

**Orthologs:**

related genes in different species

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**Figure 2**

Phylogenetic relationships and structural characteristics of predicted PR-1 proteins of rice. All 32 proteins with a predicted PR-1 domain encoded by the rice genome were aligned with *F. graminearum* FG02744 (not shown) as an outgroup for phylogenetic analysis. Three proteins are encoded by three or four identical genes, of which only one is shown in the figure (“+” signs indicate the presence of extra copies; duplicates of Os07g03280 are Os07g03370 and Os07g03460; duplicates of Os07g03320 are Os07g03410 and Os07g03500; duplicates of Os07g03290 are Os07g03380, Os07g03470 and Os07g03590). The following gene models were adjusted to optimize alignments and/or to comply with cDNA sequences: Os07g03740 (first 5 exons removed, this is now a pseudogene due to a frameshift); Os07g14070 (disrupted by insertion of a repetitive element instead of the currently proposed intron); Os04g22210 (intron removed, different start codon); Os02g27310 (split in two PR-1 genes, provisionally called Os02g27310A and Os02g27310B). Pseudogenes are indicated with an asterisk. The C-terminal extension of Os02g27310B is much larger than shown (hence the “>>>” signs) and includes a probable transmembrane segment and protein kinase domain. Boxed: the only protein with an ortholog (At4G25780) in *Arabidopsis thaliana*. See legend of **Figure 1** for details of the phylogenetic analysis and other characteristics of the proteins listed.

dimerization of the human Golgi-associated PR-1-type protein GAPR-1 may lead to the formation of a catalytic triad similar to that of serine proteases, across the dimer interface by residues from both molecules (129). However, others (70) consider the PR-1 fold to form a stable scaffold for biological interactions with other proteins. If such interactions occur in infected plants, they are not stable, because no complexes of PR-1 with other proteins have been identified upon native polyacrylamide gel electrophoresis (152).

### ACTIVITIES AND BIOLOGICAL SIGNIFICANCE OF OTHER INDUCIBLE DEFENSE-RELATED PROTEINS

In spite of the consistent association of inducible defense-related proteins with resistance reactions, most published data on constitutively expressing transgenic plants show only limited, if any, enhanced resistance against specific pathogens. Whereas many defense-related proteins have been demonstrated to inhibit growth of selected fungi and bacteria *in vitro*, in most cases such results are not predictive of suppression of pathogens in planta. This may be caused by insufficient expression or instability of the protein in the transgenic plant. Another reason may be insensitivity of the pathogen toward the protein due to secretion of inhibitors (145) or the presence of protective layers around hyphal or bacterial cell walls (125) that are not present during *in vitro* cultivation on artificial media. However, in some cases, expression of the transgenic protein may be more effective than in *in vitro* tests, for instance when elicitors that are released from the pathogen activate defense responses in the plant (10, 177). Both glucanases and chitinases could also act on endogenous plant substrates and, thereby, aid in the generation of signal molecules that may function as endogenous elicitors of further defensive mechanisms. Thus, *in vitro* only indications can be obtained of the range of fungi or bacteria that might be affected by a defense-

related protein when this is expressed in a transgenic plant.

Furthermore, the limitation of most tests to easily transformable plant species, such as tobacco, and their cognate pathogens, makes it difficult to generalize an effect observed.

Several types (classes) and isoforms of  $\beta$ -1,3-glucanases and chitinases with different substrate specificities and specific activities (13, 132) are constitutively present in plants. As a result, they can only be enhanced, rather than specifically induced, by pathogen infection, and often are increased also under other stress conditions. Although these enzymes can potentially degrade microbial cell wall components and, thereby, might contribute to the expression of resistance, causal relationships have been established only for a relatively small number of well-characterized PR-type proteins (12, 50). In tobacco, specific  $\beta$ -1,3-glucanases from alfalfa, barley, tobacco, and soybean have been shown to suppress diseases caused by *C. nicotianae* and *Phytophthora megasperma* f.sp. *medicaginis*, *Rhizoctonia solani*, *Alternaria alternata* and *P. parasitica* f.sp. *nicotianae*, and *P. tabacina* and *P. infestans*, respectively. Tobacco PR-2b did not reduce infection by viruses. In fact, transformants of tobacco and *Nicotiana sylvestris* expressing an antisense construct of a class I  $\beta$ -1,3-glucanase from the latter species (homologous to tobacco PR-2d) became less diseased than nontransformed control plants. This result is interpreted as being due to enhanced callose accumulation at the plasmodesmata, leading to more effective blockage of virus transport from cell to cell (15, 42). Overexpression of glucanases from soybean has been demonstrated to enhance protection of potato to *P. infestans* and kiwi to *B. cinerea* (50), whereas a glucanase from potato was reported to increase resistance in flax against *Fusarium oxysporum* and *Fusarium culmorum* (171).

*R. solani* was the first fungus shown to be suppressed in transgenic tobacco and canola overexpressing a basic PR-3-type chitinase from bean (12, 50). However, transgenic cucumber plants were not protected against

**$\beta$ -1,3-glucanases:**

enzymes that hydrolyze  $\beta$ -1,3 glycosidic bonds in linear or branched glucans

**Permatins:** proteins that permeabilize microbial membranes

**Osmotin:** a basic PR-5 protein that is induced by osmotic stress

**Phosphomannans:** phosphorylated mannose polymers

**AMP:** adenosine monophosphate

this fungus when transformed with the same type of chitinases from bean, petunia, or tobacco, whereas tobacco plants transformed with chitinases from barley, cucumber, or tobacco were. The tobacco chitinase counteracted *R. solani* also in *N. sylvestris* and carrot, as well as *Cercospora arachidicola* in peanut, but when overexpressed in tobacco was not effective against *C. nicotianae* (116). Chitinases from rice were effective against *R. solani* and *Magnaporthe grisea* in rice (28), against *Uncinula necator* in grapevine (175), against *B. cinerea* in cucumber (71, 136) and chrysanthemum (138), and in Italian ryegrass against *Puccinia coronata* (137), but not in alfalfa against *Stemphylium alfalfae*, *Colletotrichum trifolii*, *Phoma medicaginis*, and *P. megasperma* f.sp. *medicaginis*. The enzymes from bean and tobacco did not counteract *Alternaria cucumerina*, *B. cinerea*, or *Colletotrichum lagenarium* on cucumber. The tobacco enzyme was effective in carrot against *B. cinerea* and *Sclerotium rolfsii*, but not against *Alternaria radicina* or *Thielaviopsis basicola*, whereas a similar chitinase from petunia had no effect in either carrot or cucumber. A tomato chitinase conferred protection to oilseed rape against *Cylindrosporium concentricum*, *Phoma lingam*, and *Sclerotinia sclerotiorum* (12, 50). Finally, a chitinase from *Lycopersicon chilense* increased resistance of tomato to *Verticillium dahliae* (135).

Expression of PR-8-type chitinases from tobacco, cucumber, and sugar beet in tobacco resulted in enhanced resistance against *R. solani*. The sugar beet enzyme also protected tobacco against *C. nicotianae* and birch against natural infection by *Melampsorium betulinum* (birch rust) but not *Pyrenopeziza betulicola* (leaf spot) (103). These results indicate that effectiveness is dependent on transgene source, plant species, and pathogen sensitivity. The latter is further illustrated by observations that combinations of glucanases and chitinases can be substantially more effective in degrading fungal cell walls than each alone (12, 50, 90, 176). Notably, whereas neither glucanases nor chitinases alone were effective, simultaneous expression of tobacco PR-

2e and PR-3d rendered tomato resistant to *Fusarium oxysporum* f.sp. *lycopersici* and carrot to *Alternaria dauci*, *A. radicina*, *Cercospora carotae*, and *Erysiphe heraclei* (90).

The thaumatin-like PR-5 proteins belong to a larger family of proteins that includes permatins from monocot grains and can permeabilize fungal membranes (4). In *Arabidopsis* 24 PR-5-type genes have been annotated (82). The basic tobacco PR-5c (osmotin), which is inducible by pathogens and osmotic stress, and its homologs in tomato and potato have in vitro antioomycete activity against *P. infestans*, and transgenic tobacco and potato plants have enhanced resistance against this pathogen but not against *P. parasitica* f.sp. *nicotianae*. Overexpression of PR-5 genes from rice has been demonstrated to reduce infection of rice by *R. solani* (50), of wheat by *Fusarium graminearum* (20), of tobacco by *A. alternata* (158), and of carrot by *A. dauci*, *Alternaria petroselini*, *A. radicina*, *B. cinerea*, *R. solani*, and *S. sclerotiorum* (109). Osmotin from tomato protected transgenic orange plants against *Phytophthora citrophthora* (39).

A PR-5-type protein from corn seeds, zeamatin, has been described to inhibit mammalian trypsin and insect  $\alpha$ -amylase (123). However, several other thaumatin-like proteins lack these activities and the high molar ratio of zeamatin required for inhibition suggests either nonspecific complex formation or the presence of an impurity. Zeamatin, as well as chemically induced extracellular PR-5-type proteins from barley and pea leaves, bound polymeric  $\beta$ -1,3-glucans, whereas tobacco osmotin did not (147). Osmotin, but not zeamatin, exhibited endo- $\beta$ -1,3-glucanase activity on these substrates. Analysis of other thaumatin-like proteins confirmed a general lack of correlation between antifungal activity,  $\beta$ -1,3-glucan binding, and  $\beta$ -1,3-glucan hydrolysis (47, 91). Thus, the main antifungal action of PR-5-type proteins must reside in a different property.

An osmotin-like protein as well as a basic chitinase in suspension-cultured potato cells were found to bind actin (140), and similar

proteins from tobacco callus were implicated in binding cytokinin (74). Osmotin induces apoptosis in the yeast *Saccharomyces cerevisiae*, apparently by binding to phosphomannans in the cell wall (63), which facilitates access to a 7-transmembrane-domain receptor-like protein in the plasma membrane that regulates lipid and phosphate metabolism and is homologous to a mammalian receptor for the hormone adiponectin. Like adiponectin, osmotin activates AMP kinase in murine myocytes via adiponectin receptors, suggesting that osmotin action is receptor-mediated (94). Overexpression of the stress-related yeast PIR2 cell wall glycoprotein protected *S. cerevisiae* from the toxic action of osmotin (180), and a similar result was obtained for *Fusarium oxysporum* f.sp. *nicotianae*, allowing increased disease severity and fungal growth in tobacco seedlings (95). Thus, osmotin seems to contribute to basal resistance of tobacco against *F. oxysporum* f.sp. *nicotianae*. Other PR-5 proteins are much less active against yeast cells than osmotin but active against other fungal species (180). It seems that cell wall binding facilitates the action of osmotins and contributes to their fungal target specificity.

Four PR-5 proteins from *Arabidopsis* and seven from rice have extensions that end in a hydrophobic stretch that could serve as a membrane anchor (M. Rep, unpublished results), suggestive of a (temporary) attachment of these proteins to a membrane. Others contain a potential transmembrane segment followed by a kinase domain, suggestive of a function in extracellular sensing of perhaps  $\beta$ -1,3-glucan fragments and signal transduction. Three of these receptor-like kinases are present in *Arabidopsis* and two in rice. One of the PR-5-like receptor kinases from *Arabidopsis* is expressed constitutively at low levels in leaves and siliques and at higher levels in flower stems and roots, and has been suggested to recognize the same targets as the related PR-5 proteins (166). When expressed in creeping bentgrass, it delayed dollar spot symptoms caused by *Sclerotinia homoeocarpa* (52), indicating that it has antifungal

activity. One PR-1 protein from rice also appears to be connected to a transmembrane region followed by a kinase domain (Figure 2). A similar fusion has been described for a pathogen-inducible tobacco PR-3-type protein (CHRK1). This receptor-like kinase may bind chitin but is devoid of chitinase activity due to an amino acid change at the active site (69). CHRK1 has been shown to interact with NtPUB4, an armadillo repeat protein in tobacco (68), and appears to be involved in a developmental signaling pathway regulating cell proliferation, differentiation, and endogenous cytokinin levels (81). These findings open up the possibility that some defense-related proteins have a role in signaling in response to pathogen attack or perhaps in developmental regulation.

Similar results have been obtained for other inducible defense-related genes, including SAR 8.2 (12, 50, 90, 176) and DDR206 (167, 168). In addition, genes encoding seed-specific antimicrobial proteins or glucanases and chitinases of microbial origin have been analyzed in vitro, as well as in transgenic plants (122). The results obtained corroborate the conclusion that resistance can be enhanced against some pathogens in some plant species. Typically, disease development is slowed, or pathogen proliferation and symptom severity are reduced, but not prevented. Given the enhanced protection afforded by combinations of defense-related genes, it could be that several proteins need to act in concert for effective resistance to be manifested.

## SIGNALING IN THE INDUCTION OF DEFENSE-RELATED PROTEINS

Upon infection with various types of pathogens, defense-related genes are coordinately activated and may be expressed in both infected and noninfected tissues concomitant with the development of SAR (118). The association between accumulation of PRs, the products of the SAR genes, and SAR is often taken to represent a causal relationship, with

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**Armadillo repeat:** a 42-amino acid "arm" motif repeat first identified in the *Drosophila* segment polarity gene product armadillo ( $\beta$ -catenin)

**Induced systemic resistance:** the phenomenon that plants acquire an enhanced defensive capacity against subsequent pathogen attack as a result of root colonization by selected strains of nonpathogenic bacteria

**ISR:** induced systemic resistance

**AOS:** active oxygen species

**Lesion-mimic mutants:** plants that "spontaneously" develop necrotic lesions during development or in response to variations in environmental conditions

**Hevein:** a chitin-binding protein from rubber latex

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the proteins acting as the agents responsible for the induced resistance against subsequent infection by a wide range of pathogens. However, the limited effectiveness of the proteins in transgenic plants, as discussed above, and the results that no plant engineered to constitutively express one or more defense-related genes has been shown to be more resistant against viruses, are difficult to reconcile with the generally enhanced defensive capacity of SAR-expressing plants. Moreover, pathogens such as *B. cinerea* on tobacco and *A. brassicicola* on *Arabidopsis* are virtually insensitive to SAR, but restricted by a different mechanism of induced resistance that is independent of the presence of inducible defense-related proteins in protected tissues (143, 146). This type of enhanced defensive capacity is elicited by specific strains of nonpathogenic, root-colonizing bacteria and has been termed induced systemic resistance (ISR) (154). Like SAR, ISR is active against a broad spectrum of pathogenic fungi and bacteria but, unlike SAR, not against viruses. ISR has been studied mainly in *Arabidopsis* and found not to be associated with the expression of defense-related genes (160). In the case of challenge inoculation of induced *Arabidopsis* plants with the leaf pathogenic bacterium *P. syringae* pv. *tomato* DC3000, the protection afforded by SAR usually is slightly stronger than that by ISR. Many of the pathogens investigated can be restricted by both SAR and ISR and induction of both SAR and ISR in the same plant leads to additively increased protection (157). These observations indicate that SAR and ISR are complementary types of induced resistance with partly overlapping, partly specific actions against different types of pathogens.

Many conditions have been described to induce SAR as well as defense-related proteins (150). The expression of a PR-1 gene or protein in particular is usually taken as a molecular marker to indicate that SAR was induced. All PR-1 genes in plants appear to be inducible by SA, and endogenous production or exogenous application of SA has been shown to be both necessary and sufficient to elicit the in-

duced state (161). Pathogen-induced synthesis of SA in tobacco is considered to occur from benzoate, whereas the evidence in *Arabidopsis* points to isochorismate as the immediate precursor (35). How synthesis of SA in infected plants is activated and regulated is not known, nor is it clear which factor(s) act as elicitors of SA production during pathogen infection. It has been suggested that active oxygen species (AOS) are involved and, indeed, tissue damage invariably leads to the production of AOS and synthesis of SA. Although SA can be transported in the plant, reciprocal graftings of transgenic NahG plants, in which SA is degraded, and nontransformed plants as rootstocks or scions, demonstrated that SA is not the translocated signal in SAR (161). Similar graftings between transgenic ethylene-insensitive tobacco plants expressing a mutant ethylene receptor gene from *Arabidopsis* as rootstock and nontransformed control plants as scion showed little or no SAR induction in the scion, indicating that ethylene perception is necessary for the generation, release, or transport of the mobile signal to distant tissues. Upon arrival of the mobile signal, the latter tissues must start producing SA, which induces the defense-related proteins locally (159). The nature of the mobile signal has remained elusive so far. An *Arabidopsis* mutant, *dir1*, impaired specifically in the systemic character of SAR, implicates involvement of a lipid transfer protein (86), suggesting that the mobile signal may contain a lipid moiety.

SA production has been suggested to be part of a feed-forward loop (130). Progressive damage will amplify SA production even further, and inevitably lead to induction of defense-related genes and SAR. This is precisely what happens when cells start necrotizing, such as during a hypersensitive reaction, and explains why slowly developing and expanding necrotic lesions or spots lead to such strong expression of defense-related genes and SAR. This also explains why lesion-mimic mutants that are affected in very different genes but all exhibit necrotic leaves at

some stage of development or under some environmental conditions, express inducible defense-related genes and SAR constitutively. Typically, many *cpr* (constitutively expressing PRs) mutants are lesion mimics (22) and are likely to express marker PRs and SAR because of this phenotype rather than because of a mutation in a specific step in the defense signaling pathway.

Upon hypersensitive necrosis, not only does the level of SA increase, but also JA synthesis and ET production are strongly enhanced early on (e.g., 108, 128). As a result, in addition to SA-inducible defense-related genes, such as in *Arabidopsis* PR-1, -2, and -5, JA- and ET-inducible genes, i.e., PR-3-type basic chitinase, PR-4-type hevein-like protein, and PR-12 defensin PDF1.2, become activated (143). PDF1.2 in particular is often used as a marker for the induction of the JA- and ET-dependent defense-signaling pathway (80). Induction of PDF1.2 can be limited, however, because accumulation of SA inhibits JA synthesis and action (134). ET sensitizes the tissue to respond to SA, as evidenced by a lowering of the concentration of SA that is required for PR-1 expression when *Arabidopsis* is exposed to ET (79). On the other hand, in tobacco induction of PR-1a by SA was reduced by simultaneous application of JA (97). The nature and extent of the cross-talk between the three defense-regulating hormones depend on the timing and magnitude of their increases, which, in turn, can be modulated through the action of the attacking pathogen (32).

Biotrophic pathogens are dependent on live tissues and avoid triggering necrosis. The enhanced disease susceptibility to biotrophic pathogens, such as the oomycete *H. parasitica*, of *Arabidopsis* mutants that are impaired in SA synthesis or signaling indicates that SA-dependent defenses contribute to basal resistance against these types of pathogens (143). Exogenous application of SA leads to induction of PR-1, -2, and -5 mRNAs. Thus, SA-regulated defense-related proteins may be directed primarily against pathogens with a

biotrophic lifestyle (those forming haustoria) rather than oomycetes as such (100). Little information on the effect of these proteins on biotrophs other than oomycetes is available and further clarification is needed.

*Arabidopsis* plants impaired in JA or ET signaling are, in general, more susceptible to necrotrophic pathogens (45, 143, 164). SA may induce resistance against these pathogens also, but it is likely that SA-regulated defenses are important only at a stage in which the pathogen (still) behaves as a hemibiotroph. *B. cinerea*, a pathogen that is completely dependent on its necrotrophic lifestyle, kills the tissue in advance of tissue colonization, is insensitive to SA-regulated defenses—at least in *Arabidopsis* and tobacco—and is not affected by SAR. In contrast, necrosis in *Arabidopsis* as a result of infection by *P. syringae* pv. *tomato* follows a phase of spreading chlorosis, in which the bacterium multiplies abundantly in the infected leaves. Thus, *P. syringae* pv. *tomato* has a mixed biotrophic/necrotrophic lifestyle, and both SA- and JA/ET-regulated defenses contribute to basal resistance against this pathogen (157). JA/ET-dependent defenses in *Arabidopsis* are boosted upon challenge inoculation of plants expressing rhizobacteria-mediated ISR and are most effective against pathogenic bacteria and fungi with mixed biotrophic/necrotrophic and necrotrophic lifestyles (146, 154).

Many PR-type proteins are JA- and/or ET-inducible (150), and their occurrence can be further modulated by abscisic acid (7, 89, 113, 181). Whereas in *Arabidopsis* a distinction between the SA-inducible PR-1, -2, and -5, and the JA/ET-inducible PR-3, -4, and -12 seems clear (143), in tobacco it has been demonstrated that different members within the same protein family are differentially regulated by SA and JA/ET (97, 128). Thus, the acidic PR-1, -2, -3, and -5 proteins, which are inducible by TMV and accumulate in the apoplast, are regulated primarily by SA, with ET and/or JA acting sometimes in a synergistic manner. The basic isoforms that are developmentally regulated and are present in the

vacuole appear to be regulated and are inducible further by JA and ET, acting alone or in concert. For instance, ET-insensitive tobacco does not express basic PR-1g, -2d, and -5c in response to TMV infection, whereas local expression of the acidic isoforms is not affected (159). In other plant species, similar differential induction has been noted but not systematically investigated. In tomato, the basic orthologues of tobacco PR-1a, -1b, and -1c are present in the apoplast and inducible by SA, but in *Arabidopsis* and rice, the specific induction characteristics of the many PR-1-type proteins have not been investigated.

### RELEVANCE OF DEFENSE-RELATED PROTEINS IN SITUATIONS OTHER THAN PATHOGEN INFECTION

Secreted PRs accumulate in senescing leaves of some species when yellowing is becoming apparent and in ripening fruits (14, 30, 110), as well as in the medium of cell suspension cultures in the absence of visible necrosis (12, 139), indicating that at least some inducible defense-related proteins are produced also under specific physiological conditions. Abiotic stresses can also elicit defense-related protein induction, as in the case of osmotic stress, cold stress, or wounding (12). Typically, tobacco osmotin is induced in leaves, stems and roots by drought, high salt, or abscisic acid, as well as in leaves by wounding or UV light. Several reports show that apoplastic PR-1-type proteins, chitinases, glucanases, thaumatin-like proteins, thionins, and lipid-transfer proteins are induced during cold stress, as exemplified by results of Hon et al. (60) on cold hardening in rye showing that the accumulating PR-2, -3, and -5 proteins have antifreeze activity. The same proteins accumulated in response to cold, short daylength, and dehydration. The chitinase- and glucanase-like proteins have both enzymatic and antifreeze activities (6, 49). At freezing temperatures the proteins alter ice crystal shape and reduce freezing injury by

slowing the growth and recrystallization of ice (48). Similar proteins are induced under the same conditions in freezing-tolerant wheat and barley, but not in freezing-sensitive maize (3). In winter rye, similar PRs were induced by cold, by treatment with SA, and upon infection with the snow mold fungus *Microdochium nivale*, but only the PRs induced by cold exhibited antifreeze activity (56). Those proteins appear to be induced through the action of ethylene (178), indicating a striking parallel with the differential induction of specific PRs in other plant species.

Other abiotic stresses, such as heavy metal toxicity, are likely to induce defense-related proteins as a result of the cell-damaging action of the stimulus. In fact, induction of PRs is a common phenomenon in plants treated with millimolar concentrations of various types of chemicals that affect cell metabolism. Such results make it difficult to interpret whether induction was due to a toxic, pharmacological, or physiological effect. Even SA is toxic to plants at concentrations that are less than a factor two higher than commonly used to induce SAR. Such toxicity can be easily monitored, as stress-induced JA, ET, and enzymes such as peroxidase are increased under those conditions, in contrast to the specific induction of only some PRs by SA alone (150).

When testing for the compounds in various pollen and latex that are responsible for allergic reactions in humans, several types of PR-like proteins were found to be responsible. The major pollen allergen from birch, *Bet v 1* (16), belongs to the PR-10 family, is present in "orbicules," protrusions on the sporopollenin coat, and is induced by both biotic and abiotic stress conditions in various plant tissues. PR-10-type proteins are widespread in plants and PR-10-type allergens are present in many foods such as fruits and vegetables. Other defense-related food allergens belong to the PR-2, -3, -4, -5, -8, -12, -14, and -15 families (58, 173). Rubber latex contains several PR-type protein allergens that may serve a protective function. The main latex allergens are *Hev b 2*, a

glycosylated  $\beta$ -1,3-glucanase containing vacuolar targeting sequences, and a PR-3-type chitinase (174). The identification of several PR-like defense-related proteins as allergens in plant products is a cause of concern (59) and reduces the likelihood that transgenic plants with enhanced resistance as a result of PR gene expression will be commercially acceptable.

## CONCLUSIONS AND PERSPECTIVES

Genes encoding inducible plant defense-related proteins, particularly PRs, comprise broad, evolutionarily conserved families with individual members differing widely in occurrence and, where known, activity. Therefore, they likely have an ancient origin with subsequent diversification to serve different functions. Those proteins that are expressed during plant development in specific stages or organs may, through their specific hydrolytic activities, contribute to the generation of signal molecules that can act as morphogenetic factors, such as chitinase in somatic embryogenesis. Some may themselves act as ligands of specific receptors and have a signaling role, as deduced from the occurrence of genes encoding contiguous PR-type-transmembrane-protein kinase domains. Other PR-like proteins can serve a role in protection of cellular structures against abiotic stress, such as the PR-type proteins with antifreeze activity. Whether these are original functions from which an association with defense against pathogens was derived, or the reverse, is unclear. However, their widespread induction upon pathogen attack and their regulation by the defense regulatory hormones SA, JA, and ET suggest that they play an important role in alleviating the effects of attack by pathogens and insects, as well as some forms of abiotic stress. In several instances, quantitative resistance against pathogens has been shown to be associated with constitutively expressed PRs (84, 107, 165). Adult plants growing in the field without any signs of disease often contain readily de-

tectable levels of PRs (L.C. van Loon, unpublished observations), suggesting that they have experienced stress conditions. There are several reports that apparently nonpathogenic microorganisms present on leaves or in soil can increase activities of, for example, chitinase, glucanase, PAL, peroxidase, and/or polyphenoloxidase systemically in plants (151). Therefore, plants seem to constantly adjust their defensive status to the dynamically changing environment.

Because only some members of the families of defense-related proteins have a suppressive effect on some pathogens but not others, their role in restricting pathogen growth and tissue colonization appears limited. However, they appear to be part of a far larger array of SA-, JA-, and ET-dependent defenses in which each component could contribute more or less to basal resistance against an attacker, as well as to the enhanced resistance in plants with induced resistance. In SAR, the presence of induced PR-type proteins is likely to contribute to some extent to the enhanced defensive capacity. In contrast, in ISR, no defense-related proteins are present in induced leaves before challenge, but upon infection activation of JA-responsive genes in particular is accelerated and enhanced, a phenomenon known as priming (23).

Plants are able to recognize microbial invaders through specific surface determinants, collectively called pathogen-associated molecular patterns (PAMPs) and to react through defense signaling cascades (5, 99). Although a causal connection between recognition of nonpathogenic microorganisms and specific inducible defense-related proteins has not been established so far, broad-spectrum effectiveness of the induced resistance responses suggests that PR-type and similar proteins are part of an immune surveillance mechanism that protects the plant primarily against invasion by microorganisms that are generally perceived as nonpathogenic. Many saprophytic fungi appear more sensitive to the action of lytic enzymes, such as glucanases and chitinases, than are pathogens that are adapted

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### PAMPs:

pathogen-associated molecular patterns

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to attack living plants (124). Indeed, certain pathogens have been shown to be insensitive to the action of defense-related proteins from their host. For instance, *Cladosporium fulvum* is not sensitive to the chitinase and  $\beta$ -1,3-glucanase of its host, tomato (65). *Phytophthora sojae* specifically inhibits the glucanase activity of its host, soybean, by producing inhibitor proteins (117). *Fusarium solani* f.sp. *eu-martii* degrades PRs in the intercellular fluid from potato (101). Similarly, constitutive expression of various PRs in tobacco did not affect colonization by the beneficial mycorrhizal fungus *Glomus mosseae* (163).

Mutant or transgenic plants with reduced sensitivity to either JA or ET become sponta-

neously infected by normally nonpathogenic fungi when grown in commercial potting soil (73, 164). A surveillance system based on antimicrobial proteins appears very similar to the innate immunity of vertebrates and insects, in which there are signaling cascades comprising components very similar to those involved in plant resistance signaling (99). Although the details of these systems and the nature of the antimicrobial polypeptides differ substantially between animals and plants, there appears to be a basic similarity in the innate immune responses that allow plants to flourish and us as human beings to exploit and enjoy the world of plants around us.

#### SUMMARY POINTS

1. Microarray analyses have shown that in both compatible and incompatible plant-pathogen interactions, hundreds of genes are up- or downregulated, but only a limited number of inducible defense-related proteins have been characterized.
2. Pathogenesis-related (PR) and similar proteins have been found to be inducible by infection with various types of pathogens in many plant families and have been classified into 17 families.
3. Upon pathogen or insect attack, many defense-related proteins are inducible by the signaling compounds salicylic acid, jasmonic acid, or ethylene, whereas other plant hormones, such as abscisic acid, can modulate expression.
4. Accumulation of PRs is a hallmark of pathogen-induced systemic acquired resistance, but no defense-related proteins are detectable in plants with rhizobacteria-induced systemic resistance, in spite of a similar enhanced defensive capacity of the plants. However, in both cases, upon challenge inoculation with a pathogen, defense-related genes are expressed faster and to higher levels, a phenomenon called priming.
5. Most inducible defense-related proteins possess antimicrobial activity against fungi and bacteria *in vitro*, but when expressed in transgenic plants reduce only a limited number of diseases, depending on the nature of the protein, plant species, and pathogen involved.
6. *Arabidopsis* and rice contain 22 and 39 PR-1-type genes, but only 1 and 2, respectively, have been found to be inducible by pathogens or insect attack. Many other PR-1 genes are expressed constitutively in roots or floral tissues, indicative of a role in plant development.
7. Defense-related proteins can be induced also by wounding and cold treatment. Some PRs can modulate ice crystal formation and have antifreeze activity.

8. Many constitutively expressed PRs in fruits, vegetables, and pollen can act as allergens, raising questions as to the possible impact of transgenic plants expressing antimicrobial proteins for enhanced disease resistance.

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23. Within this short review it is outlined that JA- and ET-dependent priming of defense-related gene expression rather than accumulation of defense-related proteins is characteristic of rhizobacteria-induced systemic resistance.

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## ERRATA

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