Active defence responses associated with non-host resistance of *Arabidopsis thaliana* to the oomycete pathogen *Phytophthora infestans*

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SUMMARY

The molecular basis of non-host resistance, or species-specific resistance, remains one of the major unknowns in the study of plantmicrobe interactions. In this paper, we describe the characterization of a non-host pathosystem involving the model plant Arabidopsis thaliana and the economically important and destructive oomycete pathogen Phytophthora infestans. Cytological investigations into the early stages of this interaction revealed the germination of P. infestans cysts on Arabidopsis leaves, direct penetration of epidermal cells, formation of infection vesicles and occasionally secondary hyphae, followed by a typical hypersensitive response. P. infestans biomass dynamics during infection of Arabidopsis was monitored using kinetic PCR, revealing an increase in biomass during the first 24 h after inoculation, followed by a decrease in the later stages. Transgenic reporter lines and RNA blot analyses were used to characterize the defence responses induced following P. infestans infection. Significant induction of PDF1.2 was observed at 48 h after inoculation, whereas elevated levels of PR gene expression were detected three days after inoculation. To further characterize this defence response, DNA microarray analyses were carried out to determine the expression profiles for c. 11 000 Arabidopsis cDNAs 16 h after infection. These analyses revealed a significant overlap between Arabidopsis non-host response and other defencerelated treatments described in the literature. In particular, non-host response to P. infestans was clearly associated with activation of the jasmonate pathway. The described Arabidopsis-P. infestans pathosystem offers excellent prospects for improving our understanding of non-host resistance.

INTRODUCTION

Plants are challenged by numerous pathogens throughout their life cycles and yet are able to fend off most infections. Indeed, in interactions between plants and microbial pathogens, resistance is the rule and disease the exception. This phenomenon is known as non-host resistance or species-specific resistance and is thought to explain why a pathogen can cause disease in particular plant species but not in others. Understanding the molecular basis of non-host resistance remains one of the elusive guests in the study of plant-microbe interactions. Pre-formed barriers and compounds such as saponins are ubiquitous in plants and play an important role in non-host resistance to filamentous fungi (Morrissey and Osbourn, 1999; Osbourn, 1996). However, most contemporary models of non-host resistance evoke a complex overlay of specific resistance and nonspecific defence responses (Gomez-Gomez and Boller, 2002; Heath, 2000; Kamoun et al., 1999; Kamoun, 2001; Nurnberger and Brunner, 2002). Specific resistance has been extensively studied in host pathosystems and typically follows Flor's gene-for-gene model. In this model, resistance is determined by the simultaneous expression of a pathogen avirulence (Avr) gene with the corresponding plant resistance (R) gene, leading to the hypersensitive response (HR), a general defence response of plants that includes apoptotic cell death (Dangl and Jones, 1998; Flor, 1971; Staskawicz et al., 1995). The extent to which the gene-for-gene model can be expanded to non-host interactions remains unclear. However, we and others have speculated that in many pathosystems non-host resistance can be explained by the occurrence of an arsenal of *R* genes that recognize multiple or essential Avr genes (Heath, 2000; Kamoun et al., 1998; Kamoun et al., 1999; Kamoun, 2001; Staskawicz et al., 1995).

The oomycetes represent a diverse and phylogenetically unique branch of eukaryotic microbes that includes many important pathogens of plants (Baldauf *et al.*, 2000; Margulis and Schwartz, 2000; Sogin and Silberman, 1998). The most notorious

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oomycetes are *Phytophthora* species, arguably the most devastating pathogens of dicot plants (Erwin and Ribeiro, 1996; Kamoun, 2000; Kamoun, 2003). For example, *Phytophthora infestans* causes late blight, a devastating disease that results in multibillion-dollar losses in potato and tomato production (Fry and Goodwin, 1997a,b; Garelik, 2002; Smart and Fry, 2001). Most plants, such as weeds and various crops, are resistant to *P. infestans*, and grow unimpaired in or next to fields with a high incidence of late blight (Colon *et al.*, 1992; Kamoun *et al.*, 1999). Understanding the molecular basis of non-host resistance to *P. infestans* will provide insight into a key molecular process and will offer novel perspectives for engineering durable late blight resistance in crop plants.

A first insight into the basis of non-host resistance to Phytophthora infestans came through detailed cytological analyses. Microscope observations revealed penetration of epidermal cells by P. infestans in all examined interactions, including those with plant species unrelated to the solanaceous hosts (Gross et al., 1993; Kamoun et al., 1998; Kamoun et al., 1999; Naton et al., 1996; Schmelzer et al., 1995; Vleeshouwers et al., 2000). Fully resistant plants, such as the non-hosts Solanum nigrum, parsley and tobacco display a typical localized HR at all infection sites (Colon et al., 1992; Kamoun et al., 1998, 1999; Naton et al., 1996; Schmelzer et al., 1995; Vleeshouwers et al., 2000). The HR can be highly localized to a single epidermal cell or can affect a group of cells surrounding the penetrating hyphae, depending on the interaction examined (Kamoun et al., 1998; Vleeshouwers et al., 2000). The view that has emerged from these studies is that the HR, perhaps mediated by R genes, is associated with all known forms of genetic resistance to P. infestans including nonhost resistance (Kamoun et al., 1999; Kamoun, 2001).

Some of the Phytophthora molecules that trigger the HR or other defence responses in non-host plants are known. Speciesspecific elicitors have been described in P. infestans and other Phytophthora species and can trigger defence responses in nonhost plants. For example, an extracellular transglutaminase that is conserved in *P. infestans* and other *Phytophthora* species induces defence responses in the non-host parsley (Brunner et al., 2002; Nurnberger and Brunner, 2002). Members of the INF elicitin family induce the HR and related biochemical changes specifically in Nicotiana (Kamoun et al., 1997, 1998; Sasabe et al., 2000). P. infestans strains deficient in the elicitin INF1 induce disease lesions on Nicotiana benthamiana, suggesting that INF1 functions as an Avr factor that conditions resistance in this species (Kamoun et al., 1998). Using gene silencing, Peart et al. (2002) recently showed that the response of N. benthamiana to INF1 was dependent on the ubiquitin ligase-associated protein SGT1, which is also required for non-host resistance to bacterial plant pathogens. The N. benthamiana pathosystem holds great promise for dissecting elicitor response and resistance to P. infestans, since this plant is amenable to highthroughput functional assays using virus-induced gene silencing (VIGS) (Baulcombe, 1999). Nevertheless, the *N. benthamiana– P. infestans* interaction does not qualify as a strict non-host pathosystem since some wild-type isolates of *P. infestans* were recently found to infect this plant (F. Govers, personal communication; C. Smart & W.E. Fry, personal communication) (Kamoun, 2001).

We elected to employ Arabidopsis thaliana as a model for understanding non-host resistance to oomycete pathogens. Several biotrophic oomycetes, such as Peronospora parasitica and Albugo candida, are known to infect Arabidopsis (Holub et al., 1995; Parker et al., 1996; Rehmany et al., 2000; Reignault et al., 1996). Cabbage isolates of *Phytophthora brassicae* (previously known as *Phytophthora porri*) (Roetschi et al., 2001) and several isolates of Phytophthora cinnamomi (Robinson & Cahill, 2003) can also infect Arabidopsis, and these pathosystems are expected to facilitate the study of host infection by *Phytophthora*. However, most Phytophthora species, such as P. infestans, and the root pathogen Phytophthora sojae cannot infect Arabidopsis suggesting that this plant forms an untapped source of resistance to Phytophthora (Kamoun et al., 1999; Kamoun, 2001; Takemoto et al., 2003). Considering the impressive set of functional genomic resources that are available, Arabidopsis offers good prospects for dissecting the complex interactions that take place between a non-host plant and an oomycete pathogen and forms both an alternative and a complementary system to ongoing work on the resistance of Nicotiana to P. infestans. In this study, we describe the characterization of a non-host pathosystem involving Arabidopsis and an economically important Phytophthora species. Using cytological and molecular analyses, as well as microarray gene expression profiling, we obtained an overview of the active defence responses associated with the nonhost resistance of A. thaliana to P. infestans.

RESULTS

Interaction between Arabidopsis and P. infestans

To characterize the interaction between *Arabidopsis* and *P. infestans*, we performed repeated inoculations of *Arabidopsis* with *P. infestans* zoospores. We tested numerous inoculation parameters, including *Arabidopsis* leaves at the seedling or rosette stage, multiple combinations of *Arabidopsis* ecotypes and *P. infestans* strains, detached vs. attached leaves, and drop vs. spray inoculations. In all treatments, late blight lesions and sporulation were never observed, whereas infection of the host plant tomato was observed under most of the conditions tested. Normally, no macroscopic symptoms could be detected on *Arabidopsis*, but occasionally, discrete necrotic specks typical of the HR could be observed at the inoculation site, particularly when highly concentrated zoospore solutions were used.



Fig. 1 Hypersensitive response (HR) in *Arabidopsis*, 46 h after inoculation with *Phytophthora infestans*. Upon penetration of *P. infestans*, an infection vesicle and a secondary hypha were formed, and the HR was induced in a single epidermal cell. Characteristics of the HR cell are: (A) granular structure of the cytoplasm noted with DIC optics, (B) fluorescence of cytoplasm and cell wall with UV illumination and thickening of the cell wall; h, hyphae; HR, HR cell; iv, infection vesicle; n, nucleus, bar = 15 μ m.

Cellular responses of Arabidopsis to P. infestans

To determine the cellular responses of *Arabidopsis* to *P. infestans*, we performed microscope examinations of leaves inoculated at the rosette stage with droplets of zoospores. These analyses revealed the penetration of *Arabidopsis* epidermal cells in multiple independent infection sites. Cyst germination, penetration of epidermal cells and the formation of infection vesicles occurred as early as 46 h after inoculation, and in some cases was followed by the formation of a short secondary hyphae (Fig. 1). Penetrated epidermal cells displayed features typical of the HR, including granulated cell cytoplasm, thickened cell walls, condensed nuclei near the penetration site and autofluorescence under UV light (Fig. 1). These responses were typically limited to the penetrated epidermal cell.



Fig. 2 Kinetic PCR quantification of *Phytophthora infestans* biomass upon germination and penetration of *Arabidopsis* (non-host, open squares), tomato (host, solid squares) and in the absence of a plant substrate (no-host, open triangles). Zoospore suspensions of *P. infestans* were used to inoculate the respective plants. Four samples, each containing four leaf discs, were harvested at 0, 16, 24, 48 and 72 h after inoculation and used for DNA extraction. Four uninoculated leaf discs were added to the no-host treatment upon harvesting. Two ng of total DNA was used for every sample as a template for the PCR amplifications. DNA quantities were estimated after a natural log transformation of the obtained values. Arbitrary units of *P. infestans* DNA were used.

Phytophthora infestans biomass dynamics during infection of *Arabidopsis*

We used kinetic PCR technology to examine changes in P. infestans biomass during the interaction with Arabidopsis and the host plant tomato (Fig. 2). Primers specific to highly repetitive sequences (> ×10 000) in the P. infestans genome were previously used to quantify relative levels of P. infestans DNA in infected plant tissue, and were found to reflect an accurate and sensitive estimate of the P. infestans biomass (Judelson and Tooley, 2000). We performed kinetic PCR on DNA extracted from discs excised from Arabidopsis leaves infected with droplets of P. infestans zoospores at successive time points (0, 16, 24, 48 and 72 h after inoculation). Control treatments included inoculated leaves from tomato (host), and inoculum incubated in water in the absence of plant tissue (no-host). In both Arabidopsis and tomato, significant increases in the P. infestans biomass were observed in the initial 16 h. The Phytophthora infestans biomass continued to increase over the 3-day period on tomato, whereas it steadily declined on Arabidopsis to reach the lowest level at 72 h after inoculation (Fig. 2). In contrast, no notable changes in biomass were observed over the 3-day period for inoculum

incubated in the absence of plant tissue (Fig. 2). Statistical analyses using ANOVA were performed for each time point and suggested that the changes in biomass observed on *Arabidopsis* are statistically significant (P = 0.0011). Based on a protected mean separation, biomass increased significantly during the initial stages of *Arabidopsis* infection followed by a significant decrease in the later stages of the interaction (P < 0.05). Independent repetitions of the time course and the kinetic PCR experiments demonstrated that these biomass changes are reproducible (data not shown), however, the extent of the decrease in biomass observed at the later stages varied between experiments.

Local induction of *PR1* and *BGL2* genes during *P. infestans* infection

We used two transgenic Arabidopsis lines carrying fusions between the *PR1* and *BGL2* promoters to the β -glucuronidase (GUS) reporter gene (Cao et al., 1997; Manners et al., 1998; Stone et al., 2000) to examine the expression of these defence genes during P. infestans infection. The transgenic lines were inoculated at the rosette stage with either P. infestans zoospores or water droplets (mock treatment) and the leaves were excised and stained with X-Gluc at successive time points after inoculation. In both lines, elevated levels of PR-gene expression was detected 3 days after inoculation, as GUS staining around the inoculation sites (Fig. 3). No GUS expression was detected around mock-inoculated sites. Similar results were obtained from a series of independent experiments. Occasionally, light GUS staining was observed at some sites as early as 2 days after inoculation, but in most cases GUS staining was only observed 3 days or later after inoculation.

Induction of PDF1.2 during P. infestans infection

We assayed the expression of known defence genes during infection of Arabidopsis by P. infestans using Northern blot time-course analyses. Rosette leaves of Arabidopsis plants were sprayed with either water or *P. infestans* zoospore suspensions, and used for RNA extraction at 0, 16, 24, 48 and 72 h after inoculation. Northern blot hybridizations were performed with the defence-response genes PDF1.2, BGL2 and the constitutive gene UBQ5 (Fig. 4). A significant induction of PDF1.2 was observed at 48 h after inoculation. No induction of BGL2 was observed under Northern blot conditions over the 3-day period. No changes in transcript levels were observed in mock-inoculated plants. Independent replications of the time course suggested that the induction of PDF1.2 is significant and reproducible (data not shown), however, the timing of PDF1.2 induction varied between 16 and 48 h depending on the experiment.



Fig. 3 Induction of *Arabidopsis* (A) *PR1*::GUS and (B) *BGL2*::GUS expression by *Phytophthora infestans*. Transgenic lines were drop-inoculated with *P. infestans* zoospore suspensions. Leaves were harvested and stained with X-Gluc 3 days after inoculation with a droplet of *P. infestans* zoospores. Local expression of GUS was detected in both transgenic lines, 3 days after inoculation. Mock-inoculated sites did not show any detectable staining (left side of leaf in panel A).



Fig. 4 Time course Northern blot analysis of genes expressed in *Arabidopsis* rosette leaves 0,16, 24, 48 and 72 h after inoculation with *Phytophthora infestans* zoospores or mock inoculation with water. The probes corresponded to *PDF1.2*, a marker gene for the jasmonic acid pathway, and *BGL2*, a marker gene for the salicylic acid-mediated defence pathway. As a loading control, a probe for the constitutive ubiquitin 5 (*UBQ5*) gene was used.

DNA microarray gene-expression profiling of Arabidopsis non-host response to P. infestans

To further investigate Arabidopsis non-host response to P. infestans, we used DNA microarray analyses to determine expression profiles for c. 11 000 Arabidopsis cDNAs using the Arabidopsis Functional Genomics Consortium (AFGC) Microarray Facility (Wisman and Ohlrogge, 2000). In two hybridization experiments, RNA populations derived from mock treated and P. infestans-infected leaves were compared against each other using a dye-swap approach. Normalized data were subjected to regression analysis and subsequent outlier detection. Lists of outliers extracted from both hybridization data sets were compared to each other and a list containing an overlapping set of cDNAs was generated. A total of 89 cDNAs were identified that fall outside a 99% confidence interval (CI) in both hybridizations (supplementary data at http://www.oardc.ohio-state.edu /phytophthora /supp.htm). The cDNA identifiers were used to retrieve their respective predicted loci using the TAIR annotation database (http://www.arabidopsis.org) resulting in 54 annotated and non-redundant Arabidopsis genes (Table 1). A total of 11 genes were represented by multiple cDNAs (range 2-12) that were identified independently as being differentially expressed in both experiments. This suggests that the experiments and analyses we employed are reliable and robust.

Of the 54 *Arabidopsis* genes identified as differentially expressed during the non-host interaction with *P. infestans*, 52 were up-regulated and 2 were down-regulated. The 54 genes were classified into 7 functional categories that included physiological states related to metabolism, cell wall modification, development, as well as defence and stress responses. Genes that had known functions but that could not be placed in a particular functional category, were compiled in a separate class (Table 1).

Validation of DNA microarray analysis using Northern Blot analysis

We validated the analysis of the microarray data by a regression approach using Northern Blot hybridizations (Fig. 5). A Northern blot containing the RNA samples that were used in the microarray experiment was hybridized with probes from two genes, At1g21400 and At5g25350, that were selected as differentially expressed (Table 1), and two genes, At2g40000 and At3g43740, that were not. In addition, probes for *PDF1.2* and the constitutive gene *UBQ5* were included as controls (Fig. 5). The signals obtained with the various probes were quantified using a phosphor imager, normalized to the *UBQ5* signal, and used to calculate induction ratios. The induction levels obtained by Northern blot and microarray hybridization correlated well (Fig. 5). At1g21400 and At5g25350 were at least twofold induced in the Northern blot experiment vs. 3–5-fold in the microarray



Fig. 5 Validation of microarray analyses using Northern blot hybridization. RNA samples obtained from *Arabidopsis* rosettes 16 h after inoculation with *Phytophthora infestans* (Inf) or mock inoculation with water (H₂O) were blotted and hybridized with probes for At1g21400 and At5g25350, that were selected as differentially expressed based on microarray data analysis, and At2g40000 and At3g43740, that were not selected. *PDF1.2* and *UBQ5* were used as a positive control and a loading control, respectively. The numbers on the right correspond to the induction levels based on the Northern blot (RNA blot), and the two microarray experiments (slides I and II).

hybridizations. In contrast, At2g40000 and At3g43740 showed no significant differential expression by Northern blot hybridization. We also validated the microarray data by Northern Blot analysis using RNA isolated from an independently performed experiment (biological replicate). In this experiment, At1g21400 and At5g25350 were induced 2.8- and 2.4-fold relative to the *UBQ5* gene (data not shown). Overall, these results suggest that the microarray experiment and data processing by regression analyses are reliable.

Comparison of *Arabidopsis* **non-host response to** *P. infestans* **to other defence-related treatments**

Transcriptional changes observed during *P. infestans* infection were compared with those reported in two recent microarray analyses of defence-related treatments (Maleck *et al.*, 2000; Schenk *et al.*, 2000). We used TBLASTN searches to compare the data set of differentially expressed genes from our study to those

Table 1 *Arabidopsis* gene loci and number of representative cDNAs found differentially expressed during *Phytophthora* infection by microarray analysis. 11 000 cDNAs were used as targets for labelled cDNA from the two treatments. Genes were selected or considered significantly differentially expressed when the data points fell out of a 99% confidence interval during regression analysis in two separate comparisons (slides I and II). Genes that were found in previous studies (Maleck *et al.*, 2000; Schenk *et al.*, 2000) are indicated.

Putative function	Locus ID	Number of cDNAs	Slide I	Slide II	Identified by	
					Maleck <i>et al</i> .	Schenk <i>et al</i> .
Stress/defence						
leucine-rich repeats containing protein	At5q25350	1	3.2	4.5		
catalase 3	At1a20620	11	2.2	3.0	ves	ves
peroxidase, putative ATP2a	At2q37130	2	2.8	4.3	ves	ves
glutathione transferase	At2a30860	1	2.4	2.6	ves	ves
glutathione transferase, putative	At4a02520	1	2.2	2.7	ves	ves
family II lipase EXL3	At1a75900	1	2.3	2.6	,	,
lipoxygenase AtLOX2	At3a45140	1	-1.8	-2.3	ves	
thaumatin-like protein	At1a75030	1	-1.3	-1.3	Ves	Ves
Cell wall modification	, (cig) 5050	·	110		<i>j</i> co	yes
xylosidase (glycosyl hydrolase family 3)	At5a49360	3	24	3 1		
xylosidase (glycosyl hydrolase family 3)	At5g64570	1	2.1	4 1		
B-galactosidase (glycosyl hydrolase family 35)	At5a56870	1	2.5	4.1		
galactosidase (glycosyl hydrolase family 35)	At3a13750	2	2.5	2.7		
Development	///////////////////////////////////////	2	2.1	2.7		
ethylene response sensor ERS	At2a40940	1	2.5	3.6		ves
senescence-associated protein SEN1	At4a35770	5	2.2	3.0	ves	ves
dormancy-associated protein putative	At1a28330	2	2.4	2.7	Ves)
auxin-regulated protein	At2a33830	3	2.4	2.8	ves	
cytochrome P450 CYP83B1 indole	At4a31500	2	2.9	4.2	Ves	Ves
alucosinolate synthase	, (c1g0 1000	-	215		<i>J</i> = 5	yes
late embryogenesis abundant protein	At4a02380	1	5.0	6.2		
nodulin-like protein	At5a14120	1	2.0	3.1		
Metabolism	,	·	210	511		
branched-chain amino transferase	Δt1a10070	1	3.4	35		
branched-chain alnha keto-acid debydrogenase	At1g21400	1	3.7	3.6		
10-formyltetrahydrofolate synthetase	At1a50/80	1	23	2.8		
acetolactate synthese nutative	At7g31810	1	2.5	2.0		
nbytochalatin synthese AtPCS1	At5g//070	1	2.0	2.7		
formate debudrogenace EDH	At5a14780	1	2.2	2.0		100
aldabuda dabudroganasa bomalag	At1g54100	1	2.1	2.0		yes
aldenyde denydrogenase homolog	At1g54100	1	2.2	2.9		yes
putative thosephosphate isofferase	Atzyz 1170	1	2.1	2.2		
	ALSYSS150	1	2.5	2.1		
	ALIGITZOU	Z	2.5	2.2		
AP2 domain protein RAP2 3	At3a16770	3	2 21	3 3/	VOC	105
probable transcription regulator protein	At3q/8530	1	2.21	3.14	yes	yes
Othor	Al3940330	I	2.00	5.15		
alucing rich PNA hinding protein AtCPD7	A+2a21660	10	2 20	2 70		100
putative patatin	At2g21000	1	2.33	5 37		yes
putative patatili	At2920300	1	2.00	2.27		
villin 2 fragment	ALT920270	1	2.00	2.00		
putative myosin beauty chain	ALSY57410	1	2.7	2.0		
putative myosin neavy chain	AL2952240	1	5.5	4.7 2.E		
COS ribosomol protoin L7A	ALTG07940	1	2.5	2.5		
	Αισύοτα/Ο	I	1.9	3.4	yes	
	At1a21500	1	2.00	2 60		
annown ON	AL1931300	1	2.09	2.00		Voc
expressed protein	ALJY37033	1	1.9Z 2.10	2.1Z	VOC	yes
expressed protein	ALZY20220	1	2.10	J.41 1 J1	yes	Voc
expressed protein	AL5901290	I	2.00	4.24	yes	yes

Putative function	Locus ID	Number of cDNAs	Slide I	Slide II	Identified by	
					Maleck <i>et al</i> .	Schenk <i>et al</i> .
expressed protein	At3g15450	1	1.93	2.58	yes	
expressed protein	At1g21680	1	2.05	3.34		
unknown protein	At2g30600	1	2.04	2.44		
unknown protein	At1g73960	1	2.63	3.87		
unknown protein	At1g78110	1	2.08	2.44		
conserved hypothetical protein	At3g24860	1	2.05	3.56		
hypothetical protein	At4g16110	1	1.03	1.38		yes
putative protein; hypothetical protein	At3g49590	1	2.07	2.62		
putative protein	At4g17900	1	2.47	3.42		
putative protein	At4g24690	1	2.05	3.56		
putative protein	At5g53160	1	2.15	4.32		

(A)



Fig. 6 Cluster analysis illustrating the relatedness of transcriptional changes between *Phytophthora infestans* and other defence-related treatments. The *P. infestans* data sets corresponding to two microarray experiments (*P. infestans* I and II) were combined with overlapping data from the transcriptional profiling experiments reported by (A) Schenk *et al.* (2000) and (B) Maleck *et al.* (2000). The gene numbers are indicated at the top, and the defence treatments were described in Schenk *et al.* (2000) and Maleck *et al.* (2000). The red colour corresponds to up-regulated genes, whereas green represents down-regulated genes.

published previously (see Experimental procedures). Of the 54 *Arabidopsis* genes represented in our data set, 15 matched cDNAs identified in the study performed by Maleck *et al.* (2000) and 15 cDNAs identified by Schenk *et al.* (2000) (Table 1). The expression ratios of the respective genes were extracted from the data sets provided with the two studies and used for building two data matrices. Cluster analysis of the two matrices generated an overview of relatedness between the various treatments and *P. infestans* infection (Fig. 6). Patterns of defence responses induced

by MeJA treatment were found to be more similar to responses induced by *P. infestans* infection, whereas ethylene, SA and *Alternaria* inducing conditions resulted in less similar defenceinduction profiles (Fig. 6A). Similar comparisons to the SAR-related treatments described by Maleck *et al.* (2000) were made. Cluster analysis revealed a notable resemblance of our expression data to gene expression profiles in *cim11* mutant genotypes, as well as gene expression changes 48 h after treatment with the salicylate analogue benzothiadiazole (BTH). In addition, *P. infestans* defence responses shared some similarity to those induced by *nim1* over-expression, compatible and incompatible *Peronospora parasitica* interactions, *Pseudomonas syringae* (*AvrRpt2*) infection, and early response after BTH treatment (4 h) (Fig. 6B). Experimental treatments involving plant genotypes containing the *nahG* gene showed least similarity, together with *cim6* and *cim7* and various double-mutant genotypes.

DISCUSSION

We performed the cytological and molecular characterization of a non-host pathosystem involving the model crucifer plant A. thaliana and the destructive and economically important oomycete pathogen P. infestans. Our cytological observations confirm previous work by Vleeshouwers et al. (2000), which showed that the interaction of *P. infestans* with non-host plants, including those that are phylogenetically distant from the solanaceous hosts, is typically associated with the penetration of plant tissue and the HR. However, a more detailed cytological investigation needs to be performed to determine whether the proportion of successful penetration events and the level of HR induction following penetration differ between host and non-host interactions. Nevertheless, our results suggest that recognition of P. infestans by Arabidopsis takes place and may form one important barrier in non-host resistance. Therefore, a model that evokes an arsenal of Arabidopsis R genes that recognize multiple or essential P. infestans Avr genes is sufficient to explain non-host resistance in this pathosystem, but it cannot be ruled out that additional layers of nonspecific defence responses occur. With extensive genetic and genomic resources available, the described Arabidopsis-P. infestans pathosystem offers excellent prospects for dissecting the complex layers that may form non-host resistance.

In addition to cytological analyses, we used kinetic PCR to monitor relative levels of P. infestans DNA and consequent biomass during infection (Fig. 2). Previously, DNA and RNA blot hybridizations have been used to estimate the biomass of pathogenic oomycetes in plant tissue (Kamoun et al., 1998; Rairdan et al., 2001). However, these techniques are not sensitive enough to monitor the small changes in P. infestans biomass that are expected to occur on non-host plants. In contrast, kinetic PCR is highly sensitive, quantitative, objective and should prove ideal for non-host pathosystems. To enhance the sensitivity of the kinetic PCR quantification, we used primers corresponding to highly repetitive (> \times 10 000) sequences from the *P. infestans* genome that allow amplification of as little as 10 fg of P. infestans DNA (Judelson and Tooley, 2000). The sensitivity of these primers is obvious since we routinely obtained quantifiable signals from inoculation sites bearing as little as 1000 zoospores.

Phytophthora infestans exhibited dynamic changes in biomass over a 3-day infection of *Arabidopsis* (Fig. 2). Over the first 16 h, a significant increase in biomass was observed that was similar to the increase observed on the host tomato. This early increase may correspond to the germination of cysts, penetration of plant epidermis, and formation of infection vesicles and short secondary hyphae, as determined by cytology on both Arabidopsis and host plants. This suggests that some level of growth and nuclear division, perhaps in the infection vesicles or secondary hyphae, occurs in P. infestans during early infection of Arabidopsis. Subsequently, a gradual but significant decrease in P. infestans biomass was observed from 24 to 72 h after inoculation of Arabidopsis, and contrasted sharply with the steady increase observed on tomato. This decrease may reflect death and degradation of P. infestans hyphae caused by the HR and correlates with the termination of pathogen ingress determined by cytology. Interestingly, the dynamic changes in *P. infestans* biomass observed on Arabidopsis contrasted with the constant level of biomass observed for P. infestans cysts germinating in water in the absence of plant tissue. These results support the interpretation that a successful penetration of the plant epidermis rather than surface growth is required for the biomass increase we observed in the early stages of the interaction. Taken together, these results indicate that P. infestans is able to successfully initiate an infection on Arabidopsis and complement the cytological analyses.

To gain a first insight into the molecular aspects of *Arabidopsis* non-host response to *P. infestans*, we examined changes in the expression of defence genes using Northern blot hybridizations with probes for *PDF1.2* and *BGL2* (Fig. 4), and the transgenic lines *PR1*::*GUS* and *BGL2*::*GUS* (Fig. 3). Although no induction of *PR1* or *BGL2* was detected by Northern blot analyses, a localized expression of both genes was detected around inoculation sites in the transgenic reporter lines beginning 3 days after inoculation. Since we found *PR1* and *BGL2* to be locally induced at very late stages of the interaction, the discrepancy between the two methods may point to a difference in sensitivity. Considering that most of the cells in the inoculated leaves are not infected, a dilution effect may have reduced the sensitivity of the Northern analysis.

PDF1.2 is a marker for the jasmonate (JA)/ethylene (ET)mediated defence-response pathways, and its up-regulation has been associated with numerous pathogen or defence-related treatments (Glazebrook, 2001). On the other hand, PR1 and BGL2 are marker genes for the salicylate (SA)-mediated defence pathway that is typically induced following infection by necrotizing pathogens or the HR, and during systemic acquired resistance (SAR) (Glazebrook, 2001; Ryals et al., 1996). Taken together, our data suggest the sequential induction of the JA/ET pathway followed by the SA pathway during non-host response of Arabidopsis to P. infestans. Studies in other Arabidopsis pathosystems suggest significant cross-talk and the co-regulation of both SA and JA/ET mediated defence pathways (Clarke et al., 2000; Ellis et al., 2002; Glazebrook, 2001; Schenk et al., 2000). In addition, these pathways have been shown to work antagonistically as well as in concert, to confer enhanced resistance to fungal, bacterial and oomycete pathogens (Cohn *et al.*, 2001; Ellis *et al.*, 2002; van Wees *et al.*, 2000). The direct role of JA, ET and SA signalling in non-host resistance to *P. infestans* remains to be determined. Quantitative assays using *Arabidopsis* mutant genotypes covering the various branches of known defence pathways are currently underway to address this question.

To gain a better understanding of Arabidopsis non-host responses to P. infestans, we performed DNA microarray experiments comparing the responses of Arabidopsis plants inoculated with P. infestans to their mock inoculated counterparts. Despite the harsh selection conditions imposed on the data set, a remarkably high level of redundancy was found amongst the positive cDNAs (see supplementary data at http://www.oardc.ohiostate.edu/phytophthora/supp.htm). Subsequent Northern blot analysis validated our regression analysis strategy. There was a clear correlation in induction levels between the microarray experiment and Northern blot hybridizations using two selected genes and two non-selected genes (Fig. 5). In addition, about one-third of the cDNAs that were identified in our experiments overlapped with cDNAs identified in the defence response gene expression profiling studies of Maleck et al. (2000) and Schenk et al. (2000) (Table 1).

We classified plant responses using cluster analyses of gene expression profiles across the P. infestans treatments and other defence related treatments (Fig. 6). P. infestans-induced defence responses were most similar to gene-expression changes after MeJA treatment. From the 15 genes that were used for these comparisons, 14 genes were also induced by MeJA, according to Schenk et al. (2000), suggesting that non-host defence responses to P. infestans are associated with activation of the JA response pathway (Fig. 6A). In contrast, the expression profiles of all other treatments, such as ethylene, Alternaria and SA, had less similarity to our data set. Similarly, patterns of overlap were found in the comparison of SAR-related treatments (Maleck et al., 2000) to our data set (Fig. 6B). Cluster analysis of expression profiles indicated a similarity between treatments and the occurrence of two general groups of Arabidopsis genes within the set of P. infestans up-regulated genes. Two genes were up-regulated 48 h after BTH treatment and had higher expression levels in the *cim11* (constitutive immunity) mutant background. A second and larger group of seven genes was commonly up-regulated in our treatment and during Peronospora parasitica compatible and incompatible interactions, BTH treatment (4 h), plants over-expressing NIM1, and Pseudomonas syringae infection. The transcript levels of the genes used in this comparison were either unchanged or lowered in all NahG plants and NahG-containing mutants (cim11NahG and cim6NahG), cim6 and cim7 mutant genotypes and other related treatments.

Some notable genes that are induced in the defence-related treatments examined by Schenk *et al.* (2000) and Maleck *et al.* (2000) were identified in this study (Table 1). Among these,

Rap2.3 (At3q16770), which encodes an AP2 domain transcription factor, was up-regulated during non-host resistance and many other defence responses, suggesting that it may mediate common regulatory steps in defence pathway activation or modulation. Another gene, Cyp83B1 (At4g31500), is a member of a large family of cytochrome P450 genes, and is involved in the production of indole-glucosinolates as well as the plant hormone IAA (auxin) (Bak and Feyereisen, 2001). The expression of Cyp83B1 and related members of this gene family was found to be elevated upon SA and MeJA treatments and were associated with an increase in indole-glucosinolates (Mikkelsen et al., 2003). Smolen and Bender (2002) identified a non-functional mutant of Cyp83B1 that showed a lesion-mimic phenotype. Taken together, these data indicate a possible involvement of Cyp83B1, and perhaps indoleglucosinolates, in defence responses and possibly regulation of the HR.

Many of the Arabidopsis genes identified as up-regulated during non-host response to P. infestans could be related to cellular aspects of signalling and defence. For instance, four glycosyl hydrolase genes (At3g13750, At5g49360, At5g56870 and At5g64570), that are possibly involved in modifications of cell wall components were up-regulated two- to fivefold (Table 1). During cell stress and pathogen attack, cell wall modifications are commonly observed (Heath, 1998; Nicholson and Hammerschmidt, 1992; Vleeshouwers et al., 2000) which is also illustrated by the cell wall depositions described in Fig. 1. Therefore, cell wall alterations are likely to form a major barrier in non-host resistance. Other notable genes that were up-regulated include genes related to oxidative stress such as catalase (At1g20620), glutathione transferases (At2g30860 and At4g02520), and peroxidase (At2g37130). The occurrence of these genes during non-host HR is not surprising since the role of oxidative stress and production of active oxygen species (AOS) during the HR is well documented (Delledonne et al., 2001; Levine et al., 1994; Sasabe et al., 2000). Since the HR is observed in the Arabidopsis-P. infestans interaction, induction of these genes in concert with early defence is plausible. Overall, these data support the concept that defence responses induced by P. infestans involve the HR as well as JA-mediated signalling and defence.

In addition to genes in common between non-host and various host defence treatments, some genes uniquely up-regulated during the *P. infestans* interaction were identified (Table 1). The function of these genes in non-host resistance remains unclear, but they represent attractive candidates for functioning in processes unique to non-host resistance to *P. infestans*, and perhaps, other non-host pathogens.

In this study we characterized the interaction between *P. infestans* and the non-host plant *A. thaliana*. An integrated multifaceted approach has enhanced our understanding of this interaction and is helping us to devise future research strategies. Based on the diversity of molecular genetic tools and genomic

resources available for *Arabidopsis* and *Phytophthora*, we expect this non-host pathosystem to become of key importance in studies on molecular plant–microbe interactions. Further research on this pathosystem will provide significant insight into key molecular processes regulating non-host resistance to an economically important pathogen. The knowledge gained will result in immediate biotechnological applications and will offer novel perspectives for engineering durable resistance in crop plants.

EXPERIMENTAL PROCEDURES

Plant growth conditions

Arabidopsis (Col-3) seeds were routinely surface sterilized in 70% EtOH for 30 s, followed by incubation in 50% bleach solution for 10 min. Seeds were then washed multiple times in dH₂O before plating on to MS-Phytagar sucrose plates (1 × MS salts, 2% w/v sucrose, 0.8% w/v Phytagar). Plated seeds were incubated at 4 °C for 3–4 days prior to germination. Seven-to-10 day old seedlings were transferred to potting media and grown under controlled conditions (22 °C, 8 h photoperiod). Mature non-bolting plants at the rosette stage (4–5 weeks) were used for infection experiments.

Phytophthora infestans culturing and infection assays

Cultures of *P. infestans* isolate 90128 (A2 mating type, race 1.3.4.7.8.9.10.11, isolated from potato in the Netherlands in 1990), were routinely grown on rye agar medium supplemented with 2% sucrose (Caten and Jinks, 1968). Zoospores were produced by flooding 11–14-day-old cultures with dH₂O, followed by incubation at 4 °C for 1–3 h. Rosette leaves of 4–5 week-old *Arabidopsis* (Col-3) plants were inoculated with 10 μ L droplets of zoospore suspensions for microscopy. For the DNA microarray and Northern time course experiments, complete rosette stage plants were sprayed with zoospore suspensions. Concentrations ranging from 200 000 to 500 000 zoospores/mL were used for all experiments. Deionized water was used as a negative control in all relevant experiments.

Microscope observations

Leaf discs containing the inoculum were excised at various times after inoculation and examined by microscopy for plant response and growth of *P. infestans*. Lactophenol-trypan blue staining and destaining with chloral hydrate were performed as previously described (Colon *et al.*, 1992; Wilson and Coffey, 1980). The discs were examined using a Zeiss Axiophot microscope equipped with a high-pressure mercury vapour lamp. Autofluorescence was observed with a G365 excitation filter, FT395 interference beam splitter and LP420 barrier filters.

GUS staining procedure

Complete *Arabidopsis* leaves were immersed in a GUS staining solution (2 mM 5-bromo-4-chloro-3-indolyl-beta-D-glucuronic acid (X-Gluc) (Rose Scientific, Edmonton Alberta, Canada) in 0.2% Triton X-100, 50 mM NaHPO4 Buffer (pH 7.2), 2 mM potassium ferrocyanide, 2 mM potassium ferricyanide). A vacuum was applied for 10 min and was then gently released over several minutes. Leaves were incubated at 37 °C for 24 h and examined macroscopically for staining patterns.

Kinetic PCR quantification of P. infestans

For kinetic (real-time) PCR quantification of *P. infestans* biomass in plant tissue, four samples were taken for every experimental treatment or time point, with each sample consisting of a pool of four leaf discs corresponding to four independent inoculation sites. Total DNA was extracted from each pool of four leaf discs using a Ojagen Plant DNA extraction kit following the manufacturer's instructions. DNA was quantified using the picogreen ds DNA guantification kit (Molecular Probes, Eugene, OR) and checked by electrophoresis. Dilutions were made using volumes of 10 µL or larger to minimize pipeting errors. Kinetic PCR was performed on a Roche Lightcycler (Roche, Indianapolis, IN) using Lightcycler-FastStart DNA Master SYBR Green I reagents, primers J-08-3 and J-08-4 previously described by Judelson and Tooley (2000), and 2–10 ng of total DNA as a template. PCR conditions were 45 cycles of 1 s at 95 °C, 5 s at 50 °C, and 10 s at 72 °C, with a temperature transition rate of 20 °C per s. Phytophthora infestans relative DNA amounts were estimated using a standard curve generated using 100, 10, 1 and 0.1 pg of purified P. infestans total DNA. Natural log (In) transformation was applied on the data to obtain distributions that approached normality. Subsequently, ANOVA was performed on the normalized data using both treatment and replication in the model as fixed effects using the SAS statistical software package version 8 (SAS Institute, Cary, NC). Least significant difference mean separation was used to detect significant differences between the time points assayed.

RNA manipulations and Northern blot analysis

RNA was extracted using the Trizol reagent (Gibco-BRL, Bethesda, MD) using the manufacturer's directions. RNA samples were checked for purity and integrity using spectrophotometry and electrophoresis. When necessary, samples were cleaned further by adding 1 volume of phenol:chloroform:isoamyl-alcohol (125 : 24 : 1, pH 6.7) to the sample, high-speed centrifugation (15800 g, 15 min) and subsequent collection of the aqueous phase. This step was then followed by another extraction using equal volumes of chloroform. RNA was ethanol precipitated from the aqueous phase and dissolved in dH₂O. Membranes for Northern blot

analysis were prepared using a modified method from McMaster and Carmichael (1977) as described in Sambrook *et al.* (1989). 15–20 μ g of RNA per sample was used and RNA was blotted on Hybond N⁺ membranes (Amersham, Piscataway, NJ) following standard procedures and instructions from the manufacturer.

Hybridization probes were first prepared using Polymerase Chain Reaction (PCR) amplifications. Primers for the amplification of PDF1.2, BGL2 and UBQ5 were as described elsewhere (Glazebrook et al., 1996; Penninckx et al., 1996; Xiao et al., 2000). Primers for amplification of the selected genes for microarray validation were: At1g21400-F (5'-GAGAAGTCGATATGGACATGA-TAC-3') and At1g21400-R (5'-AACGGATGGTGGAGTGAGGAAG-3'); At5q25350-F (5'-CTTCACTCCTACTGATACTACTCA-3') and At5G25350-R (5'-CT TCGAAT TATGTCTGGAATC T TCA-3'); At2q40000-F (5'-CGAAGTTCTCAATTGAGACCAG-3') and At2g40000-R (5'-GGAACAATCCCAACAAACGGA-3'); At3g43740-F (GAGGAAGAT-GGTATCATCAG-3') and At3q43740-R (5'-TCCATTCACGGTGGTT-GATG-3'). Amplified fragments were purified from TAE-agarose gels and sequenced using an ABI Prism 377 automated sequencer (PE Applied Biosystems). Similarity searches were used to confirm amplification of the correct fragments.

All probes were labelled with α -³²P-dCTP using a random primer labelling kit (Gibco-BRL, Bethesda, MD). All Northern blot hybridizations were carried out at 65 °C in Modified Church Buffer (0.36 M Na₂HPO₄, 0.14 M NaH₂PO₄, 1 mM EDTA and 7% SDS). Membranes were washed for 15 min in 1× SSC/0.5%SDS and 0.5× SSC/0.5%SDS at 65 °C, followed by 1 wash in 0.5× SSC/ 0.5% SDS for 1 min at room temperature. Membranes were exposed to a phosphor imager screen for 24–48 h and scanned using a Molecular Dynamics Storm 840 Phosphor Imager.

DNA microarray hybridizations

Total RNA samples were subjected to standard probe preparations and microarray hybridization procedures as described in the protocol section on the Arabidopsis Functional Genomics Consortium (AFGC) website (http://afgc.stanford.edu/afgc.html/site2.htm). Briefly, two purified mRNA samples were used for synthesis of probes labelled with either CY3 or CY5 fluorescent dye. Four separately labelled cDNA samples were generated using a dye swap to distinguish replicates of the same treatment. These were hybridized to two slides in a dye-swap experimental setup (technical replication). Hybridized slides were scanned using a Scan Array 3000 (GSI Lumonics, Billerica, MA). Two data points were obtained for each spot on one slide.

Microarray data analysis

The generated data were normalized by the default method specified by the AFGC (http://afgc.stanford.edu/afgc_html/site2.htm). All data analysis and processing steps were done using the SAS statistical software package version 8 (SAS Institute, Cary, NC, USA). For analysis purposes, normalized but 'raw' expression values were downloaded for both hybridized slides (technical replicates) from the AFGC web site. The natural log of every expression value was calculated to obtain a data set with distributions that approached normality. Transformed data points were then used for comparisons, applying a regression analysis/ outlyer detection technique. In short, for every slide, normalized and transformed expression values of both treatments were plotted against each other (x-axis, expression values of mock treatment, y-axis, expression values of P. infestans treatment). A 99% confidence interval (CI) was generated and used to select cDNAs that correspond to data points falling outside of the 99% CI, representing significant differential expression. Lists of identifiers representing the cDNA spots that were detected were compiled. Only cDNAs identified as outliers at both slides were considered significantly differentially expressed and were used for further analysis. The cDNA identifiers were used to retrieve annotated locus names from the Arabidopsis Information Resource (TAIR) database (http://www.Arabidopsis.org). A non-redundant data set was generated and was used to construct cluster data files.

Data compilation and cluster analysis

To identify overlap between the genes identified in this study and published data sets, we first compiled files containing nucleotide sequences for the differentially regulated genes reported in the DNA microarray studies of Schenk et al. (2000) and Maleck et al. (2000). We then used amino acid sequences corresponding to the 54 differentially expressed genes (Table 1), to perform a TBLASTN search against the generated sequence files. E-value scores below 10⁻¹² were considered significant. Data matrices were constructed by retrieving and combining expression data of the genes common in either relevant data set. Expression ratios for each gene and treatment were calculated using the inherent control of each treatment as the denominator. Two data matrices were used for cluster analysis using the CLUSTER analysis software package (Eisen et al., 1998). Self-Organizing Map (SOM) analysis was first used to generate classes of genes based on expression ratios across all treatments. Output files generated in this procedure were then used to order the input file for cluster analysis. Complete average linkage hierarchy clustering was used for both comparisons presented. Generated results were visualized and evaluated using TREEVIEW (Eisen et al., 1998).

Microarray data availability

The entire data set can be freely obtained and searched at the AFGC website (http://afgc.stanford.edu/afgc_html/site2.htm). Data for the differentially regulated genes is also provided as a supplementary file (http://www.oardc.ohio-state.edu/phytophthora/supp.htm).

ACKNOWLEDGEMENTS

This work was supported by the OARDC Research Enhancement Grant Program. Salaries and research support were provided, in part, by State and Federal Funds appropriated to the Ohio Agricultural Research and Development Center, the Ohio State University. We are grateful to Caitlin Cardina, Shujing Dong, Diane Kinney and Antonino Testa for their expert technical assistance; Bert Bishop for his suggestions regarding statistical analysis; as well as Tea Meulia and the staff of the OARDC Molecular and Cellular Imaging Center for help with the kinetic PCR experiments. We are also grateful to Alan Shapiro and Brian Staskawicz for the *PR1-GUS* line, the *Arabidopsis* Biological Resource centre for the *Arabidopsis* lines and the *Arabidopsis* Functional Genomics Consortium for providing their DNA microarray service, expertise and assistance.

SUPPLEMENTARY MATERIAL

A Table has been provided, which is available at <http:// www.blackwellpublishing.com/products/journals/suppmat/MPP/ MPP195/MPP195sm.htm>, and which lists identified cDNAs with their respective GENBANK accession and Locus ID numbers. Redundancy represents the number of cDNAs identified from each locus. Ratios S1 and S2 were calculated using the normalized expression values from each hybridized slide.

REFERENCES

- Bak, S. and Feyereisen, R. (2001) The involvement of two P450 enzymes, CYP83B1 and CYP83A1, in auxin homeostasis and glucosinolate biosynthesis. Plant Physiol. 127, 108–118.
- Baldauf, S.L., Roger, A.J., Wenk-Siefert, I. and Doolittle, W.F. (2000) A kingdom-level phylogeny of eukaryotes based on combined protein data. *Science*, 290, 972–977.
- Baulcombe, D.C. (1999) Fast forward genetics based on virus-induced gene silencing. *Curr. Opin. Plant Biol.* 2, 109–113.
- Brunner, F., Rosahl, S., Lee, J., Rudd, J.J., Geiler, C., Kauppinen, S., Rasmussen, G., Scheel, D. and Nurnberger, T. (2002) Pep-13, a plant defense-inducing pathogen associated pattern from *Phytophthora* transglutaminases. *EMBO J.* 16, 6681–6688.
- Cao, H., Glazebrook, J., Clarke, J.D., Volko, S. and Dong, X. (1997) The Arabidopsis NPR1 gene that controls systemic acquired resistance encodes a novel protein containing ankyrin repeats. Cell, 88, 57–63.
- Caten, C.E. and Jinks, J.L. (1968) Spontaneous variability of single isolates of *Phytophthora infestans*. I. Cultural variation. *Can. J. Bot.* 46, 329–347.
- Clarke, J.D., Volko, S.M., Ledford, H., Ausubel, F.M. and Dong, X. (2000) Roles of salicylic acid, jasmonic acid, and ethylene in *cpr*-induced resistance in *Arabidopsis. Plant Cell*, **12**, 2175–2190.
- Cohn, J., Sessa, G. and Martin, G.B. (2001) Innate immunity in plants. *Curr. Opin. Immunol.* **13**, 55–62.
- Colon, L.T., Eijlander, R., Budding, D.J., van Ijzendoorn, M.T., Pieters, M.M.J. and Hoogendoorn, J. (1992) Resistance to potato late blight (*Phytophthora infestans* (Mont.) de Bary) in *Solanum nigrum*,

Solanum villosum and their sexual hybrids with *Solanum tuberosum* and *Solanum demissum. Euphytica*, **66**, 55–64.

- Dangl, J. and Jones, J.D. (1998) Affairs of the plant: colonization, intolerance, exploitation and co-operation in plant–microbe interactions. *Curr. Opin. Plant Biol.* 1, 285–287.
- Delledonne, M., Zeier, J., Marocco, A. and Lamb, C. (2001) Signal interactions between nitric oxide and reactive oxygen intermediates in the plant hypersensitive disease resistance response. *Proc. Natl Acad. Sci.* USA, 98, 13454–13459.
- Eisen, M.B., Spellman, P.T., Brown, P.O. and Botstein, D. (1998) Cluster analysis and display of genome-wide expression patterns. *Proc. Natl Acad. Sci. USA*, 95, 14863–14868.
- Ellis, C., Karafyllidis, I. and Turner, J.G. (2002) Constitutive activation of jasmonate signaling in an *Arabidopsis* mutant correlates with enhanced resistance to *Erysiphe cichoracearum, Pseudomonas syringae*, and *Myzus persicae*. *Mol. Plant-Microbe Interact.* **15**, 1025–1030.
- Erwin, D.C. and Ribeiro, O.K. (1996) *Phytophthora Diseases Worldwide*. St Paul, MN: APS Press.
- Flor, H.H. (1971) Current status of the gene-for-gene concept. *Annu. Rev. Phytopathol.* **9**, 275–296.
- Fry, W.E. and Goodwin, S.B. (1997a) Re-emergence of potato and tomato late blight in the United States. *Plant Dis.* **81**, 1349–1357.
- Fry, W.E. and Goodwin, S.B. (1997b) Resurgence of the Irish potato famine fungus. *Bioscience*, 47, 363–371.
- Garelik, G. (2002) Agriculture. Taking the bite out of potato blight. *Science*, 298, 1702–1704.
- Glazebrook, J. (2001) Genes controlling expression of defense responses in *Arabidopsis*—2001 status. *Curr. Opin. Plant Biol.* **4**, 301–308.
- Glazebrook, J., Rogers, E.E. and Ausubel, F.M. (1996) Isolation of Arabidopsis mutants with enhanced disease susceptibility by direct screening. Genetics, 143, 973–982.
- Gomez-Gomez, L. and Boller, T. (2002) Flagellin perception: a paradigm for innate immunity. *Trends Plant Sci.* 7, 251–256.
- Gross, P., Julius, C., Schmelzer, E. and Hahlbrock, K. (1993) Translocation of cytoplasm and nucleus to fungal penetration sites is associated with depolymerization of microtubules and defence gene activation in infected, cultured parsley cells. *EMBO J.* **12**, 1735–1744.
- Heath, M.C. (1998) Apoptosis, programmed cell death and the hypersensitive response. *Eur. J. Plant Pathol.* **104**, 117–124.
- Heath, M.C. (2000) Nonhost resistance and nonspecific plant defenses. *Curr. Opin. Plant Biol.* **3**, 315–319.
- Holub, E.B., Brose, E., Tor, M., Clay, C., Crute, I.R. and Beynon, J.L. (1995) Phenotypic and genotypic variation in the interaction between *Arabidopsis thaliana* and *Albugo candida*. *Mol. Plant-Microbe Interact*. 8, 916–928.
- Judelson, H.S. and Tooley, P.W. (2000) Enhanced polymerase chain reaction methods for detecting and quantifying *Phytophthora infestans* in plants. *Phytopathology*, **90**, 1112–1119.
- Kamoun, S. (2000) Phytophthora. In: *Fungal Pathology* (Kronstad, J., ed.) pp. 237–265. Netherlands: Kluwer Academic Publishers.
- Kamoun, S. (2001) Nonhost resistance to *Phytophthora*: novel prospects for a classical problem. *Curr. Opin. Plant Biol.* 4, 295–300.
- Kamoun, S. (2003) Molecular genetics of pathogenic oomycetes. Euk. Cell, 2, 191–199.
- Kamoun, S., Huitema, E. and Vleeshouwers, V.G.A.A. (1999) Resistance to oomycetes: a general role for the hypersensitive response? *Trends Plant Sci.* 4, 196–200.
- Kamoun, S., van West, P., de Jong, A.J., de Groot, K., Vleeshouwers, V.G.A.A.

and Govers, F. (1997) A gene encoding a protein elicitor of *Phytophthora infestans* is down-regulated during infection of potato. *Mol. Plant-Microbe Interact.* **10**, 13–20.

- Kamoun, S., van West, P., Vleeshouwers, V.G., de Groot, K.E. and Govers, F. (1998) Resistance of *Nicotiana benthamiana* to *Phytophthora infestans* is mediated by the recognition of the elicitor protein INF1. *Plant Cell*, **10**, 1413–1426.
- Levine, A., Tenhaken, R., Dixon, R. and Lamb, C. (1994) H₂O₂ from the oxidative burst orchestrates the plant hypersensitive disease resistance response. *Cell*, **79**, 583–593.
- Maleck, K., Levine, A., Eulgem, T., Morgan, A., Schmid, J., Lawton, K.A., Dangl, J.L. and Dietrich, R.A. (2000) The transcriptome of *Arabidopsis thaliana* during systemic acquired resistance. *Nat. Genet.* 26, 403–410.
- Manners, J.M., Penninckx, I.A., Vermaere, K., Kazan, K., Brown, R.L., Morgan, A., Maclean, D.J., Curtis, M.D., Cammue, B.P. and Broekaert, W.F. (1998) The promoter of the plant defensin gene *PDF1.2* from *Arabidopsis* is systemically activated by fungal pathogens and responds to methyl jasmonate but not to salicylic acid. *Plant Mol. Biol.* 38, 1071–1080.
- Margulis, L. and Schwartz, K.V. (2000) *Five Kingdoms: an Illustrated Guide to the Phyla of Life on Earth*. New York: WH Freeman.
- McMaster, G.K. and Carmichael, G.G. (1977) Analysis of single- and double- stranded nucleic acids on polyacrylamide and agarose gels by using glyoxal and acridine orange. *Proc. Natl Acad. Sci. USA*, 74, 4835–4838.
- Mikkelsen, M.D., Petersen, B.L., Glawischnig, E., Jensen, A.B., Andreasson, E. and Halkier, B.A. (2003) Modulation of *CYP79* genes and glucosinolate profiles in *Arabidopsis* by defense signaling pathways. *Plant Physiol.* **131**, 298–308.
- Morrissey, J.P. and Osbourn, A. (1999) Fungal resistance to plant antibiotics as a mechanism of pathogenesis. *Microbiol. Mol. Biol. Rev.* 63, 708–724.
- Naton, B., Hahlbrock, K. and Schmelzer, E. (1996) Correlation of rapid cell death with metabolic changes in fungus-infected, cultured parsley cells. *Plant Physiol.* **112**, 433–444.
- Nicholson, R.L. and Hammerschmidt, R.E. (1992) Phenolic compounds and their role in disease resistance. *Annu. Rev. Phytopathol.* 30, 369– 389.
- Nurnberger, T. and Brunner, F. (2002) Innate immunity in plants and animals: emerging parallels between the recognition of general elicitors and pathogen-associated molecular patterns. *Curr. Opin. Plant Biol.* 5, 318– 324.
- Osbourn, A. (1996) Preformed antimicrobial compounds and plant defense against fungal attack. *Plant Cell*, **8**, 1821–1831.
- Parker, J.E., Holub, E.B., Frost, L.N., Falk, A., Gunn, N.D. and Daniels, M.J. (1996) Characterization of *eds1*, a mutation in *Arabidopsis* suppressing resistance to *Peronospora parasitica* specified by several different *RPP* genes. *Plant Cell*, **8**, 2033–2046.
- Peart, J.R., Lu, R., Sadanandom, A., Malcuit, I., Moffett, P., Brice, D.C., Schauser, L., Jaggard, D.A.W., Xiao, S., Coleman, M.J., Dow, M., Jones, J.D.G., Shirasu, K. and Baulcombe, D.C. (2002) Ubiquitin ligase-associated protein SGT1 is required for host and nonhost disease resistance in plants. *Proc. Natl Acad. Sci. USA*, 99, 10865–10869.
- Penninckx, I.A., Eggermont, K., Terras, F.R., Thomma, B.P., De Samblanx, G.W., Buchala, A., Metraux, J.P., Manners, J.M. and Broekaert, W.F. (1996) Pathogen-induced systemic activation of a plant defensin gene in *Arabidopsis* follows a salicylic acid-independent pathway. *Plant Cell*, 8, 2309–2323.

- Rairdan, G.J., Donofrio, N.M. and Delaney, T.P. (2001) Salicylic acid and NIM1/NPR1-independent gene induction by incompatible Peronospora parasitica in Arabidopsis. Mol. Plant-Microbe Interact. 14, 1235–1246.
- Rehmany, A.P., Lynn, J.R., Tor, M., Holub, E.B. and Beynon, J.L. (2000) A comparison of *Peronospora parasitica* (Downy mildew) isolates from *Arabidopsis thaliana* and *Brassica oleracea* using amplified fragment length polymorphism and internal transcribed spacer 1 sequence analyses. *Fung. Genet. Biol.* **30**, 95–103.
- Reignault, P., Frost, L.N., Richardson, H., Daniels, M.J., Jones, J.D. and Parker, J.E. (1996) Four *Arabidopsis RPP* loci controlling resistance to the Noco2 isolate of *Peronospora parasitica* map to regions known to contain other *RPP* recognition specificities. *Mol. Plant-Microbe Interact.* 9, 464–473.
- Robinson, L.H. and Cahill, D.M. (2003) Ecotypic variation in the response of Arabidopsis thaliana to Phytophthora cinnamomi. Aus. Plant Path. 32, 53–64.
- Roetschi, A., Si-Ammour, A., Belbahri, L., Mauch, F. and Mauch-Mani, B. (2001) Characterization of an *Arabidopsis–Phytophthora* pathosystem: resistance requires a functional *PAD2* gene and is independent of salicylic acid, ethylene and jasmonic acid signalling. *Plant J.* 28, 293–305.
- Ryals, J.A., Neuenschwander, U.H., Willits, M.G., Molina, A., Steiner, H.Y. and Hunt, M.D. (1996) Systemic acquired resistance. *Plant Cell*, 8, 1809–1819.
- Sambrook, J., Fritsch, E.F. and Maniatis, T. (1989) *Molecular Cloning: a Laboratory Manual*, 2nd edn. Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press.
- Sasabe, M., Takeuchi, K., Kamoun, S., Ichinose, Y., Govers, F., Toyoda, K., Shiraishi, T. and Yamada, T. (2000) Independent pathways leading to apoptotic cell death, oxidative burst and defense gene expression in response to elicitin in tobacco cell suspension culture. *Eur. J. Biochem.* 267, 5005–5013.
- Schenk, P.M., Kazan, K., Wilson, I., Anderson, J.P., Richmond, T., Somerville, S.C. and Manners, J.M. (2000) Coordinated plant defense responses in *Arabidopsis* revealed by microarray analysis. *Proc. Natl Acad. Sci. USA*, 97, 11655–11660.
- Schmelzer, E., Naton, B., Freytag, S., Rouhara, I., Kuester, B. and Hahlbrock, K. (1995) Infection-induced rapid cell death in plants: a means of efficient pathogen defense. *Can. J. Bot.* 73(Suppl. 1), S426–S434.
- Smart, C.D. and Fry, W.E. (2001) Invasions by the late blight pathogen: renewed sex and enhanced fitness. *Biol. Invas.* 3, 235–243.
- Smolen, G. and Bender, J. (2002) Arabidopsis cytochrome P450 Cyp83B1 mutations activate the tryptophan biosynthetic pathway. Genetics, 160, 323–332.
- Sogin, M.L. and Silberman, J.D. (1998) Evolution of the protists and protistan parasites from the perspective of molecular systematics. *Int. J. Parasitol.* 28, 11–20.
- Staskawicz, B.J., Ausubel, F.M., Baker, B.J., Ellis, J.G. and Jones, J.D.G. (1995) Molecular genetics of plant disease resistance. *Science*, 268, 661–667.
- Stone, J.M., Heard, J.E., Asai, T. and Ausubel, F.M. (2000) Simulation of fungal-mediated cell death by fumonisin B1 and selection of fumonisin B1-resistant (fbr) *Arabidopsis* mutants. *Plant Cell*, **12**, 1811–1822.
- Takemoto, D., Jones, D.A. and Hardham, A.R. (2003) GFP-tagging of cell components reveals the dynamics of subcellular re-organization in response to infection of *Arabidopsis* by oomycete pathogens. *Plant J.* 33, 775–792.

Vleeshouwers, V.G.A.A., van Dooijeweert, W., Govers, F., Kamoun, S.

and Colon, L.T. (2000) The hypersensitive response is associated with host and nonhost resistance to *Phytophthora infestans*. *Planta*, **210**, 853–864.

- van Wees, S.C., de Swart, E.A., van Pelt, J.A., van Loon, L.C. and Pieterse, C.M. (2000) Enhancement of induced disease resistance by simultaneous activation of salicylate- and jasmonate-dependent defense pathways in *Arabidopsis thaliana*. *Proc. Natl Acad. Sci. USA*, **97**, 8711–8716.
- Wilson, U.E. and Coffey, M.D. (1980) Cytological evaluation of general resistance to *Phytophthora infestans* in potato foliage. *Ann. Bot.* **45**, 91–90.
- Wisman, E. and Ohlrogge, J. (2000) Arabidopsis microarray service facilities. Plant Physiol. 124, 1468–1471.
- Xiao, W., Sheen, J. and Jang, J.C. (2000) The role of hexokinase in plant sugar signal transduction and growth and development. *Plant Mol. Biol.* 44, 451–461.

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