RaxH/RaxR: A Two-Component Regulatory System in Xanthomonas oryzae pv. oryzae Required for AvrXa21 Activity

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Xanthomonas oryzae pv. oryzae is the causal agent of bacterial leaf blight, one of the most serious diseases in rice. X. oryzae pv. oryzae Philippine race 6 (PR6) strains are unable to establish infection in rice lines expressing the resistance gene Xa21. Although the pathogen-associated molecule that triggers the Xa21-mediated defense response (AvrXa21) is unknown, six rax (required for AvrXa21 activity) genes encoding proteins involved in sulfur metabolism and Type I secretion were recently identified. Here, we report on the identification of two additional rax genes, raxR and raxH, which encode a response regulator and a histidine protein kinase of two-component regulatory systems, respectively. Null mutants of PR6 strain PXO99 that are impaired in either raxR, raxH, or both cause lesions significantly longer and grow to significantly higher levels than does the wild-type strain in Xa21-rice leaves. Both raxR and raxH mutants are complemented to wild-type levels of AvrXa21 activity by introduction of expression vectors carrying raxR and raxH, respectively. These null mutants do not affect AvrXa7 and AvrXa10 activities, as observed in inoculation experiments with Xa7- and Xa10-rice lines. Western blot and raxR:gfp promoter-reporter analyses confirmed RaxR expression in X. oryzae pv. oryzae. The results of promoter-reporter studies also suggest that the previously identified raxSTAB operon is a target for RaxH/RaxR regulation. Characterization of the RaxH/RaxR system provides new opportunities for understanding the specificity of the X. oryzae pv. oryzae-Xa21 interaction and may contribute to the identification of AvrXa21.

Genetic factors governing interactions between pathogens and their host plants determine disease resistance or susceptibility. This genetic pattern is the basis of the “gene-for-gene” hypothesis, which states that plant disease resistance is controlled by a resistance (R) gene that encodes a product that interacts (directly or indirectly) with an effector molecule encoded by a corresponding bacterial avirulence (avr) gene (Flor 1971). The gene-for-gene hypothesis has been shown to accurately describe the interactions between plant species and a large variety of pathogen, including viruses, bacteria, fungi, nematodes, and insects (Dangl and Jones 2001).

Most plant R genes encode presumed intracellular proteins that are predicted to bind intracellular ligands possibly but not necessarily encoded by avr genes (Dangl and Jones 2001). In plant pathogenic bacteria, identification of mutant strains altered in their ability to cause disease and to induce a hypersensitive response (HR) led to the discovery of hrp (hypersensitive response and pathogenicity) genes (Alfano and Collmer 1997; Leach and White 1996; Lindgren et al. 1986). Several hrp genes encode membrane-associated proteins that form part of a Type III secretory pathway that is active during infection of the plant. Over the past several years, a number of bacterial Avr proteins have been identified that require the Type III apparatus for secretion. In contrast to the intracellular model, it appears likely that some R gene products bind pathogen-associated molecules that are present in the extracellular environment. The tomato Cf9 gene product that mediates resistance to the fungal pathogen Cladosporium fulvum (Jones et al. 1994), the flagellin receptor Fls2 (Gomez-Gomez and Boller 2000), and the product of the rice R gene Xa21 that confers resistance to several Xanthomonas oryzae pv. oryzae races (Song et al. 1995) best fit this model.

X. oryzae pv. oryzae is the causal agent of one of the most serious diseases of rice (Oryza sativa L.), bacterial leaf blight, which is responsible for significant reductions of rice yields in tropical, subtropical, and warm temperature regions. Xa21 encodes a presumed receptor kinase (RK), which confers resistance to multiple races of X. oryzae pv. oryzae (Wang et al. 1996). Xa21 belongs to the leucine-rich repeats (LRR) subclass of plant RK and contains an extracellular domain carrying imperfect units of 24-aa LRR (Song et al. 1995). The presumed extracellular LRR domain of Xa21 is responsible for recognition of specific X. oryzae pv. oryzae strains (Wang et al. 1998).

Six rax (required for AvrXa21 activity) genes from PXO99, a X. oryzae pv. oryzae Philippine race 6 (PR6) strain, were recently identified. In contrast to PXO99, which is unable to cause disease in rice lines expressing Xa21, PXO99 strains carrying rax gene mutations lose AvrXa21 activity (da Silva et al. 2004; Shen et al. 2002). Three of these genes, raxA, raxB, and raxC, encode proteins that share similarity with compo-
ments of Type I secretion systems of gram-negative bacteria. RaxA and RaxB show similarity with membrane fusion proteins and inner membrane permeases of the ATP-binding cassette (ABC) superfamily, respectively. RaxC is similar to the Escherichia coli outer membrane protein ToIC (da Silva et al. 2004). Three additional rax genes encode proteins involved in sulfur metabolism. raxP and raxQ encode ATP sulfurylase and adenosine phosphosulfate kinase activities (Shen et al. 2002), whereas raxST shares similarity with mammalian and bacterial sulfotransferases. raxST is linked to raxA and raxB, and results of sequence analyses suggest these genes are part of one operon named raxSTAB (da Silva et al. 2004). One or more presumed AvrXa21 molecules is yet to be identified.

Sequencing downstream of raxSTAB led to the identification of two open reading frames (ORF), raxR and raxH, showing similarity to response regulators (RR) and histidine protein kinases (HK) of two-component regulatory (or signal transduction) systems. Two-component systems are present in both gram-negative and gram-positive bacteria and have been shown to regulate a wide variety of biological processes, including expression of toxins and other proteins related to virulence and pathogenicity (Grebe and Stock 1999; Stock et al. 2000; West and Stock 2001). In lactic acid bacteria, two-component systems regulate synthesis and type I secretion of bacteriocin-like peptides (small peptides, some of them having antimicrobial activity) (Nes and Eijsink 1999). Indeed, the X. oryzae pv. oryzae raxB product is most similar to ABC transporters involved in secretion of bacteriocin-like peptides, and raxC complements an E. coli toIC mutant strain for secretion of the bacteriocin colicin V (da Silva et al. 2004). In light of these findings, we investigated the role of raxR and raxH in the X. oryzae pv. oryzae-Xa21 interaction. In this report, we show that the RaxH/RaxR system is required for wild-type levels of AvrXa21 activity in X. oryzae pv. oryzae.

RESULTS

Sequencing of the X. oryzae pv. oryzae genomic region downstream of raxSTAB.

A 9-kb Sau3AI fragment from clone p10.78 (Table 1), comprising the raxSTAB region from PR6 X. oryzae pv. oryzae PXO99, was used as the probe to search an available cosmids genomic library of the PR6 strain PXO86 (Hopkins et al. 1992). Both strains are avirulent on Xa21 plants and share a very high level of identity. Clone pHMX7-10 carrying a fragment of about 25 kb was selected for subcloning and sequencing. Following restriction with EcoRI and Apal, the resulting fragments were subcloned into pBlueScript. Subclones pSXX312 and pSXX115, containing a 7.194-bp Apal-EcoRI fragment and an approximately 4-kb EcoRI fragment, respectively, were partially sequenced, leading to a sequenced genomic region of about 15.5 kb (Fig. 1). The PXO99 9-kb fragment from p10.78 and the PXO86 sequenced region from pSXX312 (GenBank accession number YJ237537) overlap approximately 100 nt with 100% identity. Sequence analysis lead to the completion of ORF10 (da Silva et al. 2004) and to the identification of at least four additional ORF designated ORF11 through ORF14 (Fig. 1).

ORFs 13 and 14 encode a putative two-component system required for wild-type levels of AvrXa21 activity in X. oryzae pv. oryzae strain PXO99.

The region comprising ORFs 11 to 14 shares a high level of similarity to genomic regions of the recently sequenced xan-thomonads, X. axonopodis pv. citri and X. campestris pv. campestris. ORF11 and ORF12 encode putative proteins with unknown functions. ORF13 and ORF14 encode putative 238- and 430-aa proteins, showing significant similarity to RR and HK of prokaryotic two-component regulatory systems, respectively.

PXO99 null mutants impaired in ORF13 and ORF14 were created by insertional mutagenesis and were designated PXO99R (RR) and PXO99H (HK), respectively. Various inoculation experiments using these strains in comparison with the parental strain PXO99 (AvrXa21+) and the Korean race 1 (K1) strain DY89031 (AvrXa21+) were carried out using the Orzya sativa subsp. japonica cultivars TP309 (susceptible control) and TP309-Xa21 (a transgenic rice line carrying the Xa21 gene) plants. In all experiments, PXO99R and PXO99H showed reduced AvrXa21 activity in comparison with wild-type PXO99 (Table 2). That is, 5 to 6 days after inoculation (dai), TP309-Xa21 leaves inoculated with the mutant strains

Table 1. Bacterial strains used in this study

<table>
<thead>
<tr>
<th>Strain</th>
<th>Relevant characteristics</th>
<th>Source or reference</th>
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<tbody>
<tr>
<td>Escherichia coli</td>
<td></td>
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<tr>
<td>DH10B</td>
<td>F' mcrA Δmorr-hsdRMS-mcrBC Φ80lacZ ΔM15 Δaac74 deor recA1 endA1 araΔ139 marA, leu7697 galU galK λ- rpsl (Sm') nupG λ- tonA</td>
<td>Gibco BRL</td>
</tr>
<tr>
<td>BL21(DE3)</td>
<td>F'ompT hsdS'6 (rK- mQ-) gal dcm (DE3) pLYsS (Cm')</td>
<td>Novagen</td>
</tr>
<tr>
<td>Xanthomonas oryzae pv. oryzae</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PXO99</td>
<td>Philippine race 6 (PR6) strain, having AvrXa21 activity (AvrXa21+), Cpf'</td>
<td>S. H. Choi</td>
</tr>
<tr>
<td>PXO99R</td>
<td>PXO99 raxR::Km, AvrXa21Δ-, Km'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99H</td>
<td>PXO99 raxH::Km, AvrXa21Δ+, Km'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99RH</td>
<td>PXO99 raxR::Ω, raxH::Km, AvrXa21Δ+, Km', Sm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99R+</td>
<td>PXO99 carrying pMLaxR, AvrXa21+, Gm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99R-R*</td>
<td>PXO99R complemented with pMLaxR, AvrXa21+, Km', Gm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99R-ST*</td>
<td>PXO99R carrying pMLaxST, AvrXa21+, Km', Gm'</td>
<td>This study</td>
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<tr>
<td>PXO99H-H*</td>
<td>PXO99H complemented with pMLaxH, AvrXa21+, Km', Gm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99H-R*</td>
<td>PXO99H carrying pMLaxR, AvrXa21+, Km', Gm'</td>
<td>This study</td>
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<tr>
<td>PXO99-a7</td>
<td>PXO99 carrying plasmid pXO29-29, Sm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99-a10</td>
<td>PXO99 carrying plasmid pXO5-15, Sm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99R-a7</td>
<td>PXO99R carrying plasmid pXO29-29, Km', Sm'</td>
<td>This study</td>
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<td>PXO99R-a10</td>
<td>PXO99R carrying plasmid pXO5-15, Km', Sm'</td>
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<tr>
<td>PXO99H-a7</td>
<td>PXO99H carrying plasmid pXO5-15, Km', Sm'</td>
<td>This study</td>
</tr>
<tr>
<td>PXO99H-a10</td>
<td>PXO99H carrying plasmid pXO29-29, Km', Sm'</td>
<td>This study</td>
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a AvrXa21+ causes short lesions (1 to 2 cm) on Xa21 rice lines; AvrXa21-, causes long lesions (15 to 25 cm) on Xa21 rice lines; AvrXa21Δ+, causes intermediate lesions (5 to 10 cm) on Xa21 rice lines
b National Crop Experiment Station, Rural Development Administration, Suwon, South Korea.

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showed lesions of similar lengths as those caused by DY89031, ranging from 3 to 5 cm, and were significantly longer than lesions caused by the xa21-avirulent strain PXO99 (data not shown). While lesions induced by DY89031 continued to extend up to 15 to 25 cm at 14 dai, lesions induced by the mutant strains extended to 5 to 10 cm (Fig. 2; Table 2). Lesions caused by wild-type PXO99 were significantly shorter, with averages from most experiments ranging from 1 to 2 cm (Fig. 2; Table 2).

To better quantify these results, we carried out in planta growth curve experiments. By 4 dai, PXO99R and PXO99H grew to higher levels in TP309-Xa21 as compared with PXO99 (Fig. 3A). Also by this time, the observed growth for the mutant strains was similar to that of DY89031 (Fig. 3A) and correlated with lesion length data (data not shown). At 12 dai, growth of PXO99R and PXO99H in TP309-Xa21 leaves was intermediate to growth of PXO99 and DY89031 (Fig. 3A). While differences in growth of PXO99 in TP309 vs. TP309-Xa21 started to appear 4 dai, PXO99R and PXO99H did not display differences in growth between these rice varieties until 6 dai. All strains were virulent on TP309 plants and grew to equally high levels in these plants (Fig. 3B). An additional growth curve experiment was carried out using the non-transgenic O. sativa subsp. indica cultivars IRBB21 and IR24 (carrying and lacking Xa21, respectively), and similar results for bacterial growth and lesion lengths were obtained (data not shown).

The above findings indicate that ORF13 and ORF14 are required for wild-type levels of AvrXa21 activity; therefore they were named raxR (RR) and raxH (HK), respectively. A PXO99 mutant strain impaired in both raxR and raxH (PXO99RH) also was created. Inoculation experiments showed that this strain did not display decreased AvrXa21 activity as compared with that of the single-mutant strains (Table 2).

**RaxR belongs to the OmpR family of RR and RaxH to the HPK subfamily of HK.**

Based on similarity searches, ORF predictions, and comparative analysis with similar two-component systems, it is likely that raxR and raxH are part of a single operon. BlastP revealed that RaxR and RaxH are highly similar to putative

### Table 2. Summary of inoculation experiments of Xanthomonas oryzae pv. oryzae strains on TP309-Xa21 plants

<table>
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<tbody>
<tr>
<td>1</td>
<td>2.3 ± 0.6</td>
<td>5.6 ± 0.6</td>
<td>5.5 ± 1.2</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>19.6 ± 0.5</td>
</tr>
<tr>
<td>2</td>
<td>1.9 ± 0.9</td>
<td>5.8 ± 1.2</td>
<td>5.9 ± 1.9</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>16.4 ± 3.9</td>
</tr>
<tr>
<td>3</td>
<td>1.8 ± 1.1</td>
<td>6.4 ± 1.9</td>
<td>–</td>
<td>1.1 ± 0.5</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>16.0 ± 2.8</td>
</tr>
<tr>
<td>4</td>
<td>2.1 ± 1.4</td>
<td>7.4 ± 3.3</td>
<td>6.7 ± 2.1</td>
<td>1.1 ± 0.5</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>25.7 ± 2.9</td>
</tr>
<tr>
<td>5</td>
<td>0.9 ± 0.4</td>
<td>9.3 ± 2.7</td>
<td>10.0 ± 3.1</td>
<td>10.1 ± 3.2</td>
<td>–</td>
<td>1.9 ± 1.7</td>
<td>–</td>
<td>2.4 ± 1.7</td>
</tr>
<tr>
<td>6</td>
<td>3.2 ± 0.9</td>
<td>9.9 ± 2.5</td>
<td>9.7 ± 3.2</td>
<td>10.9 ± 3.3</td>
<td>3.0 ± 1.2</td>
<td>3.4 ± 1.6</td>
<td>3.6 ± 2.3</td>
<td>3.9 ± 2.1</td>
</tr>
<tr>
<td>7</td>
<td>1.9 ± 0.8</td>
<td>–</td>
<td>5.4 ± 2.6</td>
<td>–</td>
<td>–</td>
<td>2.1 ± 1.8</td>
<td>–</td>
<td>25.5 ± 4.2</td>
</tr>
</tbody>
</table>

*Lesion lengths (average cm ± standard deviation) were measured 14 days after inoculation (dai). In each experiment, four to eight leaves per plant (three to six plants per treatment) were inoculated. In planta growth curves were carried out in experiment 2. TP309 plants (susceptible controls) were used in all experiments, and all strains were similarly virulent on this variety, with lesion lengths ranging between 15 and 25 cm among the different experiments.

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**Fig. 1.** Schematic representation of a 15.5-kb region from the *Xanthomonas oryzae pv. oryzae* chromosome, containing genes required for AvrXa21 activity. A. Sequences AF389909 and AY237358 (GenBank accession numbers) compose the 9-kb fragment from strain PXO99 in clone p10.78, which confers AvrXa21 activity to strain DY89031. AY237357 was sequenced from subclones of a PXO86 genomic library. Horizontal arrows indicate the predicted coding sequences of the various open reading frames. B. pSKX312 clone containing an *ApdI-EcoRl* 7,194-bp fragment from PXO86. C. pSKX115 clone containing an approximately 4-kb *EcoRl* fragment from PXO86. Dotted lines indicate the *AccI-EcoRl* 775-bp fragment from pSKX312 used to generate pUC115K+, for insertional mutagenesis of *raxH*. Triangles indicate sites of insertion of Km' or Sm' cassettes for insertional mutagenesis. Relevant restriction sites: E = *EcoRl*, Ap = *ApdI*, B = *BglII*, A = *Apol*, Ac = *AccI*, Ag = *AgerI*, K = *KpmI.*
two-component elements of the related *X. axonopodis* pv. *citri*, and *X. campestris* pv. *campestris* RaxR shows 90 and 87% identity to NP_641557 (*X. axonopodis* pv. *citri*) and NP_636495 (*X. campestris* pv. *campestris*), respectively. RaxR also shows high similarity (63 to 66%) to the putative RR NP_638454 and NP_636170 from *X. campestris* pv. *campestris* and to NP_643558 and NP_641186 from *X. axonopodis* pv. *citri*. Similarly, RaxH is 94 and 85% identical to NP_641558 from *X. axonopodis* pv. *citri* and NP_636496 from *X. campestris* pv. *campestris*, respectively. It also shares similarity (46 to 49%) with the putative HK NP_638453 and NP_636171 from *X. campestris* pv. *campestris* and to NP_643557 and NP_641187 from *X. axonopodis* pv. *citri*.

Among characterized two-component systems, the one giving the highest homology scores for the RaxH/RaxR system is the ColS/ColR system of the plant-growth-promoting rhizobacterium *Pseudomonas fluorescens*, which is involved in bacterial root colonization (Dekkers et al. 1998); RaxR and ColR (CAA70931) share 69% similarity and 51% identity, whereas the levels of similarity and identity for RaxH and ColS (CAA70932) are 48 and 30%, respectively. Other two-component systems showing similarity to RaxH/RaxR (48 to 53% for RaxR; 45 to 50% for partial length of RaxH, respectively) are the *Burkholderia pseudomallei* IrlR/IrlS, which is involved in invasion of eukaryotic cells and heavy-metal resistance (Jones et al. 1997), and the *Salmonella typhimurium* BasR/BasS (or PmrA/PmrB), for which a role in virulence has been attributed (Roland et al. 1993).

Sequence analysis indicates that RaxR belongs to the well-studied OmpR family of RR. For instance, RaxR shows 44% similarity with *E. coli* OmpR. PFAM analysis revealed the presence of typical N-terminal RR receiver and C-terminal transcriptional regulatory (effector) domains in RaxR. RaxH has typical histidine kinase A (dimerization and phosphoacceptor) and HATPase_c (ATP-binding kinase) domains in the C-terminus. In addition, a HAMP domain (histidine kinases, adenyl cyclases, methyl binding proteins and phosphatases) with unknown function is found in RaxH. A SMART search also revealed the presence of two transmembrane segments, suggesting the presumed RaxH is a typical transmembrane HK. Based on multiple sequence alignment of 348 kinase domains, Grebe and Stock (1999) designated 11 HK subfamilies, named HPK1 to HPK11. According to homology analysis, RaxH belongs to the HPK5 subfamily that also contains ColS and EnvZ, the cognitive HK of ColR and OmpR, respectively.

**PXO99R and PXO99H are complemented to wild-type levels of AvrXa21 activity by pML122 overexpressing raxR and raxH, respectively.**

The *raxR* and *raxH* coding sequences from PXO99 were cloned into the expression vector pML122, generating pMLraxR and pMLraxH, respectively. Sequencing of these genes from PXO99 revealed 100% identity with the corresponding genes from PXO86. PXO99R (*raxR* mutant) carrying pMLraxR displayed lesion lengths similar to wild-type PXO99 in TP309-Xa21 leaves. Similarly, PXO99H (*raxH* mutant) was complemented for AvrXa21 activity by pMLraxH (Table 2).

Interestingly, PXO99H carrying pMLraxR was also shown to have an increased AvrXa21 activity in comparison with PXO99H, as observed by the reduced lesion lengths in TP309-Xa21 leaves induced by this strain (Table 2). Complementation of the *raxH* mutation by pMLraxR not only confirmed RaxR overexpression in planta but also indicates that at least part of the overexpressed RR protein is activated in the absence of the HK.

**The RaxH/RaxR system does not affect AvrXa7 and AvrXa10 activities.**

The *X. oryzae* pv. *oryzae-Xa21* interaction is unique in that it requires a functional bacterial type I secretion system and likely involves extracellular recognition of AvrXa21 by the Xa21 gene product. Therefore, we investigated whether the RaxH/RaxR system is relevant for activity of other *X. oryzae* pv. *oryzae* avr genes, avrXa7 and avrXa10. These genes are presumed to encode intracellular effectors delivered into the plant cell via a type III secretion system (Yang et al. 2000; Zhu et al. 2000). Since Xa7 and Xa10 rice lines are susceptible to PXO99, plasmids pXO5-15 and pXO29-29 expressing avrXa10 and avrXa7 from strain PXO86 (Hopkins et al. 1992), respectively, were introduced into wild-type PXO99 and the mutant strains PXO99R and PXO99H.

As expected, PXO99, PXO99R, and PXO99H were virulent on Xa7 and Xa10 lines (Fig. 4), and all strains were virulent on IR24 plants (susceptible control; data not shown). PXO99R and PXO99H carrying avrXa7 or avrXa10 caused lesions with lengths similar to those caused by PXO99 expressing the corresponding avr gene in Xa7 or Xa10 plants (Fig. 4). In other words, the RaxH/RaxR system has no effect on AvrXa7 and AvrXa10 activities.

**Expression and purification of recombinant RaxR from *E. coli*.**

The *raxR* coding sequence from PXO99 was cloned into pET-15b, to create recombinant RaxR fused to a histidine tag (His 6× tag) in its N-terminal. The resulting vector, pETraxR, was transformed into *E. coli* BL21(DE3)pLysS, in which overexpression of recombinant His-RaxR protein was induced by the addition of isopropyl-β-d-thiogalactoside (IPTG). The calculated molecular mass for the recombinant His-RaxR is 2.88314 kDa, thus coinciding with the mobility observed in sodium dodecyl sulfate-polyacrylamide gel electrophoresis.
(SDS-PAGE) (data not shown) and in Western blots using anti-His 6x monoclonal antibodies (Fig. 5A). The double-band pattern following induction with IPTG is frequently observed for highly overexpressed recombinant proteins and seems to be the result of partial premature translation termination or protein degradation (The QIAexpressionist, a handbook for high-level expression and purification of 6X His-tagged proteins, Qiagen, Valencia, CA, U.S.A.).

For production of anti-RaxR antibodies, the recombinant protein was purified under denaturative conditions, using Ni²⁺-NTA columns according to manufacturer’s procedures (The QIAexpressionist; data not shown).

**X. oryzae pv. oryzae** expresses wild-type RaxR in culture.

An XbaI-HindIII fragment from pETrAxR containing the His-RaxR coding sequence was cloned into pML122 to generate pMLrAxR (the same construct was used for complementation of PXO99R, as previously described). pMLrAxR was introduced into PXO99, generating PXO99-R*. Overexpression of His-RaxR in this strain was clearly observed by comparing its protein profiles with those of wild-type PXO99 and PXO99R (raxR mutant strain) grown in different media (nutrient broth [NB], M9 minimal medium, and peptone sucrose) in SDS-PAGE (data not shown). These results were confirmed by Western blot analyses using anti-His 6x and anti-RaxR antibodies (Fig. 5). Note that, for detection with anti-RaxR antibodies (Fig. 5B), the overexpressed signal observed in PXO99-R* extracts was obtained from a loaded sample containing significantly less total protein as compared with that in PXO99 and PXO99R samples. His-RaxR was detected at a similar intensity in extracts of M9 minimal medium- and peptone sucrose–grown PXO99-R* cells (data not shown).

Western blots with anti-RaxR antibodies led to the detection of a band with the expected size, the same as that of wild-type RaxR (2.685682 kDa) in protein extracts of NB-grown PXO99 cells (Fig. 5B). Similar bands were also detected in protein extracts of PXO99 grown in M9 minimal medium and peptone sucrose broth, although with a lower intensity than in NB (data not shown). The results discussed above suggest that raxR is constitutively expressed in PXO99 cultures.

To verify the above results, we created a raxR promoter-reporter construct using the promoterless-gfp (green fluorescent protein)-containing vector pPROBE-NT, for which efficiency as a promoter-probe vector was recently demonstrated for other plant pathogens, such as *Erwinia herbicola* and *Pseudomonas syringae* (Miller et al. 2000). PXO99 cells carrying pPROBE-NT-pR1 (containing a 381-bp fragment upstream of the raxR coding sequence fused to promoterless-gfp) were grown for 24 and 48 h in NB (supplied with kanamycin [Km]) and compared for GFP expression with wild-type PXO99 as well as with PXO99 carrying “empty” pPROBE-NT. A pPROBE-NT construct, in which a DNA region located within the raxR coding sequence was fused to the promoterless-gfp, was used as an additional negative control. PXO99 cells carrying pPROBE-NT-pR1 gave significantly higher fluorescence values after 24 and 48 h of growth, as compared with the other strains, which did not show significant differences among each other (data not shown). Observation of bacterial cultures using a confocal microscope correlated with these results, since fluorescence was only observed in PXO99 cells carrying pPROBE-NT-pR1. These findings support Western blot results showing that RaxR is expressed in culture.

PXO99 carrying the above constructs were also used to inoculate 3- to 4-cm pieces of TP309-Xa21 leaves. The leaves were inoculated by infiltration of bacterial suspensions (10⁷ CFU per ml of water) through the main vein and were incubated at 30°C for 5 days in petri dishes containing a layer of wet filter paper. In bacteria grown in culture, only cells carrying pPROBE-NT-pR1 were fluorescent (data not shown), thus supporting results from inoculation experiments in which the RaxH/RaxR system is expressed in planta.

**The raxR mutation affects expression of the raxSTAB promoter in PXO99.**

Due to the chromosomal proximity of raxSTAB and raxRH, as well as the apparent uniqueness of the raxSTAB region in *X. oryzae pv. oryzae* strains in A, TP309-Xa21- and B, TP309-inoculated leaves. PXO99 (■, solid lines), PXO99R (●, dashed lines), PXO99H (▲, dashed lines), and DY89031 (●, solid lines). Bacteria were extracted from leaves every two days, and bacterial concentration was determined by serial dilution counts. Each value represents averages ± standard deviation of three sampled leaves per treatment.
oryzae pv. oryzae (da Silva et al. 2004), we tested whether the *raxSTAB* operon is a target for RaxH/RaxR regulation. Preliminary observations indicated that expression of the *raxSTAB* operon in culture occurs at very low levels. Indeed, we were not able to detect *raxST*, *raxA*, or *raxB* RNA in Northern blot experiments in either PXO99R or PXO99 (data not shown). This could be due to either low expression of these genes, RNA instability, or both, known to be important limiting factors for detection of bacterial RNA.

As an alternative to Northern blot analysis, a 309-bp DNA fragment upstream of the *raxST* coding sequence, presumably containing the *raxSTAB* promoter (named pST), was cloned into pPROBE-AT upstream of the promoterless-*gfp* for promoter-reporter studies. The resulting plasmid, pPROBE-AT-pST was electroporated into PXO99 and PXO99R cells. pPROBE-AT-pNm, carrying a pML122 fragment containing the pNm promoter (from *nptII*; constitutively expressed in *X. oryzae pv. oryzae*) was also introduced into the wild-type and *raxR* mutant strains, to allow determination of relative activity of the *pST* promoter. As a negative control, pPROBE-AT (carrying promoterless-*gfp*) was also introduced in these strains. Similar values of fluorescence were obtained when comparing PXO99 and PXO99R for the pNm-*gfp* fusion and pPROBE-AT for the negative control. No differences were observed in growth rate between the different strains (data not shown). Figure 6 shows that the *raxR* mutation in PXO99R clearly suppresses the activity of pST. Interestingly, in PXO99R, pPROBE-AT-pST gave higher fluorescence values than pPROBE-AT (data not shown), suggesting the occurrence of residual pST activity in the background of the *raxR* mutation.

Inoculation experiments showed that PXO99R overexpressing *raxST* (cloned into pML122) induced shorter lesions in TP309-Xa21 leaves as compared with PXO99R (Table 2). This finding is in agreement with the results from promoter expression analysis and further supports that the *raxSTAB* operon is likely a target of RaxH/RaxR regulation.

**Fig. 4.** Lesion lengths in leaves of IRBB7 and IRBB10 rice plants inoculated with strains PXO99, PXO99R, and PXO99H, and the same strains expressing either *avrXa7* (a7) or *avrXa10* (a10), 14 days after inoculation. Three or four plants (four to six leaves per plant) were inoculated with each strain. Bars represent average ± standard deviation from one of two independent experiments with comparable results. No differences in lesion lengths were found among the strains in IRBB24 plants (data not shown).

**Fig. 5.** Expression of RaxR in *Escherichia coli* and in *Xanthomonas oryzae pv. oryzae*. **A.** Western blot analysis of various *X. oryzae pv. oryzae* strains, and *E. coli* BL21(DE3)pLysS carrying pETRaxR. *X. oryzae pv. oryzae* strains were grown in nutrient broth (NB) (with gentamycin for PXO99 carrying pMlaxR) for 24 h. Detection with anti-His 6x monoclonal antibodies (1:5,000; Sigma) and horseradish peroxidase-anti-mouse IgG (1:10,000; Jackson ImmunoResearch Laboratories) as primary and secondary antibodies, respectively. Lane 1, BL21-pETRaxR, 1 mM IPTG, lane 2, BL21-pETRaxR, no IPTG, lane 3, PXO99, and lane 4, PXO99-R* (PXO99 carrying pMLRaxR). Membranes were exposed for 15 s to an X-ray film. **B.** Detection of RaxR from various *X. oryzae pv. oryzae* strains grown in nutrient broth for 24 h, using anti-RaxR polyclonal antibodies (1:3,000) and HRP-anti-rabbit IgG (1:10,000, Cappel Laboratories) as primary and secondary antibodies, respectively. Lane 1, PXO99, lane 2, PXO99R (*raxR* mutant), and lane 3, PXO99-R*. For PXO99 and PXO99R extracts, 15 µl were loaded, and for PXO99-R*, 3 µl were loaded. Membranes were exposed for 1 min to an X-ray film.

**Fig. 6.** Relative activity of the *raxSTAB* promoter (pST) in wild-type (PXO99) and *raxR* mutant (PXO99R) strains. Cells carrying pPROBE-AT-pST were grown in nutrient broth with ampicillin for 24 h to the middle exponential phase. After two washings with PBS (20 mM NaHPO4, 150 mM NaCl, pH 8.0), cells were resuspended with the same buffer to an optical density at 600 nm = 1.0. Fluorescence was measured at 490- and 510-nm excitation and emission wavelengths, respectively. pST relative activities in PXO99 and PXO99R were calculated relative to PXO99 and PXO99R cells expressing pPROBE-AT-pNm, respectively (pNm = 100%). Results from three independent experiments are shown. In each experiment, data represent average ± standard deviation of three independent replicates (cultures).
**DISCUSSION**

The rice Xa21 resistance gene encodes a RR with a presumed extracellular domain that is responsible for race-specific recognition of *X. oryzae pv. oryzae* (Song et al. 1995). Recent characterization of six *X. oryzae pv. oryzae* rax genes indicate that type I secretion (raxA, raxB, and raxC) and sulfation (raxST, raxP, and raxQ) are required for Xa21-mediated recognition of *X. oryzae pv. oryzae* (da Silva et al. 2004; Shen et al. 2002).

In the present study, two ORF encoding proteins with similarity to RR and HK of the OmpR/EmrZ class of bacterial two-component regulatory systems were identified downstream of the raxSTAB genomic region. Because several two-component regulatory systems regulate virulence and pathogenicity (Grebe and Stock 1999; Stock et al. 2000; West and Stock 2001), we investigated whether this presumed two-component system is involved in the *X. oryzae pv. oryzae-Xa21* interaction.

Null mutant strains for the RR and HK ORF were generated by insertional inactivation in strain PX099 carrying AvrXa21 activity. Inoculation experiments showed that these ORF are required for wild-type levels of AvrXa21 activity, and the genes were consequently named raxR and raxH. Lesion length analysis and in planta growth curve experiments showed that the PXO99 raxR and raxH mutants possess an intermediate phenotype between wild-type strains PXO99 and DY89031, the latter lacking AvrXa21 activity. Also, both mutant strains induce shorter lesions in Xa21 leaves as compared with those of some of the other PXO99 rax mutants, such as the raxST knockout strain (data not shown). A double-mutant strain impaired in both raxR and raxH did not display a further decrease in AvrXa21 activity. These results suggest that the raxH and raxR mutant strains do elicit a defense response in Xa21 plants but that this response is delayed or weaker, or both, as compared with that in wild-type PXO99.

A possible explanation for the intermediate phenotype caused by inactivation of the RaxH/RaxR system may be that other factors, including additional two-component systems, are involved in regulation of AvrXa21 activity and that inactivation of one of these systems is not sufficient for its complete inhibition. As an example of redundant two-component regulation, two-component regulatory systems (named System 1 and System 2) that regulate virulence of the gram-negative human pathogen *Vibrio cholerae* were recently identified (Miller et al. 2002). Moreover, using the *Vibrio harveyi* luciferase (lux) operon as a heterologous reporter, Miller and associates (2002) showed that a *V. cholerae* double mutant, inactivated in both System 1 and System 2, still retains density-dependent lux expression, thus suggesting the involvement of at least one additional sensory circuit. Indeed, sequencing of the genomes of *X. axonopodis pv. citri* and *X. campestris pv. campestris* revealed the presence of more than 30 two-component regulatory systems in these species, with one being identical and two highly similar (in each species) to the *X. oryzae pv. oryzae* RaxH/RaxR system. It is not yet known if the *X. oryzae pv. oryzae* genome encodes two-component systems similar to RaxH/RaxR, as the *X. oryzae pv. oryzae* genome has not yet been published. If present in *X. oryzae pv. oryzae*, these additional systems could also play a role in AvrXa21 regulation. In addition, we cannot discount the possibility that the intermediate phenotype is due to low-level constitutive expression of *rax* genes regulated by the RaxH/RaxR system.

In the absence of the HK or of the required environmental signal, some RR are able to catalyze phosphoryl transfer from small molecules such as acetyl phosphate; moreover, this kind of RR phosphorylation seems to be more biologically relevant than phosphoryl transfer from noncognitive HK (Lukat et al. 1992; McCleary et al. 1993). Inoculation experiments indicated that the raxH mutation is complemented by overexpression of raxR. This finding has an important implication for future studies on gene regulation by the RaxH/RaxR system, since PXO99 cells overexpressing raxR could be used for expression studies in the absence of the RaxH-activating environmental signal (which is still unknown). Recently, Noël and associates (2001) compared cDNA-AFLP profiles of a *X. campestris pv. vesicatoria* wild-type strain and a strain expressing a mutated form of HrpG (a RR that belongs to the OmpR family) that leads to constitutive expression of *hrpg* genes in normally noninducing medium. Using this approach, 30 *hrpg*-induced and five *hrpg*-repressed cDNA fragments were successfully detected.

An important question regarding the role of the RaxH/RaxR system in regulation of avirulence is if this two-component system is relevant for expression of other *X. oryzae pv. oryzae* avr genes. The present study showed that inactivation of raxR or raxH does not affect AvrXa7 and AvrXa10 activities. In contrast to avrXa7 and avrXa10 that encode typical Avr effectors, which are likely delivered by a type III secretion system and recognized by plant intracellular receptors (Yang et al. 2000; Zhu et al. 2000), the recognition of AvrXa21 is predicted to occur in the extracellular environment. The fact that the RaxH/RaxR system affects AvrXa21 but not AvrXa7 and AvrXa10 activities emphasizes the unique aspects of the *X. oryzae pv. oryzae-Xa21* system within the context of avr–R gene interactions.

The investigation into how the RaxH/RaxR system controls AvrXa21 activity is in its infancy. However in this study, we show evidence from promoter-reporter studies and inoculation experiments suggesting that the raxSTAB locus is a target for RaxH/RaxR regulation. Moreover, the residual pST activity observed in the background of the raxR mutation is in agreement with the intermediate AvrXa21 activity observed for the raxR/raxH mutant strains in inoculation experiments. Whether RaxR up-regulates the raxSTAB locus directly or through an intermediate transcription factor and whether RaxR is involved in regulation of other rax genes are questions that remain to be answered. It is particularly intriguing that both *X. campestris pv. campestris* and *X. axonopodis pv. citri* possess orthologs for *raxB* and *raxH* (and a *RaxST* locus is a target for RaxH/RaxR regulation). These observations suggest that RaxH/RaxR system may be involved in regulation of a broad spectrum of genes, not all necessarily related to AvrXa21 activity. Identification of additional RaxH/RaxR targets in *X. oryzae pv. oryzae* as well as in the related xanthomonads is, therefore, of interest.

To date, we have isolated eight *X. oryzae pv. oryzae* genes required for AvrXa21 activity. These genes fall into three classes: type I secretion, sulfur metabolism, and two-component regulation. Further characterization of the interaction among the encoded products of these genes and identification of new AvrXa21 determinants will contribute to the understanding of the *X. oryzae pv. oryzae-Xa21* interaction as well as possible isolation of AvrXa21 itself. The unique aspects of the *X. oryzae pv. oryzae-Xa21* system suggest that AvrXa21 may represent an entirely new class of bacterial signaling molecules.

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and growth conditions.**

Bacterial strains and plasmids used in this work are described in Tables 1 and 3, respectively. *X. oryzae pv. oryzae* strains were grown in peptone sucrose media (Tsuchiya et al. 1998); molecular plant-microbe interactions.
1982), NB (Difco Laboratories, Detroit), or M9 minimal medium at 28°C. For solid media, 16 g of agar per liter was added, E. coli strains were cultured in Luria-Bertani medium at 37°C or as otherwise stated. Antibiotics used in this study were: Km, 50 µg ml⁻¹; streptomycin (Sm), 50 µg ml⁻¹; ampicillin (Ap), 100 µg ml⁻¹; cephalaxin, 20 µg ml⁻¹; chloramphenicol, 34 µg ml⁻¹; and gentamicin (Gm), 30 µg ml⁻¹ for E. coli and 15 µg ml⁻¹ for X. oryzae pv. oryzae.

Molecular techniques.

Plasmid DNA preparations, restriction enzyme digestions, and other routine DNA manipulations were performed using standard procedures (Sambrook et al. 1989). Genomic DNA from X. oryzae pv. oryzae was prepared according to Ausubel and associates (1994). For Southern blot hybridization, [³²P]-dCTP (NEN Life Science Products, Boston) was used to label DNA probes with a random labeling kit (Amer sham Life Science, Arlington Height, IL, U.S.A.). DNA sequencing of DNA fragments cloned in pUC18 or pBluescript was performed by the dyeodeoxy chain termination method, using an automated sequencer (Model 400 I; Li-Cor, Lincoln, NE, U.S.A.). DNA sequences were analyzed and assembled using Sequencher (Gene Codes Corp., Ann Arbor, MI, U.S.A.). DNA and protein homology analysis and searches for ORF were performed with the Blast algorithm (Altschul et al. 1997) and ORF Finder, respectively, through the National Center for Biotechnology Information. Domain analyses were carried out using PFAM and SMART software. Multiple alignments were carried out using ClustalW (Thompson et al. 1994).

Construction of razR and razH null mutants by insertional mutagenesis.

To generate a PXO99 razR mutant (PXO99R), an Apol 1,021-bp fragment from pSKX312 (Fig. 1) was subcloned into pUC18, and a Km resistance cassette was inserted into the

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<td><strong>Plasmid</strong></td>
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Cloning of raxR and raxH for complementation and overexpression in X. oryzae pv. oryzae.

The raxR and raxH coding sequences from PXO99 were PCR-amplified, using PXO99 genomic DNA as template and the following primers: RR_F (forward primer), 5'-CATATGCGCCTGCTTGATCAGTTG-3' (NdeI site shown in bold), and RR_R (reverse primer), 5'-GGATCCATGATACCCTGTGCTGCTGTC-3' (BamHI shown in bold) for raxR, and HK_FE (forward), 5'-GCTAGTACCGCATTGCTTACAACGAGGCCCGCTTTG-3' (KpnI and NdeI shown in bold), and HK_R2 (reverse), 5'-CGTACAGGCTTCCAGGCGCCGGCGCTAG-3' (BamHI shown in bold) for raxH. The PCR reactions were run in 1% agarose gels, and the 726-bp and 318-bp products were excised and purified. The raxR product was cloned into the pCR2.1 vector using the Original TA cloning kit (Invitrogen Corp., Carlsbad, CA, U.S.A.) to create pCRraxR. The raxH product was treated with KpnI and BamHI and was cloned into KpnI-BamHI cut pUC18 to create pUCraxH. Following verification by sequencing, the raxR and raxH fragments were isolated using NdeI and BamHI, were purified, and were ligated into NdeI-BamHI cut pET-15b, to generate pETraxR and pETraxH, respectively. Using this strategy, a tag of six sequential copies of histidine (His 6× tag) was fused to the N-terminus of the coding sequences. The fused fragments were excised from these constructs using XbaI and HindIII and were cloned into the broad-range expression vector pML122 treated with the same restriction enzymes. In the resulting constructs, pMLraxR and pMLraxH, expression of the recombinant His 6×-tagged proteins is under control of the pNm promotor (npII, neomycin resistance), which is constitutively expressed in X. oryzae pv. oryzae. pMLraxR was electroporated into PXO99, PXO99R, and PXO99H. pMLraxH was electroporated into PXO99H. Transformants were selected by Gm′, and plasmid insertion was confirmed by mini-preps and Western blots.

Expression of raxR in X. oryzae pv. oryzae cultures.

For detection of RaxR in culture, X. oryzae pv. oryzae strains were grown for 24 or 48 h in 2 ml of NB, peptone sucrose broth, or M9 minimal medium with appropriate antibiotics at 28°C. Cells were centrifuged (5,000 x g, 10 min, 4°C), and pellets were resuspended in 200 μl of 1× SDS-PAGE loading buffer. Following a brief sonication (15 s in ice), samples were incubated at 90°C for 2 min and were stored at −20°C until used for SDS-PAGE and Western blotting. All experiments were carried out twice with comparable results.

Protein electrophoresis and Western blotting.

Proteins were resolved in SDS-PAGE (with 12% polyacrylamide in the separating gel) and were stained with R250 Coomassie blue or were transferred to Hybond ECL nitrocellulose membranes (Amersham Pharmacia Biotech, Freiburg, Germany) by standard procedures (Sambrook et al. 1989). Membranes were incubated overnight in blocking solution consisting of 5% (wt/vol) skimmed milk powder in T-TBS (10 mM Tris-HCl, pH 8.0, containing 150 mM NaCl and 0.1% [vol/vol] Tween 20). The membranes were then incubated in the presence of anti-His 6× monoclonal antibodies (Sigma) or anti-RaxR polyclonal antibodies at a dilution of 1:5,000 and 1:3,000, respectively, in T-TBS for 1 h at room temperature. Following three 10-min washes in T-TBS, the membranes were incubated for 1 h at room temperature in T-
ACKNOWLEDGMENTS

PXO99R carrying pPROBE-NT or pPROBE-AT were used according to Miller and associates (2000). Cells were also antibiotics and fluorescence was assessed using an RF-1501

gfp

pNm were also electroporated into PXO99R. To determine verification by restriction assays, all plasmids were elec-

tronic

pt

px

resulting transformants were selected on Km

 promoter-reporter constructs.

to clone the region containing the

raxR promoter (pR1), a 381-bp fragment upstream of the

raxR putative translation start codon was PCR-amplified, using pSKX312 as template and the following primers: PR1_F (forward primer), 5'-CGCGGATCCACTGACGGGCTTGACCAT3'- and PR1_R (reverse primer), 5'-CCGGAATTCTTGGGGCCGGCCCGGCAAGCA-3' (BamHI and EcoRI sites, respectively, are shown in bold). To clone the pST region, a 309-bp fragment upstream of the

raxST coding sequence was PCR-amplified, using p10.78 as template and the following primers: PST_F (forward), 5'-CGGATCCGCGTTCATCTGC-3' and PST_R (reverse), 5'-CCGATTCCACCAAGCGTGCATCGT-3' (BamHI and KpnI sites, respectively, are shown in bold). The pR1 and pST PCR products were purified and cloned into the promoterless-

gfp

vectors pPROBE-NT (for pR1) and pPROBE-AT (for pST) treated with appropriate restriction enzymes, to generate pPROBE-NT-pR1 and pPROBE-AT-pST, respectively. A 212-bp fragment containing the

pNm promoter (from nptII) was obtained by treatment of pML122 with SaII and XbaI and was cloned into pPROBE-AT cut with the same enzymes to create pPROBE-AT-pNm. The constructs were electroporated into

E. coli

DH10B, and the resulting transformants were selected on Km or Ap. Following verification by restriction assays, all plasmids were elec-

troporated into PXO999_pPROBE-AT-pST and pPROBE-AT-pNm were also electroporated into PXO999R. To determine

gfp

expression, cells were grown in NB with appropriate antibiotics and fluorescence was assessed using an RF-1501 Spectrofluorophotometer (Shimadzu Co., Kyoto, Japan), ac-

ccording to Miller and associates (2000). Cells were also observed in a Zeiss LSM 5 Pascal confocal microscope (Carl Zeiss, Inc., Thornwood, NY, U.S.A.), using standard settings for GFP detection (excitation and emission wavelengths of 488 and 505 nm, respectively). Wild-type PXO99 and PXO999R carrying pPROBE-NT or pPROBE-AT were used as negative controls.

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X. oryzae pv. oryzae

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**AUTHOR-RECOMMENDED INTERNET RESOURCES**

Xanthomonas Genome Projects website: genoma4.iq.usp.br/xanthomonas


Washington University in St. Louis Pfam protein search: pfam.wustl.edu/hmmsearch.shtml

SMART database: smart.embl-heidelberg.de