Rice Pti1a Negatively Regulates RAR1-Dependent Defense Responses

Akira Takahashi,a,1 Ganesh Kumar Agrawal,a,1,2 Muneeo Yamazaki,a Katsura Onosato,a Akio Miyao,a Tsutomu Kawasaki,b Ko Shimamoto,b and Hirohiko Hirochika,a,3

a Department of Molecular Genetics, National Institute of Agrobiological Sciences, Tsukuba, Ibaraki 305-8602, Japan
b Laboratory of Plant Molecular Genetics, Nara Institute of Science and Technology, 8916-5 Takayama, Ikoma 630-0101, Japan

Tomato (Solanum lycopersicum) Pto encodes a protein kinase that confers resistance to bacterial speck disease. A second protein kinase, Pti1, physically interacts with Pto and is involved in Pto-mediated defense signaling. Pti1-related sequences are highly conserved among diverse plant species, including rice (Oryza sativa), but their functions are largely unknown. Here, we report the identification of a null mutant for the Pti1 homolog in rice and the functional characterization of Os Pti1a. The rice pti1a mutant was characterized by spontaneous necrotic lesions on leaves, which was accompanied by a series of defense responses and resistance against a compatible race of Magnaporthe grisea. Overexpression of Pti1a in rice reduced resistance against an incompatible race of the fungus recognized by a resistance (R) protein, Pish. Plants overexpressing Pti1a were also more susceptible to a compatible race of the bacterial pathogen Xanthomonas oryzae pv oryzae. These results suggest that Os Pti1a negatively regulates defense signaling for both R gene–mediated and basal resistance. We also demonstrated that repression of the rice RAR1 gene suppressed defense responses induced in the pti1a mutant, indicating that Pti1a negatively regulates RAR1-dependent defense responses. Expression of a tomato Pti1 cDNA in the rice pti1a mutant suppressed the mutant phenotypes. This contrasts strikingly with the previous finding that Sl Pti1 enhances Pto-mediated hypersensitive response (HR) induction when expressed in tobacco (Nicotiana tabacum), suggesting that the molecular switch controlling HR downstream of pathogen recognition has evolved differently in rice and tomato.

INTRODUCTION

Plants have evolved surveillance and defense response systems to protect themselves from pathogen attack. The first step of defense against attempted microbial invasion is achieved by a pattern recognition receptor that detects a pathogen-associated molecular pattern (PAMP) (Zipfel and Felix, 2005; Chisholm et al., 2006). Pathogenic microbes have specialized systems that suppress or evade plant PAMP-triggered defenses and facilitate tissue invasion by secreting several effector proteins (Nurnberger et al., 2004; Chisholm et al., 2006). When a plant resistance (R) protein directly or indirectly recognizes a specific pathogen effector, which is often the product of a pathogen avirulence (avr) gene, the plant exhibits heightened defense against the pathogen (Jones and Takemoto, 2004; Jones and Dangl, 2006). The recognition of different pathogens by several R proteins appears to amplify a common set of defense responses and triggers rapid and strongly localized generation of reactive oxygen species, pathogen-related (PR) gene expression, and accumulation of antimicrobial compounds (Dangl and Jones, 2001; Durrant and Dong, 2004). These responses are often accompanied by localized programmed cell death known as the hypersensitive response (HR) at the site of pathogen invasion (Greenberg and Yao, 2004).

In the last decade, a large number of R genes from several species have been identified by map-based cloning, insertional mutagenesis, or various high-throughput methods (Hammond-Kosack and Parker, 2003). Sequence comparisons among these genes reveal a remarkable conservation of structural features, despite the diversity of the pathogens which their products recognize (Nimchuk et al., 2003). The largest class of R genes, termed the NB-LRR class, encodes a cytoplasmic protein with a Leu-rich repeat (LRR) and a nucleotide binding (NB) site. Although the signal components downstream of R proteins are thought to be conserved, only a few components that regulate the fundamental aspects of R protein–triggered responses have been isolated (Hammond-Kosack and Parker, 2003). Among those identified, RAR1 (required for Mla12 resistance), HSP90 (heat shock protein 90), and SGT1 (suppressor of the G2 allele of skp1) are required for resistance mediated by various NB-LRR R proteins (Shirasu and Schulze-Lefert, 2003; Piffanelli et al., 2004). The RAR1 protein is required by particular R proteins that are effective against bacterial, oomycete, and viral pathogens reported in barley (Hordeum vulgare), Arabidopsis thaliana, and tobacco (Nicotiana tabacum) (Freialdenhoven et al., 1994; Liu et al., 2002; Muskett et al., 2002; Tornero et al., 2002). It interacts with both HSP90 and

1 These authors contributed equally to this work.
2 Current address: Department of Biochemistry, University of Missouri, 204 Life Sciences Center, Columbia, MO 65211.
3 Address correspondence to hirohiko@nias.affrc.go.jp.
The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (www.plantcell.org) is: Hirohiko Hirochika (hirohiko@nias.affrc.go.jp).
Online version contains Web-only data.
Open Access articles can be viewed online without a subscription.
www.plantcell.org/cgi/doi/10.1105/tpc.106.047142
SGT1 and is considered to function as a molecular chaperon, in association with HSP90, to stabilize certain R proteins (Azevedo et al., 2002; Takahashi et al., 2003).

To date, there have been several reports suggesting a link between PAMP-triggered basal resistance and R protein–mediated resistance at the molecular level. In Arabidopsis, RIN4 is a negative regulator of PAMP signaling and is targeted by Pseudomonas syringae type-III effector AvrRpt2 for degradation, leading to the activation of an R protein, RPS2 (Kim et al., 2005). AvrB, a P. syringae effector protein, suppresses PAMP-triggered immunity through RAR1, which is indispensable for the stabilization of RPM1, the R protein corresponding to AvrB (Shang et al., 2006). rar1 mutations in Arabidopsis allowed enhanced growth of the virulent bacterial strain P. syringae DC3000 (Holt et al., 2005). Nb SGT1 is required not only for R protein–mediated HR induction but also for some non-host resistance responses (Peart et al., 2002). These observations suggest that the signaling pathways for PAMP-triggered immunity and R protein–mediated race-specific resistance substantially share common regulatory components.

The tomato (Solanum lycopersicum) R protein Pto confers race-specific resistance to the bacterial pathogen P. syringae pv tomato carrying the avirulence effector proteins AvrPto or AvrPtoB (Pedley and Martin, 2003). The Pto gene encodes a Ser/Thr protein kinase and is unique among several classes of known R proteins. Pto was shown to directly interact with both the bacterial effector proteins in a yeast two-hybrid system assay. However, little is known about the signal transduction mechanism downstream of the recognition event. A number of potential downstream components of the Pto signaling pathway have been reported, such as the protein kinase Pti1, and transcription factors Pti4, Pti5, and Pti6 (Pedley and Martin, 2003). Pti1 interacts with Pto and is phosphorylated by Pto in vitro. Overexpression of Pti1 in tobacco causes enhanced HR in leaves when challenged with P. syringae pv tabaci expressing AvrPto (Zhou et al., 1995). However, there is no direct evidence to support the involvement of Pti1 in Pto-mediated disease resistance.

Despite considerable efforts to find and characterize gain-of-function or loss-of-function mutants in several plant species, it is still unclear how R proteins transmit signals to downstream factors, what the limiting factors in evoking defense responses might be, or how the relationship between basal resistance and race-specific resistance is established. To develop more insight into plant defense signaling, we screened for mutants that had enhanced resistance to rice (Oryza sativa) blast disease from a collection of mutant lines generated by rice endogenous retrotransposon Tos17 insertion (Hirochika, 2001; Hirochika et al., 2004) and designated it ttm1 (for Tos17 triggered mutation1). The ttm1 lesion is recessively inherited, and homozygous ttm1 plants have stunted growth with spontaneous small and obscure lesions over both leaf surfaces (Figures 1A and 1B). Lesions appeared at ~30 d after sowing in the field and after ~40 to 50 d after sowing in a greenhouse. The variation in the timing of lesion appearance may be due to differences in growth conditions. Because the lesion pattern was similar to that seen during HR, we presumed that lesion formation results from induction of

**RESULTS**

**Identification of the ttm1 Mutant and Its Phenotype**

We identified a lesion mimic mutant (ND5001) among stable insertion mutant lines of Japonica rice cv Nipponbare (NB) produced by endogenous retrotransposon Tos17 (Hirochika, 2001; Hirochika et al., 2004) and designated it ttm1 (for Tos17 triggered mutation1). The ttm1 lesion is recessively inherited, and homozygous ttm1 plants have stunted growth with spontaneous small and obscure lesions over both leaf surfaces (Figures 1A and 1B). Lesions appeared at ~30 d after sowing in the field and after ~40 to 50 d after sowing in a greenhouse. The variation in the timing of lesion appearance may be due to differences in growth conditions. Because the lesion pattern was similar to that seen during HR, we presumed that lesion formation results from induction of
the HR pathway triggered by the ttm1 mutation. To examine whether the ttm1 mutation activates defense responses, we inoculated mutant and wild-type plants with the rice blast fungus *Magnaporthe grisea*. The cultivar NB has Pish, an R protein that is active against rice blast fungus isolates containing avrPish, such as race 102.0 (incompatible), but not against race 003.0 (compatible) (Imbe and Matsumoto, 1985). Disease resistance of the ttm1 mutant line against the incompatible race was comparable to that of NB (data not shown). However, ttm1 plants exhibited strong resistance against the compatible race after lesion formation (Figure 2A) but not before lesion formation (data not shown). The ttm1 plants expressed defense-related genes, PR1b, PR5, PR10α, and PAL, after the appearance of lesions (Figure 2B). Furthermore, momilactone A, the major phytoalexin of rice (Cartwright et al., 1977), accumulated in uninfected leaves with lesions to a level ~150-fold higher than NB at the same developmental stage but was negligible in ttm1 leaves without lesions (Figure 2C). Thus, a series of defense responses were activated in the ttm1 mutant at developmental stages after the appearance of lesions.

**Figure 2.** Defense-Related Phenotypes of ttm1.

(A) NB and ttm1 were inoculated with a compatible race (003.0) of *M. grisea*. Lesions are shown on leaf blades 10 d after inoculation.

(B) Total RNA was extracted from lesion-negative (−) and lesion-positive (+) leaves of ttm1 mutants and NB. RNA gel blots of 10 μg total RNA were hybridized with radiolabeled probes of defense-related genes as indicated. rRNA is shown by staining with methylene blue as a loading control.

(C) A rice phytoalexin, momilactone A, was extracted from lesion-negative (−) and lesion-positive (+) leaves of ttm1 mutants and NB and quantified by the method described in Methods. FW, fresh weight.

These results suggest that the causative gene of the ttm1 mutant negatively regulates the defense signaling leading to HR induction.

### Cloning of Ttm1

To isolate the Ttm1 allele, we extracted genomic DNA from the progeny of ttm1 heterozygotes and subjected it to DNA gel blot analysis to examine the cosegregation of *Tos17* with the mutant phenotype. Genomic DNA flanking cosegregating *Tos17* was isolated by thermal asymmetric interlaced PCR with a *Tos17*-specific primer and degenerate primers (Liu et al., 1995). The PCR product was used as a probe for DNA gel blot hybridization to confirm the cosegregation with the lesion mimic phenotype. As expected, only the plants with the mutant phenotype carried a homozygous insertion (data not shown). A search of the GenBank nucleotide database using the BLAST program with the flanking sequence revealed that *Tos17* was inserted in the third exon of *Os Pti1a* (Figure 3A). This gene had high similarity to *Sl Pti1* (87% similarity at the amino acid level), *Sl Pti1* encodes a cytoplasmic protein kinase and was originally identified by a yeast two-hybrid screen as a protein that interacts with Pto, which is a tomato R protein to *P. syringae pv tomato*, the causative agent of bacterial speck disease (Zhou et al., 1995). Because tomato Pti1 is phosphorylated by Pto, but Pto is not phosphorylated by Pti1, Sl Pti1 seems to function downstream of Pto in a phosphorylation cascade (Sessa et al., 2000). The predicted *Os Pti1a* gene would encode 361 amino acids, and its deduced *M* was 39.3 kD. RNA gel blots showed that *Os Pti1a* transcripts were not detected in *Os pti1a* (ttm1) homozygous mutants but were detected in wild-type and *Os pti1a* heterozygous mutants (Figure 4A). Therefore, *Os pti1a* is a null mutation. A database search also revealed that there is another homolog of *Sl Pti1* in rice. It was designated *Os Pti1b*, and its predicted product has an 83% similarity to *Sl Pti1* and 81% to *Os Pti1a* (Figures 3B and 3C). An amino acid sequence alignment of deduced Pti1 proteins showed that the protein kinase domain was highly conserved, but the N-terminal regions were highly variable (Figure 3B). The Thr residue at 233 (Thr-233) in Sl Pti1, which is the major site phosphorylated by Pto, was conserved in both *Os Pti1a* and *Os Pti1b* (Figures 4B). Its predicted product has an 83% similarity to *Sl Pti1* and 81% to *Os Pti1a* (Figures 3B and 3C). An amino acid sequence alignment of deduced Pti1 proteins showed that the protein kinase domain was highly conserved, but the N-terminal regions were highly variable (Figure 3B). The Thr residue at 233 (Thr-233) in Sl Pti1, which is the major site phosphorylated by Pto, was conserved in both *Os Pti1a* and *Os Pti1b* (Sessa et al., 2000). The highly similar protein sequences of *Os Pti1a* and *Os Pti1b* suggest that these proteins are functionally redundant. Nevertheless, the disruption of just one of them, *Os Pti1a*, was adequate to trigger spontaneous cell death and defense responses. To further explore this observation, we analyzed the transcript levels of *Os Pti1a* and *Os Pti1b* (Figure 4B). *Pti1a* was detected abundantly in the roots, young leaves, adult leaves, and preemergent panicles but not in ripening panicles. By contrast, *Pti1b* transcripts were barely detectable in each of the organs tested. The negligible expression of *Pti1b* is thus the likely reason that the loss of *Pti1a* is sufficient to trigger cell death and defense responses despite the presence of paralog *Pti1b*. To examine *Pti1b* function, we isolated an *Os pti1b* knockout line (ND4512). This line grew as healthily as the wild type without lesion formation and did not exhibit any enhanced resistance (data not shown). We then crossed this line with the *Os pti1a* knockout line to produce a *pti1a pti1b* double knockout mutant. The double mutant was morphologically and developmentally indistinguishable from the *pti1a* single mutant (data not shown).
These results suggest that Os Pti1b has only a minor function, if any, which is consistent with its extremely low levels of expression.

Os Pti1a Is a Functional Protein Kinase

To determine whether the protein encoded by Os Pti1a is a functional protein kinase, it was expressed as a polyhistidine-tagged protein in *Escherichia coli*. Incubation of the purified fusion protein with $\gamma$-32P-ATP in an in vitro kinase assay showed that Os Pti1a was capable of strong autophosphorylation (Figure 3D). The K96N mutation, which is known to completely abolish the autophosphorylation activity of Sl Pti1 (Zhou et al., 1995), abolished the autophosphorylation activity of Os Pti1a (Figure 3D), indicating that Os Pti1a encodes a functional protein kinase similar to Sl Pti1.

Os Pti1a and Sl Pti1 Complement the Os pti1a Mutant Phenotype

To confirm that the null mutation of Os Pti1a causes lesion formation and the induction of defense reactions, we screened for other allelic mutants from Tos17 insertion mutant lines by a PCR-based method using specific primers for Tos17 and Os Pti1a. Unfortunately, however, no allelic mutants were identified. We then transformed a full-length cDNA of Os Pti1a under the control of the cauliflower mosaic virus 35S promoter into the pti1a homozygous mutant. We obtained three independent transgenic lines and used their T1 and T2 generations for the following analysis. None of the transgenic plants was stunted, nor did they have lesions, and they were as healthy as the wild type (Figure 5A). The upregulation of PR1b expression observed in the pti1a line was
lost when Pti1a was overexpressed in this mutant (Figure 5B). These transgenic lines were not resistant to a compatible race of M. grisea, unlike the background pti1a mutant, indicating that the expression of Os Pti1a cDNA complemented the pti1a mutant phenotypes. The loss of Pti1a function thus results in Os pti1a mutant phenotypes.

Transgenic tobacco plants that overexpress Sl Pti1 cDNA show enhanced HR in leaves when challenged with P. syringae pv tabaci strains carrying the avirulence gene avrPto, suggesting that Sl Pti1 functions as a positive regulator of Pto-mediated cell death and disease resistance (Zhou et al., 1995). This contrasts strikingly with the observed phenotypes of the Os pti1a mutant. To investigate this apparent discrepancy, we expressed Sl Pti1 cDNA in the Os pti1a homozygous mutant under the control of the 35S promoter. Interestingly, the expression of Sl Pti1 cDNA blocked lesion formation and PR1b expression in the Os pti1a mutant (Figures 5A and 5B). Similar results were obtained with other three independent lines. These results indicate that the two Pti1 proteins, Sl Pti1 and Os Pti1a, are functionally equivalent. Nevertheless, the contrasting phenotypes of the mutant and transgenic plants indicate that a downstream molecular switch controlling HR has evolved differently in monocotyledonous rice and dicotyledonous tomato.

**Os Pti1a Overexpression Reduces Plant Resistance**

As described, the Os pti1a mutant produces spontaneous lesions resulting from the activation of defense responses in the absence of any pathogenic signal. This result implies that Os Pti1a is involved in the negative regulation of the defense signaling pathway. This prompted us to overexpress full-length Os Pti1a cDNA (Os Pti1a-OE) under the control of the 35S promoter in the NB background to determine what effect it would have on defense responses with rice blast fungus. Accumulation of Os Pti1a proteins in the T1 generation was measured using a specific antibody against the nonconserved N-terminal region of Pti1a (Figure 6B). Pti1a protein was detected in non-transgenic and vector control (25.3) plants but not in the pti1a homozygous mutant, confirming that this antibody is specific to Os Pti1a. We selected three individuals each from the T1 progeny of two independent transgenic lines (30 and 34) and inoculated them with...
effects are presented as average with a lower level of Os Pti1a accumulation. Results from three independent lines, 30 and 34) or empty vector (vector) and the pish mutant. The plus sign indicates transgenic plants in Os Pti1a OE lines against this incompatible race (see Supplemental Figure 1 online). Pi19-mediated defense response is much stronger than that mediated by Pish; therefore, it is likely that the effects of Os Pti1a overexpression on compatible pathogen interactions were masked by the effective defense reaction. However, we cannot exclude the possibility that Os Pti1a is not involved in all R protein–mediated defense responses.

Effects of Os Pti1a Overexpression on Compatible Pathogen Interactions

To investigate the effects of Os Pti1a overexpression on compatible interactions, we first tested a compatible race of the rice blast fungus (race 003.0). However, there was no enhanced susceptibility in Os Pti1a-OE lines to this race of the fungus (data not shown). Presumably, the effect of Os Pti1a overexpression on the compatible interaction is too small to be detectable against the strong pathogenicity of this fungus. We then tested the rice pathogen Xanthomonas oryzae (Xoo), the causal agent of rice bacterial blight disease. Because wild-type NB exhibits moderate levels of resistance against compatible races of Xoo (race 1), this pathogen seemed to provide a suitable system to assess the effect of Os Pti1a overexpression on compatible pathogen interactions. We used the T2 generation of Os Pti1a-OE plants for the inoculation of Xoo, after examining Pti1a protein levels by immunoblotting (Figure 6E). Xoo-induced lesions were 1.5-fold longer in the leaves of Os Pti1a-OE lines than in the vector control plants (Figures 6D and 6F), indicating that Pti1a suppresses basal resistance against the bacteria. The Os pti1a null mutant has severe growth defects, which made it difficult to evaluate its resistance against Xoo. Among 35S:Os Pti1a lines, in leaves of generation T2 overexpression lines (Os Pti1a-OE), cosuppression lines (Os Pti1a-CS), and vector control.

(E) Os Pti1a protein in total protein was detected with anti-Os Pti1a antibody. The bottom panel shows the Coomassie blue–stained gel. (F) Lesion length measurements for the T2 generation of each line. The graphs depict the mean ± s.d from three independent experiments (n > 20).
however, we were able to obtain Os Pt1a-cosuppressed lines (Os Pt1a-CS) that did not exhibit growth defects. In Os Pt1a-CS lines, Pt1a protein accumulated at the levels comparable with those in Os Pt1a-OE lines during early developmental stages (data not shown). The proteins declined to negligible levels as the plants developed (Figure 6E), suggesting that the cosuppression occurred in the middle of development. This delayed occurrence of cosuppression is a likely mechanism for the normal growth of the cosuppressed lines. When tested after Os Pt1a protein levels had declined, Os Pt1a-CS plants showed strong resistance against Xoo compared with wild-type and Os Pt1a-OE lines. These results indicate that Os Pt1a also has a negative effect on rice plant resistance to compatible pathogens.

Os Pt1a Negatively Regulates RAR1-Mediated Defense Signaling

The recognition of pathogen invasion by R proteins is followed by rapid activation of the defense signal cascade. Some regulatory proteins involved in gene-for-gene resistance have been characterized in Arabidopsis, barley, and tobacco. One such regulatory protein, RAR1, is required for the functioning of various R proteins and acts upstream of HR induction (Shirasu et al., 1999). To examine the genetic interaction between Pt1a and RAR1 in rice, we silenced Os RAR1 expression by RNA interference (RNAi) in the Os pt1a homozygous mutant (Os pt1a-rar1i) and in NB (NB-rar1i). RT-PCR analysis demonstrated that RAR1 transcript levels decreased to ~10% of the wild type in both Os pt1a-rar1i and NB-rar1i (Figure 7B). Silencing of RAR1 in NB caused no visible phenotype without pathogen challenge; however, the effect of RAR1 silencing was striking in Os pt1a-rar1i plants. In these plants, the spontaneous lesion formation due to pt1a mutation was completely abolished. In addition, RT-PCR revealed that the expression of PR1b was reduced in pt1a-rar1i lines compared with the pt1a control line carrying empty vector (Figure 7B). The dwarf phenotype of the pt1a mutant was also suppressed, although incompletely. Similar phenotypes were observed in >10 pt1a-rar1i plants from four independent lines. These results indicate that rar1i is genetically epistatic to Os pt1a.

To investigate RAR1 dependence of cell death induction in other lesion mimic mutants, we suppressed Os RAR1 in rice mutants cdr1 and cdr2 (Takahashi et al., 1999), which are characterized by spontaneous cell death and a series of defense responses similar to the pt1a mutant. Os RAR1 suppression, however, affected neither the spatial pattern nor the timing of lesion formation in cdr mutants (see Supplemental Figure 2 online), indicating that RAR1-dependent cell death induction is not general with lesion mimic mutants.

In rice, RAR1 silencing did not compromise the resistance mediated by three R genes for blast fungus (N.P. Thao, L. Chen, A. Nakashima, S. Hara, K. Umemura, A. Takahashi, K. Shirasu, T. Kawasaki, and K. Shimamoto, unpublished data). These observations are in agreement with previous reports that RAR1 is not required for the functioning of all R genes (Shirasu and Schulze-Lefert, 2003). We found that both NB-rar1i and Os pt1a-rar1i plants retained resistance against incompatible blast fungus races carrying avrPish (data not shown), indicating that Os RAR1 is not required for Pish-mediated resistance either. Then, we
examined the effects of RAR1 silencing on the enhanced resistance of the pt1a mutant against a compatible race of blast fungus. The enhanced resistance against the compatible race of blast fungus observed in the pt1a mutant was largely cancelled in Os pt1a-rar1i plants (Figures 7C and 7D). These results clearly indicate that the function of Os Pt1a in the negative regulation of blast resistance is also dependent on Os RAR1.

Recent biochemical studies suggest various requirements for RAR1 in an R protein–triggered signaling pathway seem to reflect its role as a protein chaperone to stabilize or protect R protein complexes from degradation (Nimchuk et al., 2003; Jones and Takemoto, 2004). If Os Pt1a functions in close proximity to R proteins, the stability of the Os Pt1a protein could be regulated by a chaperone activity involving RAR1. To examine this hypothesis, we measured the levels of Pt1a protein accumulation in NB-rar1i. However, we found no significant difference in Pt1a levels between NB and NB-rar1i plants (Figure 7D), indicating that the stability of Os Pt1a protein does not depend on RAR1 activity.

DISCUSSION

SI Pt1 was originally identified as a protein that interacts with an R protein Pto in tomato. Because SI Pt1 is phosphorylated by Pto and encodes a Ser/Thr protein kinase, it is thought that SI Pt1 functions downstream of Pto and transmits a defense signal to downstream components through its protein kinase activity (Pedley and Martin, 2003). However, the genetic data supporting a direct involvement of SI Pt1 in Pto-dependent disease resistance is very limited. Here, we provide genetic evidence that a rice homolog of Pt1, Os Pt1a, negatively regulates both R protein–mediated resistance and basal resistances in a Os RAR1–dependent manner.

Os Pt1a Functions as a Negative Regulator of R Protein–Mediated and Basal Resistance

Loss of Os Pt1a induced resistance against compatible races of both M. grisea and Xoo (Figures 2 and 6D). Overexpression of Os Pt1a reduced Psh-mediated resistance to an incompatible race of the fungus. Os Pt1a overexpression also reduced basal resistance to a compatible race of Xoo (Figure 6). Thus, Os Pt1a appears to negatively regulate plant resistance to both incompatible and compatible pathogens. Recent studies suggest that the R protein–mediated signaling pathway shares some components with the PAMP-triggered signaling pathway for basal resistance. Our results seem to be consistent with this notion. A possible explanation for our results would be that Pt1a lies at the point shared by both R protein– and PAMP receptor–mediated signaling pathways. Pt1a presumably suppresses defense signal transduction through modification of the common signaling components by its phosphorylation activity. Some results with regard to disease resistance, however, are apparently inconsistent with this conclusion. One of the inconsistencies is that no enhanced resistance was observed when the pt1a null mutant was challenged with incompatible M. grisea (containing avrPsh). It could be that the strong Psh-mediated resistance masked the enhanced resistance in the pt1a mutant. Another inconsistency is that Os Pt1a-OE plants did not show reduced resistance to a compatible race of M. grisea. This is presumably because the strong pathogenicity of this pathogen overcomes Pt1a function.

Os Pt1a function Is Dependent on RAR1

We have shown that the silencing of Os RAR1 cancels the lesion formation, PR gene expression, and acquired resistance against the compatible race of the blast fungus induced in the pt1a mutant, suggesting that Pt1a functions as a negative regulator of the rice defense signaling pathway genetically upstream of RAR1. In several plant species, RAR1 is required for the functioning of particular R proteins. In addition, rar1 mutation allowed enhanced susceptibility against the virulent bacterial strain P. syringae DC3000 in Arabidopsis and against the virulent fungus M. grisea in barley (Holt et al., 2005; Jarosch et al., 2005). Thus, RAR1 functions as a positive regulator of both basal resistance and gene-for-gene resistance. Indeed, in rice, RAR1-RNAi plants impaired basal resistance to blast fungus and bacterial blight (N.P. Thao, L. Chen, A. Nakashima, S. Hara, K. Umemura, A. Takahashi, K. Shirasu, T. Kawasaki, and K. Shimamoto, unpublished data). RAR1 is known to function as a molecular chaperone to stabilize NB-LRR protein in Arabidopsis, barley, and tobacco, although it is unclear whether Os RAR1 interacts with NB-LRR proteins in rice. On the basis of our observations, we propose two models for the defense signaling pathway in rice featuring Pt1a and RAR1. Since RAR1 functions as a molecular chaperone in other plant species, we postulate that Os RAR1 stabilizes an unknown protein, X, a presumptive essential component in the defense signaling pathway. In one model (Figure 8A), we propose that Os RAR1 positively regulates the signaling pathway through X, and Os Pt1a negatively regulates the signaling indirectly by repressing RAR1. In the second model (Figure 8B), Pt1a negatively regulates the defense signaling, which is dependent on RAR1 through the stabilization of X, by directly acting on the pathway upstream or downstream of X. In either model, upregulation of Pt1a and downregulation of RAR1 should

![Figure 8](image.png)
result in the same outcome with respect to defense reactions. Our data are not necessarily consistent with this prediction: suppression of Os \textit{RAR1} in NB did not affect Pish-mediated resistance, whereas overexpression of \textit{Pti1a} reduced the gene-for-gene resistance. However, given that RNAi-mediated down-regulation is usually leaky, the downregulation of Os \textit{RAR1} expression in our Os \textit{RAR1}-RNAi transformants could be less effective than the effects of \textit{Pti1a} overexpression, irrespective of whether the negative regulation of the defense signaling pathway by \textit{Pti1a} is indirect (Figure 8A) or direct (Figure 8B).

One possibility for constitutive activation of defense responses including cell death in the absence of Os \textit{Pti1a} is explained by the guard hypothesis (Dangl and Jones, 2001). A knockout mutant of \textit{Arabidopsis RIN4}, which is a negative regulator of basal resistance, is embryo-lethal, and the lethality was suppressed by elimination of the \textit{R} gene \textit{RPS2} or delayed by a \textit{rar1} mutation leading to reduction of \textit{RIN4} protein accumulation, indicating that the elimination of \textit{RIN4} results in inappropriate \textit{RPS2-RAR1} activation (Mackey et al., 2003; Belkhadir et al., 2004). The relationship between Os \textit{Pti1a} and X-\textit{RAR1} is reminiscent of that between \textit{RIN4} and \textit{RPS2-RAR1}. A possible model based on this consideration would be that the unknown protein X is an NB-LRR protein, and the elimination of Os \textit{Pti1a} invokes the NB-LRR-RAR1 complex, leading to activation of the signaling pathway that results in cell death and defense induction.

### Defense Signal Transduction Mediated by Protein Phosphorylation

Protein phosphorylation appears to play a fundamental role in the early response of disease resistance. Some \textit{R} proteins, including tomato \textit{Pto} and rice \textit{Xa21}, or the PAMP receptor \textit{FLS2} have protein kinase activity (Martin et al., 1993; Song et al., 1995; Gomez-Gomez and Boller, 2000). Calcium-dependent protein kinase and mitogen-activated protein kinase are well known as important regulators of the defense signaling cascade (Romeis, 2001). Furthermore, pharmacological analyses demonstrated that many protein kinase activities are required for the induction of both \textit{R} protein- and PAMP receptor-mediated defense reactions (Lamb and Dixon, 1997; Takahashi et al., 1999). Recently, phosphor proteomics approaches have identified many proteins that undergo phosphorylation after treatment by elicitors or chemical inducers (Peck et al., 2001). However, in contrast with animals, only a few phosphorylation cascades have been characterized in plants. Therefore, identification of a protein kinase, or kinases, that phosphorylates Os \textit{Pti1a} could provide some very useful clues to understanding phosphorylation-mediated signaling in the negative regulation of defense. The Thr residue at 233 (Thr-233) in SI \textit{Pti1}, which is the major phosphorylation site of \textit{Pto} (Sessa et al., 2000), is conserved at the corresponding positions in both Os \textit{Pti1a} (Thr-233) and Os \textit{Pti1b} (Thr-236). This may imply that an as yet unidentified rice ortholog of \textit{Pto} may be an upstream protein kinase that phosphorylates \textit{Pti1a}.

### Genetic Screening in Rice

Using transgenic tobacco, SI \textit{Pti1} was shown to be a positive regulator of HR induction triggered by \textit{Pto-avrPto} interaction (Zhou et al., 1995). However, there are no loss-of-function data to support the involvement of SI \textit{Pti1} in Pto-dependent disease resistance. This may be explained by its functional redundancy because SI \textit{Pti1} appears to be a member of a gene family that encodes a group of closely related protein kinases (Mysore et al., 2002). \textit{Arabidopsis} has at least five \textit{Pti1} homologs in its genome. Rice has only two highly conserved \textit{Pti1} isoforms, \textit{Pti1a} and \textit{Pti1b} (Figure 3). \textit{Pti1a} alone was isolated in our genetic screens despite the apparent genetic redundancy likely because of the extremely low expression level of \textit{Pti1b} compared with \textit{Pti1a} (Figure 4). This case illustrates the merits of using rice instead of \textit{Arabidopsis} in a genetic approach to find novel proteins in a conserved signal transduction pathway when redundancy could be a problem in \textit{Arabidopsis} and other plant species. Interestingly, expression of SI \textit{Pti1} complements the Os \textit{Pti1a} phenotype (Figure 5), indicating that SI \textit{Pti1} acts as a negative regulator of the HR response in rice, while it behaves as a positive regulator in tobacco. Therefore, although protein function is conserved beyond plant species, the signal cascade leading to HR downstream of \textit{Pti1a} may have evolved differently in rice and tomato.

Despite the passage of more than a decade since the first molecular cloning of \textit{R} proteins, there is very little definitive understanding of how they transduce pathogen recognition signals or activate defense responses, including HR. Os \textit{Pti1a} functions as a negative regulator of defense responses associated with HR downstream of the \textit{R} protein and PAMP signaling. Therefore, further understanding of Os \textit{Pti1a} function should lead to a fuller understanding of the molecular mechanisms of HR induction and defense signaling pathways.

### METHODS

#### Plant and Pathogen Materials

The \textit{pTi1a} mutant is derived from rice (\textit{Oryza sativa} \textit{Japonicum} cultivar NB mutant lines induced by insertion of the rice endogenous retrotransposon \textit{Tos17} (Hirochika, 2001). NB carries the blast resistance genes \textit{Pish} and \textit{P19}. Strains of \textit{Magnaporthe grisea}, Kyu89-246 (MAFF101506; race 003.0) as compatible and Kyu77-07A (avrPish; Race 102.0) and CHNOS8-3-1 (avr19; Race 000.0) as incompatible races, were used in this experiment. Thus, NB is resistant to Kyu89-246 and CHNOS8-3-1 and susceptible to Kyu-77-07A. \textit{M. grisea} was grown on oatmeal agar medium (30 g/L oatmeal, 5 g/L sucrose, and 15 g/L agar) at 22°C. Seedlings were inoculated at the four- to six-leaf stage by spraying an aqueous spore suspension containing 10^8 to 10^9 spores per mL to run off. Inoculated seedlings were kept in a dark chamber with a moisture-saturated atmosphere at 24°C for 20 h and then maintained at 27°C and 70 to 80% relative humidity in a greenhouse. Disease development was monitored 1 week after inoculation. Lesion size per each leaf was measured and calculated using a digital microscope VXHS00 system (KEYENCE). Methods for the punch infection of the leaf blade with the blast fungus have been described (Takahashi et al., 1999). Bacterial blight inoculation experiments were performed with the Japanese \textit{Xoo} race 1 using a scissors-dip method (Kauffman et al., 1973). Lesion development was scored on rice leaves 21 d after inoculation by measuring margin progression with a ruler.

#### RNA Analysis

Total RNA was isolated from rice seedling roots, leaves, or panicles as described previously (Agrawal et al., 2001), separated on 1.2% (w/v) formaldehyde-denaturing agarose gels, and blotted onto nylon membranes.
(Hybond N+; Amersham). The cDNA fragments corresponding to OsPt1a, Ptib, Pr1b, Pr5, Pr10a, and PAL were amplified by PCR from wild-type leaf cDNAs using gene-specific primers (Takahashi et al., 1999; Agrawal et al., 2000). The cDNA fragment corresponding to Sl Pt1 was amplified by PCR from tomato leaf cDNA using the specific primers L (5'-CCACATTCTCAGAAGGTTAGAAC-3') and R (5'-CACAATT-CAGATCCTCTTGG-3'). Primers for RT-PCR analyses were 5'-AGCTCCTAGTAAGAGCCAGACC-3' and cDNA875R (5'-AGGTGTCGTAATTCCAGCAGTCATG-3') for Os Pt1a, AOL33 (5'-TCATGACGCATGAAACAGTTGGAAGG-3') and AOL34 (5'-GTGGTGACTAAGCTTCTCAA-3') for Os Ra1, AOL37 (5'-AGGTATCCAGTGCCATTG-3') and AOL38 (5'-TATGGACCCTGACCTGTTAC-3') for Pr1b, and Os Act1L (5'-GCCATCTCTGCTAGG-3') and Os Act1L (5'-GTACCCGATCAGGC-ATCTG-3') for Actin.

Phytoalexin Measurement
For measuring the accumulation of the phytoalexin momilactone A, leaves (the middle portion only) from three to four individual plants were used. Leaf samples were harvested from wild-type plants and from mutant plants before and after lesion formation. Harvested leaves were immediately frozen in liquid nitrogen to prevent touch- or wound-induced accumulation of phytoalexins. Quantification of momilactone A was performed as described previously (Takahashi et al., 1999). Briefly, leaves were cut into small pieces, transferred to a glass test tube containing 5 mL of 80% aqueous methanol, and boiled for 5 min. Three microliters of the crude extract was injected onto an HPLC and analyzed by liquid chromatography–tandem mass spectrometry (ibidi).

Isolation of Tos17 Insertion Sites and Reverse Genetic Analysis
Sequences flanking Tos17 insertions were amplified by thermal asymmetric interlaced PCR as previously described (Yamazaki et al., 2001). For reverse genetic analyses, Tos17-specific primers were used in combination with the Os Pt1a and Os Pr1b gene-specific primers. Two Tos17 (T17F-1 [5'-ACCACTTCAGAGTTGTTGTCG-3'] and T17R-1 [5'-CGACCAACAGTGTAAGTGTCGAGC-3']) and two Os Pt1a– or two Os Pt1b–specific primers were used in all possible combinations for PCR amplification of genomic DNA from each pooled DNA sample.

Sequence Analysis
Multiple sequence alignments were produced with a Web-based version of ClustalW (http://crick.genes.nig.ac.jp/homology/clustalw-e.shtml) using default settings (Matrix = blossom; GAPOPEN = 0, GAPEXT = 0, GAPDIST = -8, and MAXDIV = 40). The phylogenetic tree was calculated using the neighbor-joining method and bootstrap analysis (1000 replicates) using PHYLIP via the same website and visualized with Treeviewer version 1.6.6 (http://taxonomy.zoology.gla.ac.uk/rod/rod.html).

Rice Transformation
To overexpress Os Pt1a and Sl Pt1 cDNA, the coding sequences were cloned into the Ti-based vector pZPP2Ha3(+) downstream of the cauliflower mosaic virus 35S promoter, and Agrobacterium tumefaciens–mediated transformation of rice callus was performed according to a published protocol (Hiei et al., 1994; Fuse et al., 2001). Plants regenerated from hygromycin-resistant calluses were grown in an isolated greenhouse. For the complementation experiment, seeds harvested from the hygromycin-resistant calluses were derived from leaves in 100 mM Tris-HCl, pH 8.5, 4% (w/v) SDS, 20% (w/v) glycerol, and 2% (w/v) 2-mercaptoethanol and separated on 10% (w/v) SDS-PAGE gels.

Expression of Proteins and in Vitro Kinase Assay
Os Pt1a and its mutantized form (K96N) were expressed as fusion proteins with an N terminus poly-histidine tag using a bacterial expression system (in vitro) following the supplier’s instructions. Proteins were purified by immobilized metal ion affinity chromatography and applied for autophosphorylation assay as described (Zhou et al., 1995) with small modifications (addition of NaCl to the reaction buffer to a final concentration of 100 mM). Proteins were fractionated by SDS-PAGE and stained with Coomassie Brilliant Blue, and the 32P-labeled fractions were detected by autoradiography.

Acknowledgments
We thank Morifumi Hasegawa for measuring momilactone A and Mayuko Yamazaki for her technical assistance. We gratefully acknowledge critical comments from Hiroshi Tatsuki. This work was supported by a grant from the Ministry of Agriculture, Forestry, and Fisheries of Japan (Green Technology Project IP-4002).

Received September 5, 2006; revised September 3, 2007; accepted September 10, 2007; published September 21, 2007.

REFERENCES
Agrawal, G.K., Rakwal, R., and Jwa, N.S. (2000). Rice (Oryza sativa L.) OsPR1b gene is phytohormonally regulated in close interaction with light signals. Biochem. Biophys. Res. Commun. 278: 290–298.
Azevedo, C., Sadanandom, A., Kitagawa, K., Freialdenhoven, A., Shirasu, K., and Schulze-Lefert, P. (2002). The RAR1 interactor SGT1, an essential component of R gene-triggered disease resistance. Science 295: 2073–2076.

Belkhadir, Y., Nimchuk, Z., Hubert, D.A., Mackey, D., and Dangl, J.L. (2003). Arabidopsis RIN4 is a target of the type III virulence effector AvrRpt2 and modulates RPS2-mediated resistance. Cell 112: 379–389.

Cartwright, D., Langcake, P., Pryce, D.P., Leworthy, D.P., and Ride, J.P. (1977). Chemical activation of host defence mechanisms as a basis for crop protection. Nature 261: 153–156.

Chisholm, S.T., Coaker, G., Day, B., and Staskawicz, B.J. (2000). FLS2: An LRR receptor-like kinase involved in the perception of the bacterial elicitor flagellin in Arabidopsis. Cell 101: 165–175.

Chomela, B.K., and Dangl, J.L. (2001). Plant pathogens and integrated disease resistance. Annu. Rev. Phytopathol. 39: 1–27.

Chou, M.C., and Dangl, J.L. (2001). Innate immunity in plants and animals: Striking similarities and obvious differences. Annu. Rev. Immunol. 19: 589–636.

Cirilli, M., and Martinoia, E. (2001). The rice blast fungus Magnaporthe grisea in barley. Mol. Plant Microbe Interact. 14: 631–640.

Clouse, S.D., and Dangl, J.L. (2001). Directed proteomics identifies a plant-specific protein rap2 involved in programmed cell death. Proc. Natl. Acad. Sci. USA 98: 14633–14638.

Collinge, D.B., Thordal-Christensen, H., and Schulze-Lefert, P. (1994). Nec and Nar-2, two loci required for Mla12-specified race-specific resistance in barley. Plant Cell 6: 118–122.

Cottyn, M., Molk, D., and Corbineau, R. (2000). Arabidopsis RAR1, EDS1 and NPR1/NIM1 like genes are required for N-mediated resistance to tobacco mosaic virus. Plant J. 23: 415–429.

Cox, J.F., and Dangl, J.L. (2001). Plant innate immunity - Direct and indirect recognition of general and specific pathogen-associated molecules. Curr. Opin. Immunol. 16: 119–126.

Cox, M.A., Zhang, K., and Dangl, J.L. (2001). Arabidopsis RAR1 exerts rate-limiting control of R gene-mediated defenses against multiple pathogens. Plant Cell 14: 979–992.

Cui, Y., and Dangl, J.L. (2001). Green gene discovery. Plant Mol. Biol. 42: 179–183.

Dangl, J.L., and Jones, J.D. (2001). Plant pathogens and integrated disease resistance. Annu. Rev. Plant Physiol. Plant Mol. Biol. 48: 251–275.

Dangl, J.L., and Jones, J.D. (2001). Plant innate immunity - Direct and indirect recognition of general and specific pathogen-associated molecules. Curr. Opin. Immunol. 16: 1–8.

Dangl, J.L., and Jones, J.D. (2001). Plant pathogens and integrated disease resistance. Annu. Rev. Plant Physiol. Plant Mol. Biol. 48: 251–275.

Davison, T.S., and Dangl, J.L. (2001). Innate immunity in plants and animals: Striking similarities and obvious differences. Annu. Rev. Immunol. 19: 589–636.

Davison, T.S., and Dangl, J.L. (2001). Innate immunity in plants and animals: Striking similarities and obvious differences. Annu. Rev. Immunol. 19: 589–636.

Davison, T.S., and Dangl, J.L. (2001). Innate immunity in plants and animals: Striking similarities and obvious differences. Annu. Rev. Immunol. 19: 589–636.

Davison, T.S., and Dangl, J.L. (2001). Innate immunity in plants and animals: Striking similarities and obvious differences. Annu. Rev. Immunol. 19: 589–636.
Shang, Y., Li, X., Cui, H., He, P., Thilmony, R., Chintamanani, S., Zwiesler-Vollick, J., Gopalan, S., Tang, X., and Zhou, J.M. (2006). RAR1, a central player in plant immunity, is targeted by *Pseudomonas syringae* effector AvrB. Proc. Natl. Acad. Sci. USA 103: 19200–19205.

Shirasu, K., Lahaye, T., Tan, M.W., Zhou, F., Azevedo, C., and Schulze-Lefert, P. (1999). A novel class of eukaryotic zinc-binding proteins is required for disease resistance signaling in barley and development in *C. elegans*. Cell 99: 355–366.

Shirasu, K., and Schulze-Lefert, P. (2003). Complex formation, promiscuity and multi-functionality: Protein interactions in disease-resistance pathways. Trends Plant Sci. 8: 252–258.

Song, W.Y., Wang, G.L., Chen, L.L., Kim, H.S., Pi, L.Y., Holsten, T., Gardner, J., Wang, B., Zhai, W.X., Zhu, L.H., Fauquet, C., and Ronald, P. (1995). A receptor kinase-like protein encoded by the rice disease resistance gene, Xa21. Science 270: 1804–1806.

Takahashi, A., Casais, C., Ichimura, K., and Shirasu, K. (2003). HSP90 interacts with RAR1 and SGT1 and is essential for RPS2-mediated disease resistance in Arabidopsis. Proc. Natl. Acad. Sci. USA 100: 11777–11782.

Takahashi, A., Kawasaki, T., Henmi, K., Shi, I.K., Kodama, O., Satoh, H., and Shimamoto, K. (1999). Lesion mimic mutants of rice with alterations in early signaling events of defense. Plant J. 17: 535–545.

Tornero, P., Merritt, P., Sadanandom, A., Shirasu, K., Innes, R.W., and Dangl, J.L. (2002). RAR1 and NDR1 contribute quantitatively to disease resistance in Arabidopsis, and their relative contributions are dependent on the R gene assayed. Plant Cell 14: 1005–1015.

Yamazaki, M., Tsugawa, H., Miyao, A., Yano, M., Wu, J., Yamamoto, S., Matsumoto, T., Sasaki, T., and Hirochika, H. (2001). The rice retrotransposon Tos17 prefers low-copy-number sequences as integration targets. Mol. Genet. Genomics 265: 336–344.

Zhou, J., Loh, Y.T., Bressan, R.A., and Martin, G.B. (1995). The tomato gene Pti1 encodes a serine/threonine kinase that is phosphorylated by Pto and is involved in the hypersensitive response. Cell 83: 925–935.

Zipfel, C., and Felix, G. (2005). Plants and animals: A different taste for microbes? Curr. Opin. Plant Biol. 8: 353–360.

Os *PtI1a* Negatively Regulates Rice Defense 2951
Rice Pti1a Negatively Regulates RAR1-Dependent Defense Responses
Akira Takahashi, Ganesh Kumar Agrawal, Muneeo Yamazaki, Katsura Onosato, Akio Miyao, Tsutomu Kawasaki, Ko Shimamoto and Hirohiko Hirochika

Plant Cell 2007;19;2940-2951; originally published online September 21, 2007;
DOI 10.1105/tpc.106.047142

This information is current as of July 19, 2018

Supplemental Data /content/suppl/2007/09/13/tpc.106.047142.DC1.html
References This article cites 50 articles, 13 of which can be accessed free at:
/content/19/9/2940.full.html#ref-list-1
Permissions https://www.copyright.com/ccc/opemar.do?sid=pd_hw1532298X&issn=1532298X&WT.mc_id=pd_hw1532298X
CiteTrack Alerts Sign up for CiteTrack Alerts at:
http://www.plantcell.org/cgi/alerts/ctmain
Subscription Information Subscription Information for The Plant Cell and Plant Physiology is available at:
http://www.aspb.org/publications/subscriptions.cfm

© American Society of Plant Biologists
ADVANCING THE SCIENCE OF PLANT BIOLOGY