Induced Resistance for Plant Disease Control: Maximizing the Efficacy of Resistance Elicitors

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ABSTRACT

Plants can be induced to develop enhanced resistance to pathogen infection by treatment with a variety of abiotic and biotic inducers. Biotic inducers include infection by necrotizing pathogens and plant-growth-promoting rhizobacteria, and treatment with nonpathogens or cell wall fragments. Abiotic inducers include chemicals which act at various points in the signaling pathways involved in disease resistance, as well as water stress, heat shock, and pH stress. Resistance induced by these agents (resistance elicitors) is broad spectrum and long lasting, but rarely provides complete control of infection, with many resistance elicitors providing between 20 and 85% disease control. There also are many reports of resistance elicitors providing no significant disease control. In the field, expression of induced resistance is likely to be influenced by the environment, genotype, and crop nutrition. Unfortunately, little information is available on the influence of these factors on expression of induced resistance. In order to maximize the efficacy of resistance elicitors, a greater understanding of these interactions is required. It also will be important to determine how induced resistance can best fit into disease control strategies because they are not, and should not be, deployed simply as "safe fungicides". This, in turn, will require information on the interaction of resistance elicitors with crop management practices such as appropriate-dose fungicide use.

It is now well documented that treatment of plants with various agents (e.g., virulent or avirulent pathogens, nonpathogens, cell wall fragments, plant extracts, and synthetic chemicals) can lead to the induction of resistance to subsequent pathogen attack, both locally and systemically (74). This induced resistance rarely leads to complete control of pathogens following subsequent inoculation, but rather results in, for example, a reduction in lesion numbers and size (34). Most notably, the expression of this induced resistance does not require the presence of major pathogen-specific resistance genes, although the defense mechanisms activated are those used in other forms of plant resistance to pathogens (21). Following application of an inducing treatment to a plant, defense mechanisms may be triggered directly or they may be triggered only once pathogen challenge has occurred (72). The defense responses activated include an oxidative burst, which can lead to cell death (21), thereby trapping the pathogen in dead cells; changes in cell wall composition that can inhibit pathogen penetration; and synthesis of antimicrobial compounds such as phytoalexins (20). Induction of systemic resistance can lead to the direct activation of defense-related genes, but also can lead to the priming of cells, resulting in stronger elicitation of those defenses or, indeed, other defenses following pathogen attack (11).

Induced resistance can be split broadly into systemic acquired resistance (SAR) and induced systemic resistance (ISR). SAR develops locally or systemically in response to, for example, pathogen infection or treatment with certain chemicals (e.g., 2,6-dichloroisonicotinic acid [INA]) is effective against a wide range of pathogens and is mediated by a salicylic acid [SA]-dependent process (74). In contrast, ISR develops as a result of colonization of plant roots by plant-growth-promoting rhizobacteria (PGPR) and is mediated by a jasmonate- or ethylene-sensitive pathway (50).

The prospect of broad-spectrum disease control using the plant's own resistance mechanisms has led to increasing interest in the development of agents which can mimic natural inducers of resistance (74). Research has focused on the elicitor molecules released during the early stages of the plant–pathogen interaction, and on the signaling pathways used to trigger defenses locally and systemically. The elicitors examined include carbohydrate polymers, lipids, and glycoproteins, and are either secreted by microorganisms or derived from the cell walls of fungi, bacteria, or plants (74) (e.g., elicitors derived from yeast cell walls) (52). Compounds which might mimic the action of SA include INA and S-methylbenzo[1,2,3]thiadiazole-7-carbothiate (acibenzolar-S-methyl) (ASM). Neither INA nor ASM possess antimicrobial activity in vitro, and they activate the same genes as does biological or SA induction of resistance (33). Indeed, ASM is the first synthetic chemical developed and marketed as a SAR activator and is marketed in Europe as BION and as ACTIGARD in the United States (74).

INDUCED RESISTANCE AND DISEASE CONTROL: EXAMPLES FROM CONTROLLED ENVIRONMENT AND FIELD STUDIES

There is a growing body of information on the efficacy of induced resistance under field conditions (70). In their review of induced resistance in conventional agriculture, Vallad and Goodman (70) highlighted 32 examples where ASM was found to provide disease control. In 28 of these studies, reductions in disease severity provided by ASM ranged from 4 to 80%, with only 3 studies reporting disease control in excess of 80% (70). ASM was marketed originally for the control of powdery mildew on wheat and barley in Europe (16) and, indeed, has been shown to reduce mildew infection on wheat by between 64 and 77% in...
field experiments (65,66). However, some of the best levels of disease control were demonstrated on dicotyledonous crops. For example, on tobacco, ASM provided 99% control of *Pseudomonas syringae* pv. *tabaci*, 91% control of *Cercospora nicotiana*, and 89% control of *Alternaria alternata* (10,49). They also highlighted 60 examples where PGPR were used to control crop diseases, with reductions in disease severity of less than 80% reported in 57 of the studies (70). Particularly high levels of disease control were achieved in cucumber, with *Bacillus pumilis* INR-7 and *Serratia marcescens* 90-166 providing 86 and 89% control, respectively, of *Erwinia tracheiphila* (78).

But what about the diverse range of other agents that have been shown to induce resistance? These include phosphates, amino acids, fatty acids, cell wall fragments, and avirulent pathogens (74) which characteristically are either far less toxic than fungi-cides or nontoxic because they do not act directly against pathogens but entirely through the plant’s defense mechanisms.

**Induced resistance using phosphate salts.** Dibasic and tribasic phosphate salts were shown to induce systemic protection against anthracnose in cucumber caused by *Colletotrichum lagenarium* (17) and later work demonstrated the broad spectrum of disease control achieved in cucumber using phosphates (40). It was speculated that basic phosphates applied to plants could sequester apoplastic calcium, altering membrane integrity and influencing the activity of apoplastic enzymes such as polygalacturonases, thereby releasing elicitor-active oligogalacturonides from plant cell walls (17,73). Indeed, subsequent work by Orober et al. (48) showed that phosphate-mediated resistance induction in cucumber was associated with localized cell death, preceded by a rapid generation of superoxide and hydrogen peroxide. These workers also detected local and systemic increases in levels of free and conjugated SA following phosphate application (48). Recent work on barley showed that application of phosphate to first leaves reduced powdery mildew infection by 89% in second leaves (39). Application of phosphate to first leaves led to significant increases in activities of phenylalanine ammonia-lyase, peroxidase, and lipoxygenase in second leaves, and activities of these enzymes were increased further following pathogen challenge (39). Phosphates also have been shown to provide disease control under field conditions. Thus, K₂HPO₄ applied to rice as a 50-mM spray reduced neck blast caused by the fungus *Pyricularia oryzae* by between 29 and 42%, with increases in grain yield of between 12 and 32% (36). Phosphate (K₂PO₄, 25 mM) applied to barley in a field trial reduced powdery mildew infection by up to 70% and gave an increase in grain yield of 12% compared with untreated controls (39). In cucumbers grown hydroponically, phosphate at 20 ppm applied to the hydroponic solution reduced powdery mildew infection by between 80 and 92%, with reductions of up to 91% in numbers of conidia produced on infected leaves (53).

**Induced resistance using β-aminobutyric acid.** The non-protein amino acid β-aminobutyric acid (BABA) has been shown to induce broad-spectrum resistance in a range of crops (32). BABA applied to tobacco as a 1-mM spray causes small necrotic lesions (8) and, when applied at 10 mM, led to the formation of reactive oxygen species, lipid peroxidation, induction of callose around lesions, and an increase in the SA content of leaves (61). Treatment with BABA has been reported to lead to induction of pathogenesis-related (PR) proteins. Treatment with BABA induced PR-1a, chitinase, and glucanase in tobacco, tomato, and pepper (9,61), but not in *Arabidopsis*, cauliflower, or tobacco (8,32,62). This suggests that induction of PR proteins may not be the only mode of action of BABA, which also leads to callose deposition, lignification, and hypersensitivity in some plants (9,61). BABA has been shown to move systemically in tomato, tobacco, and grape plants (9), and this may explain the systemic protection against diseases observed in these and other plants (9,61). In field trials with grapevines, BABA reduced infection by the downy mildew fungus *Plasmopara viticola* by 57% on cv. Chardonnay and by 98% on cv. Cabernet Sauvignon (54). Interestingly, mixtures of BABA and fungicides were even more effective in reducing infection in both cultivars.

**Induced resistance using oligosaccharides.** Oligosaccharides such as N-acetylchito-oligosaccharides and β-1,3-glucans are well known to act as elicitors of plant defenses. Chitosan is a deacetylated form of N-acetylchito-oligosaccharides containing polyβ-glucosamine and is a common polymer in shells of crustaceans, exoskeletons of insects, and cell walls of fungi (19). There are numerous reports of the protective effects of chitosan against pathogen infection in a range of crops. For example, chitosan seed treatment has been shown to protect tomato plants from crown rot and root rot (6) and to protect tomato seedlings against *Fusarium oxysporum* when applied as a foliar spray (7). A commercial formulation of chitosan developed by Glycogenesys Inc. (Boston), Elexa, contains 4% chitosan as its active ingredient and has been shown to protect a range of crops against important pathogens (1). In field trials with pear millet, Elexa was shown to reduce downy mildew severity by 58% when used as a seed treatment, by 75% when used as a foliar spray, and by 77% when used as a combined seed treatment and foliar spray (59). Elicitors derived from the yeast *Saccharomyces cerevisiae* also have been shown to control plant diseases, providing up to 95% control of powdery mildew infection in barley in field trials (51,52). In that work, elicitor applied with a reduced rate of fungicide gave better disease control than the elicitor used on its own.

**Induced resistance using probenazole.** The synthetic compound probenazole has been used to control rice blast caused by *Magnaporthe grisea* in Asia for more than 20 years and also protects rice from other diseases, including bacterial blight caused by *Xanthomonas oryzae pv. oryzae* (31). The probenazole-containing product Oryzemate provides long-lasting control of rice blast when applied to paddy fields or to seedling boxes. It is absorbed by roots and distributed throughout the plant, and the disease control it provides lasts for up to 70 days after application (31). Importantly, despite its extensive use since the 1970s, there have been no reports of resistance development in the blast fungus (31). Early studies showed that probenazole possessed only weak antimicrobial activity, leading workers to suggest that it activated defense responses in rice (75). Indeed, recent work has confirmed this and shown that probenazole and its active metabolite 1,2-benzisothiazole-1,1-dioxide induce SAR by triggering signaling at a point upstream of SA accumulation (42).

**FACTORS INFLUENCING THE EXPRESSION OF INDUCED RESISTANCE**

From what we have considered so far, it is clear that, although there are cases where elicitors of induced resistance can provide very high levels of disease control (>85%), there are many more examples of induced resistance providing lower levels of disease control (38). Indeed, there are also many reports of induced resistance not providing disease control. Thus, following a 2-year field trial with eight different cultivars of winter barley, Huth and Balke (29) concluded that ASM did not induce resistance to *Barley yellow dwarf virus* and, under controlled conditions, ASM did not induce resistance to *Phytophthora brassicae* in *Arabidopsis* or *P. infestans* in potato (60). More recently, in an evaluation of the effects of different agents on control of *X. axonopodis pv. citrulli* and *X. axonopodis pv. citrisc* on sweet oranges, ASM and the harpin protein (marketed as Messenger) failed to provide significant disease control (18). Because induced resistance is a plant response to attempted infection, it stands to reason that the expression of this response will be affected by a range of factors, including genotype and the environment. Therefore, how much do we know about the influence of these factors on the expression of induced resistance?
Effects of genotype on the expression of induced resistance. In monocots, resistance activated by ASM typically is very long lasting, whereas the lasting effect is less pronounced in dicots (47). Comparatively little information exists on the influence of genotype on induced resistance. In some of the first work on this topic, Steiner et al. (67) reported that the reduction in powdery mildew on barley following treatment with culture filtrate of 
B. subtilis
was cultivar specific and was most marked in partially resistant cultivars. In some later work, Martenelli et al. (37) studied resistance in barley to powdery mildew, induced by prior inoculation with an avirulent isolate of the powdery mildew fungus, 
Blumeria graminis
f. sp. hordei. They used three sets of near-isogenic lines of barley, each of which included cultivars with four different race-specific resistance genes. Martenelli et al. (37) found that the extent of induced resistance, expressed as a reduction in the number of colonies formed, differed in lines carrying different race-specific resistance genes, being most effective in lines with the Mla7 gene and least effective in lines with the Mla13 gene. When induced resistance was measured at a later stage of infection, as a reduction in spore production, the effect was greatest in Mla7 lines and was weakest in Mla6 and Mhi lines (37). A year later, Hijewegen and Verhaar (28) showed that resistance in cucumber to the powdery mildew fungus 
Sphaerotheca fuliginea, induced by treatment with INA, also was cultivar dependent, with the highest levels of induced resistance expressed in a partially resistant cultivar and much lower levels of resistance in susceptible cultivars. Different results were obtained by Dann et al. (12) working on soybean. They found that resistance to 
Sclerotinia sclerotiorum
induced by treatment with INA or ASM was greatest in susceptible cultivars, thereby showing that induced resistance is not related to major gene resistance. Cultivar-dependent differences in the expression of induced resistance also have been found in winter wheat, where increases in yield following ASM treatment were observed only in a susceptible cultivar (66), and in bell peppers expressing resistance induced by ASM treatment (56). In some interesting recent work, Romero and Ritchie (57) examined the effect of ASM in reducing the occurrence of race-change mutants of 
X. axonopodis
pv. 
vesicatoria
that overcome major resistance (R) genes in bell pepper. In field experiments, they found that decreased disease severity brought about by ASM treatment was associated with a reduction in the number of race-change mutants and a suppression of disease caused by such mutants. Based on these data, Romero and Ritchie (57) suggest that induced resistance agents may be useful for increasing the durability of genotype-specific resistance conferred by major R genes.

Genotypic effects on the expression of ISR by PGPR also have been demonstrated. Thus, in 
Arabidopsis thaliana,
the PGPR strain 
Pseudomonas fluorescens
WCS417 elicited ISR on all ecotypes examined, except ecotypes Wassilewskija and RLD (69,71). Subsequent research demonstrated the presence of a locus (ISR1) involved in the ethylene signaling pathway of 
A. thaliana. The ecotypes Wassilewskija and RLD carried a recessive trait that affected ISR by perturbing ethylene signaling, although these plants could still express SAR (68). Clearly, in ecotypes of 
A. thaliana, allelic variability exists in genes that exert an influence on ISR pathways.

The cost of induced resistance. Following an inducing treatment, there is usually a short lag period before local or systemic resistance is expressed (34). In fact, following the lag period, there are several possible outcomes in terms of resistance expression, the major ones being (i) defenses are triggered and there is no further change in defenses following pathogen challenge, (ii) defenses are triggered and there is a further increase in these defenses or the activation of a different set of defenses following pathogen challenge, and (iii) defense mechanisms are not expressed until pathogen challenge has occurred (72). Whichever of these scenarios prevails following pathogen challenge of an induced plant, there can be a gap of perhaps several days before the plant is protected by its induced defenses. This is a risky form of defense compared with constitutive resistance, where defenses are always present. The continued and indeed widespread existence of induced resistance suggests a selective advantage over constitutive resistance or that induced resistance is more targeted at the invading organism than we sometimes think (i.e., is more specific and less general). For example, the response of plants to the closely related elicitors ASM and SA is different, with different genes being upregulated (22). A possible explanation for this selective advantage lies with fitness costs, where resistant plants would have decreased reproductive success (e.g., seed production) than nonresistant plants under conditions where there was no pathogen pressure (24). These costs can include allocation costs arising from the diversion of metabolites and energy from growth and other processes toward defense; negative effects of the resistance against symbiotic interactions such as formation and functioning of mycorrhizas; and trade-offs where increased resistance to pathogens, for example, might compromise resistance to insect pests (24).

There is evidence that induced resistance to insect herbivores incurs costs (77). However, the situation with respect to pathogens is much less clear. In some pioneering work, Smedegaard-Petersen and Stolen (63) found that, when barley was heavily inoculated with an avirulent isolate of powdery mildew, there was a 7% reduction in grain yield and a 4% reduction in grain size and grain protein content compared with uninoculated control plants. They suggested that the greatly increased respiration rates observed early in the interaction were required to provide energy for resistance to the attempted powdery mildew infection, and that this was responsible for the reductions in grain yield and quality (63). In contrast, later work reported yield increases associated with induced resistance to powdery mildew infection in barley, compared with plants receiving no inducing treatment (46,67) or no effect on yield when yeast-derived elicitors were used to induce resistance to mildew in barley (51). However, none of these studies was designed to quantify allocation costs, because they all were conducted in the presence of pathogen challenge (24). When two such studies were conducted, using wheat and bean, differing results were obtained. In one study, using ASM applied to wheat in the absence of pathogen pressure, treated plants were shown to have reduced biomass and reduced numbers of ears and grains, with the most pronounced effects detected when nitrogen supply was limited (25). This work demonstrated quite clearly that use of ASM incurred allocation costs, supporting the “growth-differentiation balance” hypothesis, which assumes a metabolic competition between processes involved in plant growth and those necessary for plant differentiation (e.g., the synthesis of chemicals for plant defense) (26). In the second study, Iriti and Faoro (30) found that, in bean treated with ASM, the efficiency of photosystem II was not affected, as was seed quality, although there was a small, statistically insignificant reduction in seed yield. The authors concluded that ASM did not induce fitness costs in bean (30). The different results obtained in these two studies (25,30) probably reflect differences between species and in environmental conditions.

Some of the strobilurin group of fungicides, notably the BASF product F500, which contains pyraclostrobin as the active ingredient, have effects on the plant’s metabolism in addition to direct toxin activity against pathogens (27). These include claims of disease resistance induction. Strobilurins also often result in yield benefit beyond that attributable to disease control, partially accounted for by “leaf greening” effects in common with other fungicides such as the triazoles, but this again is genotype and environment dependent (5). Whereas in the absence of fungicide treatments, pathogen-challenged (diseased) and unchallenged genotypes still show high yield correlation, Newton and Thomas (43) found that fungicide-treated pathogen-challenged (no disease) and...
unchallenged genotypes showed no yield correlation. How much more effect on yield–genotype interactions might induced resistance or metabolic change agents have than the conventional fungicides used in this work?

Environmental effects and induced resistance. It stands to reason that, if induced resistance requires the diversion of resources away from growth and development toward defense, constraints on the availability of such resources should exert an impact on the expression of induced resistance. This also could affect the magnitude of any costs associated with induced resistance. Surprisingly, however, this aspect of induced resistance has received little attention until recently. As noted above, Heil et al. (25) found that allocation costs associated with use of ASM on wheat were greatest when nitrogen supply was limited. In later work, nitrogen supply was shown to exert a marked effect on the expression of both constitutive and induced resistance in *Arabidopsis* (13). In this work, constitutive levels of chitinase, chitosanase, and peroxidase and levels of chitinase and peroxidase in plants induced by ASM treatment were significantly lower under limiting nitrogen supply. Although the maximum activity of chitosanase was not significantly different in induced plants, the induction kinetics of the enzyme were severely affected by nitrogen limitation. Further, total soluble protein content decreased significantly in the first 12 h following ASM treatment (13), an observation that agrees with a number of studies demonstrating reductions in the expression of genes related to primary metabolism following elicitation of resistance (64). Such a downregulation of primary metabolism may be necessary in order to make available substrates and metabolites required for the de novo synthesis of resistance compounds (23). This concept has received further support recently following work by Dietrich et al. (14), which showed that *Arabidopsis* plants treated with ASM exhibited a growth reduction during the week following induction. Induced plants appeared to compensate for these growth reductions subsequently by increasing growth rates. These data are interesting in the light of previous work demonstrating increased rates of photosynthesis in upper, uninfected leaves of plants with lower leaves infected by a biotrophic pathogen (55). Murray and Walters (41) showed that upper, uninfected leaves of otherwise rust-infected broad bean plants exhibited not just increased rates of photosynthesis but also enhanced resistance to rust infection. Zeier et al. (79) showed that SAR development in *Arabidopsis* in response to infection by avirulent bacteria was completely lost when the primary infection process occurred in the absence of light. It would be interesting and useful to determine whether the initial downregulation of primary metabolism observed rapidly following resistance induction is followed thereafter by large increases in primary metabolism (e.g., photosynthesis). In the work of Dietrich et al. (14), the effect of resistance induction on seed production (i.e., whether fitness costs were induced or not) was determined by environmental conditions such as nitrogen supply, water stress, and competition with other plants. They observed costs, no costs, or even higher seed production by ASM-induced compared with uninduced controls, under different combinations of these environmental factors. Induced plants were most likely to incur fitness costs if grown in a competitive environment (14).

It has been known for some time that exposure to abiotic stress can influence plant resistance to pathogens. The interaction between environmental stress and plant disease was reviewed by Ayres in 1984 (2), when he proposed two possible outcomes of such an interaction. The first is the possibility that the negative effects of pathogens and abiotic stress might be additive, and the second possibility is that abiotic stress might alter plant resistance to pathogen infection, leading, for example, to enhanced resistance (2). In fact, water stress has been reported to increase susceptibility to foliar pathogens (45), but also has been shown to enhance resistance to powdery mildew in older leaves of barley grown in dry soil (3). In some interesting work demonstrating the importance of crop growth conditions for the expression of pathogen resistance, Newton and Young (44) and Baker et al. (4) found that relief of water stress leads to a breakdown of mlo resistance in spring barley, and sudden cold relief can have the same effects (K Stewart, S Gurr, and A Newton, unpublished data). More recently, Wiese et al. (76) showed that osmotic stress and proton stress led to the induction of active defenses against powdery mildew in barley. The defense was found to be dependent on the intensity of the stress and the pathogen was stopped by the formation of papillae (76). Interestingly, resistance to mildew was induced in the same concentration-dependent manner by application of abscisic acid to the rooting medium (76). These reports demonstrate the fact that the expression of host resistance can be influenced by abiotic stress, and highlight the need for such interactions to be taken into account when induced resistance is used in a field situation.

OTHER FACTORS THAT MAY INFLUENCE THE EFFECTIVENESS OF INDUCED RESISTANCE

The ultimate aim of research on induced resistance is its use to control disease in practical, commercial situations. As already indicated, because induced resistance is a host response, it is likely to be affected by the environment and by genotype. But we must not forget that induced resistance will need to fit into crop-protection programs. Therefore, it also will be important to consider the timing of application and frequency of application, and use it in conjunction with appropriate-dose fungicides. These are not trivial issues, because they will affect not only the performance of induced resistance but also its cost effectiveness. Should a resistance elicitor be applied before or after a fungicide application? The answer to this question may depend upon the particular pathogen or pathogens being targeted and the stage of crop growth when they are most prevalent and troublesome. Combinations of agents that induce resistance (e.g., ASM) with fungicides or biological control agents has been shown to provide effective disease control, especially in situations where achieving acceptable disease control is difficult. For example, a mixture of a strobilurin fungicide and ASM was shown to be effective in controlling *Albugo occidentalis* and increasing leaf quality in spinach (35), while a mixture of ASM and mancozeb was shown to have potential to provide protection against *Claviceps africana* on sorghum, especially where fungal isolates resistant to the usual fungicide treatment, triadimenol, were present (58). More recently, Gent and Schwartz (15) found that integration of ASM and biological control agents with copper hydroxide could be used to replace less desirable fungicides without compromising effective control of Xanthomonas leaf blight on onion. If agents that induce resistance are to be used in integrated disease control programs, then other questions need to be answered. Will incorporation of induced resistance into disease control programs be more expensive than current approaches? Will induced resistance represent a sustainable approach to disease control? These are all issues that need to be addressed, because it is not good enough to show that a new approach to disease control works under experimental conditions. However, their different mode of action dictates that they cannot and should not be applied in the same way as fungicides and, therefore, they represent a challenge to conventional marketing and agronomic approaches. Ultimately, farmers and growers need to be convinced that induced resistance really will provide a useful, practical, and sustainable approach to dealing with their disease problems.

CONCLUSION

Plant disease control in the 21st century faces considerable challenges. There are the continuing problems of pathogen adaptability leading to fungicide resistance and breakdown in the effec-
tiveness of host genetic resistance. Then there are the newer problems of a slowing down in the rate of delivery of new fungicides to the market and the increasing public concern related to the environmental effects of widespread fungicide use. There is also the potential problem of climate change and its impact on pathogen spread. The aim of plant pathologists always has been to keep one step ahead of the pathogens, through understanding every aspect of the plant–pathogen interaction and the factors that influence it. Because of these issues and the fact that plant–pathogen interactions continue to evolve, the need for understanding will not diminish. There clearly is a great deal we understand about induced resistance, but there is equally a great deal still to understand. In our haste to realize the great potential offered by induced resistance, we have paid too little attention to the costs after induction of pathogen resistance depend on environmental conditions. Plant Cell 8:629-643.


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