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# Transcript profiling of chitosan-treated Arabidopsis seedlings

Giovanni Povero · Elena Loreti · Chiara Pucciariello · Antonietta Santaniello · Donata Di Tommaso · Gianluca Di Tommaso · Dimos Kapetis · Francesca Zolezzi · Alberto Piaggesi · Pierdomenico Perata

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Abstract In nature, plants can recognize potential pathogens, thus activating intricate networks of defense signals and reactions. Inducible defense is often mediated by the detection of microbe or pathogen associated molecular pattern elicitors, such as flagellin and chitin. Chitosan, the deacetylated form of chitin, plays a role in inducing protection against pathogens in many plant species. We evaluated the ability of chitosan to confer resistance to Botrytis cinerea in Arabidopsis leaves. We subsequently treated Arabidopsis seedlings with chitosan and carried out a transcript profiling analysis using both ATH1 GeneChip microarrays and quantitative RT-PCR. The results showed that defense response genes, including camalexin biosynthesis genes, were up-regulated by chitosan, both in wild-type and in the chitin-insensitive cerk1 mutant, indicating that chitosan is perceived through a CERK1-independent pathway.

**Electronic supplementary material** The online version of this article (doi:10.1007/s10265-010-0399-1) contains supplementary material, which is available to authorized users.

G. Povero · C. Pucciariello · A. Santaniello · P. Perata (⊠) PlantLab, Scuola Superiore Sant'Anna, Piazza Martiri della Libertà 33, 56127 Pisa, Italy e-mail: p.perata@sssup.it

#### E. Loreti

Istituto di Biologia e Biotecnologia Agraria, National Research Council, Via Moruzzi 1, 56100 Pisa, Italy

D. Di Tommaso · G. Di Tommaso · A. Piaggesi Valagro, Zona Industriale, 66041 Atessa, Chieti, Italy

D. Kapetis · F. Zolezzi

Genopolis Consortium, Università di Milano Bicocca, Piazza della Scienza 2-4, 20126 Milan, Italy **Keywords** Arabidopsis · *Botrytis cinerea* · CERK1 · Chitosan · Microarray · Transcript profiling

#### Introduction

Plants can activate defence responses upon the perception and recognition of numerous pathogen-derived molecules (Dangl and Jones 2001; Nürnberger and Scheel 2001; Parker 2003; Zipfel 2009). The detection of pathogens leads to metabolic reprogramming and the production of an array of antimicrobial compounds (Denoux et al. 2008).

Inducible defense against various pathogenic microorganisms is mediated by the detection of microbe or pathogen associated molecular patterns (MAMPs/PAMPs), which are produced by infectious agents but not by host cells (Asai et al. 2002; Nürnberger et al. 2004; Mackey and McFall 2006; Schwessinger and Zipfel 2008). These pathogen-derived molecules trigger a suite of immune responses that limit pathogen growth and damage to the host (Aziz et al. 2004; Ferrari et al. 2007; Denoux et al. 2008). MAMPs perception triggers an early induction of regulatory factors (e.g. transcription factors) and genes that encode enzymes for the synthesis of antimicrobial compounds, together with genes that encode proteins involved in signal perception and transduction, including kinases and phosphatases (Moscatiello et al. 2006; Zipfel et al. 2006).

Chitin, a major component of fungal cell walls, serves as a MAMP (Gust et al. 2007; Libault et al. 2007; Miya et al. 2007; Denoux et al. 2008). Application of purified chitin oligomers to plants or plant cell cultures elicits various defence reactions (Khan et al. 2003; Tanabe et al. 2006). Chitin pre-treatment of plants also reduces the susceptibility to subsequent fungal pathogen challenge (Tanabe et al. 2006). A large number of *Arabidopsis thaliana* genes are induced upon chitin elicitation (Ramonell et al. 2002; Ramonell et al. 2005; Libault et al. 2007; Miya et al. 2007; Wan et al. 2008).

Chitosan, a  $\beta$ -1,4-linked glucosamine, is a deacetylated derivative of chitin that induces phytoalexin synthesis in pea pods (Hadwiger and Beckman 1980), in suspensioncultured soybean cells (Köhle et al. 1984), and parsley cells (Conrath et al. 1989). It also induces the lignification response in wounded wheat leaves (Pearce and Ride 1982), and triggers callose formation in suspension-cultured parsley (Conrath et al. 1989), tomato (Grosskopf et al. 1991) and *Catharanthus roseus* cells (Keen 1975). Moreover, chitosan activates reactive oxygen species (ROS) production (Lee et al. 1999), inhibits fungal growth and elicits a variety of defence reactions in higher plants, therefore enhancing plant resistance against pathogens (Amborabé et al. 2008; Benhamou and Thériault 1992; El Ghaouth et al. 1994).

These abilities highlight the potential importance of chitosan as an elicitor and as a component of host-fungal interactions (Young et al. 1982). For instance, several studies have demonstrated the role of chitosan in protecting different species against Botrytis cinerea (Ben-Shalom et al. 2003; Barka et al. 2004: Trotel-Aziz et al. 2006). Among pathogenic fungi, B. cinerea is the causal agent of grey mould, which is responsible for worldwide yield losses (Aziz et al. 2004). Fungicides can reduce the damage to plants caused by this necrotrophic pathogen, but repeated fungicide applications are harmful to the environment (Siedlecka and Krupa 1996). Furthermore, fungicides have only been partially successful due to the continual appearance and establishment of resistant Botrytis spp. strains to these chemicals (Barka et al. 2004). Consequently, the possibility of stimulating internal plant defences has become an interesting alternative for enhancing natural resistance against B. cinerea and other pathogens. Among the elicitors known to date, chitosan has one of the greatest potentials as a biocontrol agent (El Ghaouth et al. 1994).

Although the role of chitosan in physiological and metabolic processes has been described, transcriptional reprogramming due to chitosan is still largely unknown. Little is known about the mechanisms by which plants can sense chitosan and it is unclear whether the chitin receptor CERK1 could be involved in chitosan perception. Chitosan was recently shown to bind weakly to the chitin receptor CERK1, but this is likely due to the presence of acetylated glucosamine residues (Petutschnig et al. 2010).

In this study, we carried out a transcript profiling analysis in order to evaluate the ability of chitosan to elicit transcriptional defense responses in Arabidopsis seedlings. Microarray results demonstrated the inductive role of chitosan on several genes involved in defense responses. Analysis of the expression of chitosan induced genes using the *cerk1* mutant revealed that chitosan perception is uncoupled from CERK1.

# Materials and methods

Plant materials and growth condition

Arabidopsis thaliana, ecotype Columbia (Col-0) and cerk1-2 (N409189) seeds were purchased from the European Arabidopsis Stock Centre. Seeds were sterilized for 7 min in 1.7% (v/v) bleach solution, and incubated over night in 4% PPM (Plant Preservative Mixture, Plant Cell Technology, Washington, DC, USA) in a full strength sterilized Murashige-Skoog (MS) salt solution (0.5% sucrose) in Hepes-KOH 50 mM (pH 6.8) with gentle shaking. The seeds were then rinsed in abundant sterile water and transferred into 2.5 ml liquid growing media (MS half strength solution in Hepes-KOH 50 mM pH 6.8 containing 0.5% sucrose) with 0.05% PPM in 6-well plates. The plates were incubated in the dark at 4°C for 2 days and finally transferred to continuous light (90 µmol photons/  $m^{-2}$ ) with gentle swirling for 4 days in a plant growth chamber at 22°C. We used adult plants for chitosan pretreatment and B. cinerea inoculation. Plants were grown for 4 weeks in soil, in a growth chamber with a 12-h photoperiod (120  $\mu$ mol photons/m<sup>-2</sup>).

# Chitosan preparation

Chitin was recovered by successive extraction of the concomitant substances from the chitin-containing raw material (exoskeletons of crab, provided by Algea A.S., Kristiansund, Norway). Exoskeletons of crab were washed, dried and pulverized with pestle and mortar into fine powder. Chitin was extracted from the shell by demineralising and deproteinising the powder. Demineralization was achieved using HCl 1.3 M for 2 h. Deproteinization was carried out with NaOH 4 M, at 70°C for 3 h. Depigmentation was achieved using  $H_2O_2$  (1.4 M), for 1.5 h at 70°C. After each phase, the sample was filtered using a G1 gooch filter, to recover the supernatant, which was washed with water until the pH returned to neutrality. Deacetylation was carried out in NaOH (8 M) at 120°C for 3 h (raw material: alkali ratio = 1:5). The molecular weight of the chitosan was determined viscometrically in 0.3 M aqueous acetic acid containing 0.2 M sodium acetate at 30°C. The measurements were performed on an Ubbelohde viscometer with a capillary diameter of 0.3 mm at 30°C. The degree of deacetylation in chitosan samples was measured using conductometric titration. The molecular weight of chitosan obtained was 5-10 kDa and the deacetylation was 80%.

# Plant treatments

Chitosan was used at a concentration of  $150 \mu g/ml$ , whereas oligogalacturonides (OGs) were used at a concentration of 50  $\mu g/ml$ . In some experiments we used the solvent used to solubilise chitosan (final concentration 0.02% acetic acid) as an additional control. Chitosan pretreatment of Arabidopsis adult leaves was performed by spraying a chitosan solution 6 h before the inoculation of *B. cinerea* spores. The samples used for the microarray experiment were 4 days old seedlings treated for 3 h with a chitosan solution (150  $\mu g/ml$ ). Chitosan was solubilised using acetic acid. Consequently, we used an acetic acid solution to treat (3 h) the control samples.

# Expression analysis

The total RNA, extracted using an RNAqueous kit (Ambion, Austin, TX, USA) according to the manufacturer's instructions, was subjected to DNase treatment using a TURBO DNA free kit (Ambion, Austin, TX, USA). Two micrograms of each sample were reverse transcribed into cDNA with a "High capacity cDNA archive kit" (Applied Biosystems, Foster City, CA, USA). Quantitative RT-PCR amplification (qPCR) was carried out using an ABI Prism<sup>®</sup> 7000 Sequence Detection System (Applied Biosystems, Foster City, USA), with primers described in Table S1. Ubiquitin10 (UBQ10) was used as an endogenous control. qPCR reactions were carried out using a SYBR Green PCR Master Mix (Applied Biosystems, Foster City, USA), 50 ng cDNA template, and gene-specific primers in a final reaction volume of 15 µl. The relative quantitation of each single gene expression was performed using the comparative  $C_{\rm T}$  method as described in the ABI PRISM 7700 Sequence Detection System User Bulletin #2 (Applied Biosystems).

RNA isolation, cRNA synthesis, and hybridization to Affymetrix GeneChips

Two independent, replicated experiments were performed for each experimental condition. Each independent experiment consisted of four replicated seedling cultures pooled after RNA extraction. RNA was extracted using an Ambion RNAqueous kit (Ambion, Inc., Austin, TX, USA). RNA quality was assessed by agarose gel electrophoresis and spectrophotometry. RNA was processed for use on Affymetrix GeneChip<sup>®</sup> Arabidopsis ATH1 Genome Arrays, as previously described (Loreti et al. 2005). Hybridization, washing, staining, and scanning procedures were performed by Genopolis (University of Milano-Bicocca), as described in the Affymetrix technical manual. A microarray analysis was performed using an R/Bioconductor (Gentleman et al. 2004). Expression measures were obtained using a GeneChip Robust Multi-Array (Wu and Irizarry 2005), a multi-array analysis method estimating probe set signals, taking into account the physical affinities between probes and targets. Normalization was done using a quantile method (Bolstad et al. 2003). To reduce the number of noninformative genes we applied an interquantile filter (interquantile range, IQR = 0.14). 10,840 probesets passed the filtering. To identify a statistically reliable number of differentially expressed genes among the two conditions, a linear model was performed (Wettenhall and Smyth 2004). To assess a differential expression, an empirical Bayesian method (Smyth and Gordon 2004) was used to moderate the SE of the estimated log-fold changes. Limma, the Bioconductor package that has been used to identify the differentially expressed genes, produces false discovery rate (FDR)-adjusted "p values" that are called "q values". The technique used for multiple testing correction was the Benjamini and Hochberg (Reiner et al. 2003). What is reported in this paper is the Limma FDR "q value" that for clearness where defined as "adjusted p value". The differentially expressed genes (DEG, adjusted P value <0.001) were further filtered by selecting genes showing a fold change ( $\geq 2$  or  $\leq -2$ ) in a hybridization signal (namely the ratio between the intensity in the treated spot vs the control). Microarray datasets were deposited in a public repository with open access (accession no. GSE17193; http://www. ncbi.nlm.nih.gov/projects/geo).

In order to build gene expression maps the MapMan software (Thimm et al. 2004) from http://gabi.rzpd.de/ projects/MapMan/ has been used. Gene categories were used as defined by MapManx BINs.

The chitosan dataset GSE17193 was compared with the publicly available chitin dataset described by Wan et al. (2008) (GSM206278, GSM206277, GSM206279: controls; GSM206274, GSM206275, GSM206276: chitooctaose treatment) using the microarray analysis pipeline described above.

Fungal growth and plant inoculation

*B. cinerea* (as described by Ferrari et al. 2003) was grown on potato dextrose agar plates for 7–15 days at 22°C with a 12-h photoperiod before collecting the spores as described in Ferrari et al. (2007). Rosette leaves from 4-week-old soil grown Arabidopsis plants were placed in Petri dishes containing 0.8% agar, with the petiole embedded in the medium. Inoculation was performed by placing 5  $\mu$ l of a suspension of 5 × 10<sup>5</sup> conidiospores ml<sup>-1</sup> in 24 g l<sup>-1</sup> potato dextrose broth (PDF; Difco, Detroit, USA) on each side of the middle vein. The plates were incubated at 22°C with a 12-h photoperiod for 5 days. High humidity was maintained by covering the plates with a clear plastic dome. Under these experimental conditions, most

Fig. 1 Effect of chitosan on B. cinerea infection of Arabidopsis leaves. a Wild-type (Col-0) Arabidopsis adult leaves were sprayed with a chitosan solution (150 µg/ml) or with the solvent used to solubilise chitosan as a control. After 6 h, leaves were detached and inoculated with B. cinerea spores. Photos of the lesions induced by B. cinerea were taken 5 days after the inoculation of B. cinerea spores. Enlarged pictures were taken using a Nikon TMS-F microscope (type 104) connected with the NIS-elements F2.20 imaging software (Laboratory Imaging, Nikon, Tokyo, Japan). b Lesion areas were measured 5 days after the inoculation of B. cinerea spores. Data are mean of 10 measurements  $\pm$  SD. c Expression analysis of Arabidopsis seedlings treated with active oligogalacturonides (OGs). Fourday old Arabidopsis seedlings were treated with 50 µg/ml OGs for 3 h. WRKY40 mRNA level was analyzed by qPCR. Relative expression levels are shown as Fold Change values (1 = control). Data are means of three replicates  $\pm$  SD. **d** Effect of chitosan on the expression of WRKY40 and Cyp81f2. mRNA levels were measured in 4-day Arabidopsis seedlings treated with water (control), a solution containing the solvent used to solubilise chitosan, and chitosan (150 µg/ml). Relative expression level, measured by qPCR, is shown. Relative expression levels are shown as Fold Change values (1 = control, time 0). Data are means of three replicates  $\pm$  SD

inoculations resulted in rapidly expanding water-soaked lesions of a comparable diameter. Lesion size was determined by measuring their area.

# Peroxidases activity

Arabidopsis leaves were collected after being treated with chitosan, frozen in liquid nitrogen, and stored at  $-70^{\circ}$ C. Samples were extracted by grinding in 2 volumes of icecold sodium phosphate buffer (50 mM, pH 6.5). Homogenates were centrifuged in microfuge tubes at 4°C for 10 min at 14,000g. The supernatant was used to measure enzyme activity. Protein concentration was determined as described by Bradford (1976) with bovine serum albumin (BSA) as the standard. Peroxidase activity was estimated using guaiacol as the substrate. The oxidation of the substrate was measured spectrophotometrically at 450 nm as previously described (Summermatter et al. 1995). The assay solution contained sodium phosphate (10 mM, pH 6), 0.25% (v/v) guaiacol, 100 mM H<sub>2</sub>O<sub>2</sub>, and leaf extract.

# Results

# Chitosan protects Arabidopsis from Botrytis cinerea

Chitosan has been proposed as a strong elicitor of plant defences (Amborabé et al. 2008). We evaluated the ability of this compound to confer resistance to *B. cinerea* in Arabidopsis (Col-0) adult leaves. Four-week-old leaves were inoculated with a *B. cinerea* spore suspension. Lesions induced by *B. cinerea* were measured 5 days after inoculation. Infection of chitosan pre-treated leaves resulted in lesions six times smaller than those on leaves that were not treated with chitosan (Fig. 1a, b). This indicates



that chitosan rapidly stimulates defense reactions in Arabidopsis leaves and inhibits the development of *B. cinerea*.

Chitosan elicits rapid expression of defence-related genes

To test the effect of chitosan at a molecular level, we selected WRKY40 (At1g80840), a transcription factor (TF) involved in the early response to MAMPs (Xu et al. 2006; Denoux et al. 2008). WRKY40 is responsive to oligogalacturonides (OGs), which can increase its mRNA level (Denoux et al. 2008). OGs were found to be very effective in the experimental set-up used in this study (Fig. 1c). We analysed the pattern of WRKY40 expression in a

Category	Gene ID	Gene name	Function (TAIR)	Induction (fold change)
Jasmonate response	At2g24850	TAT3	Tyrosine aminotransferase	786.48
	At3g49620	DIN11	Similar to 2-oxoacid-dependent dioxygenase	282.78
	At1g15520	PDR12	ABC transporter family involved in resistant to lead	126.91
	At1g17380	JAZ5	Jasmonate-ZIM-Domain Protein 5	120.89
	At1g61120	GES/TPS04	Geranyllinalool synthase	107.87
Defence response	At3g44860	FAMT	Farnesoic acid carboxyl-O-methyltransferase. Insect defence	710.95
	At2g30770	CYP71A13	Cytochrome P450. Camalexin pathway	649.08
	At2g26560	PLA IIA	Lipid acyl hydrolase. Pathogen response	338.57
	At1g64160	F22C12.8	Disease resistance-responsive family protein	232.65
	At3g26830	CYP71B15	Conversion of dihydrocamalexic acid to camalexin	187.35
	At1g33960	AIG1	Pathogen response	171.66
	At4g33720	T16L1.210	Pathogenesis-related protein. Pathogen response	107.31
	At2g35980	YLS9	Virus response	105.42
Others	At5g22300	NIT4	Nitrilase isomer. Cianide detoxification	474.12
	At1g26380	T1K7.24	FAD-binding domain-containing protein. Berberine-bridge	402.43
	At1g10585		bHLH transcription factor	211.22
	At4g36700	AP22.80	Cupin family protein	197.91
	At1g80820	CCR2	Cinnamoyl CoA reductase. Lignin biosynthesis	162.69
	At1g26420	T1K7.20	FAD-binding domain-containing protein. Berberine-bridge	139.93
	At4g23700	CHX17	Member of putative Na+/H+ antiporter family	120.53
	At3g46660	UGT76E12	UDP-glucoronosyl/UDP-glucosyl transferase	116.94
	At3g23550	MDB19.3	MATE efflux family protein	113.36
	At5g61160	AACTI	Anthocyanin 5-aromatic acyltransferase 1	111.44
	At5g05600	MOP10.14	Oxidoreductase. 20G-Fe(II) oxygenase family protein	109.97
	At5g20230	BCB	Al-stress-induced gene. Oxidative stress	109.50

Table 1 List of genes induced >100-fold by chitosan (p value <0.001) based on the transcript profiling performed using ATH1 GeneChip microarrays

time-course experiment using chitosan (Fig. 1d). In addition to *WRKY40*, the expression pattern of *CYP81f2* (*At5g57220*), another gene involved in the early response to MAMPs (Clay et al. 2009) was also tested (Fig. 1d). Both *WRKY40* and *CYP81f2* were induced by chitosan after a 2 h treatment, although their expression peaked after 3 h (Fig. 1d) and this time-point was chosen to carry out a transcript profiling analysis.

#### Chitosan elicits a change in the transcript profile

Our microarray results indicated that a treatment with chitosan exerted a dramatic effect on the transcriptome, with 617 probe sets showing an increased expression ( $\geq 2$ -fold, FDR *p* value <0.001) and 388 probe sets indicating a decreased expression ( $\leq -2$ -fold, FDR *p* value <0.001) after data filtering and the selection of differentially expressed genes (Table S2). The list of genes up- regulated >100-fold is reported in Table 1. MapMan (Thimm et al. 2004) highlighted that many chitosan-modulated genes in our experiment are involved in biotic stress responses

(Fig. 2a), including defense-related genes and transcription factors related to hormone signalling (Fig. 2b).

TFs are the upstream triggers in signalling pathways. Chitosan induces TF genes belonging to different families, including ERF, MYB and WRKY domain transcription factor families (Fig. 2b).

In Arabidopsis over 70% of the entire WRKY gene family of 74 members responds to pathogens or pathogen-mimicking stimuli (Lippok et al. 2007). Eight of them responded to chitosan (Fig. 2b). For example, WRKY40 (Fig. 1c, d) is a pathogen-induced WRKY-TF which is also one of the modulators of jasmonate pathways, functioning as an activator of jasmonate-dependent defence (Xu et al. 2006). Among the differentially expressed WRKY-TF factors, we also identified *At5g13080* (*WRKY75*), *At3g01970* (*WRKY45*), and *At2g46400* (*WRKY46*) and *At4g31800* (*WRKY18*) (Table S3). These WRKY transcription factors are involved in pathogen responses (Dong et al. 2003).

The expression of some *MYB* TFs was also affected by chitosan (Fig. 2b). *MYB31* (*At1g74650*), one of the MYB factors known to be responsive to chitin (Libault et al. 2007)



Fig. 2 MapMan screenshots showing the effects of chitosan on the expression of chitosan-modulated genes. **a**, **b** The MapMan software was queried with the list of differentially regulated genes following chitosan treatment (Table S2). *Red shades* indicate induction; *blue shades* indicate repression of gene expression

and *MYB15* (*At3g23250*), a jasmonate responsive gene, were induced by chitosan (Table S3).

At5g47220 (*ERF2*), a member of the AP2/ERF family of transcription factors, was chitosan-induced (Table S3), in line with previous studies demonstrating the involvement of At5g47220 in early chitin responses and jasmonic acid treatment (Libault et al. 2007).

A large number of PR-proteins were induced by chitosan (Fig. 2b). These include *PLP2* (*At2g26560*, induced 338 fold), an integral component of the plant cell death machinery (La Camera et al. 2009), *GH3.12* (*At5g13320*, induced 42-fold), a gene involved in the regulation of salicylic acid (SA) metabolism (Nobuta et al. 2007), *AIG1* (*At1g33960*, induced 171-fold) induced by avirulence genes *avrRpt2* and *RPS2* after infection with *Pseudomonas syringae* (Wang and Li 2009), and the plant defensin *PDF1.2B* (*At2g26020*, induced 95-fold).

Among the enzymes involved in biotic stress responses, we found a large group of peroxidases (POXs; Figs. 2b, 3a), usually considered enzymes involved in detoxification of ROS (Kawano 2003). The differential expression of



Fig. 3 Effect of chitosan on peroxidases. a Effect of chitosan on the induction of genes encoding peroxidases or enzymes with peroxidase activities in the microarray experiment. b Peroxidase activity after chitosan treatment. Results are mean  $\pm$  SD of three replicates

genes encoding for peroxidases was mirrored by a corresponding increase in enzyme activity (Fig. 3b).

Chitosan induces genes involved in camalexin biosynthesis

We examined the transcriptional reprogramming induced by chitosan and compared it with the effects of *B. cinerea*. Arabidopsis eFP browser software (Winter et al. 2007) allowed us to identify a set of genes inducible after *B. cinerea* inoculation (Ferrari et al. 2007), which are also induced by chitosan treatment. The results of this comparison are summarized in Table S4.

Interestingly, this list includes genes involved in the biosynthesis of camalexin, the indole phytoalexin of *Arabidopsis thaliana* (Fig. 4a). Camalexin biosynthesis is essential for resistance to *B. cinerea* in Arabidopsis (Ferrari et al. 2007). Almost all the genes codifying for enzymes which are involved in the biosynthesis of tryptophan, the precursor of camalexin, such as *At5g05730* (*ASA1*), *At5g17990* (*PAT1*), *At2g04400* (*IGPS*), and *At3g54640* (*TSA1*) were induced (Fig. 4a). Camalexin is synthesized from tryptophan via indole-3-acetaldoxime (IAOx) in a



Fig. 4 Effects of chitosan on the camalexin biosynthetic pathway. a Chitosan-dependent differentially expressed genes encoding enzymes involved in the camalexin pathway are shown. The fold change measured in the microarray experiment is shown. b Venn diagram showing the overlap between genes differentially regulated by chitin (Wan et al. 2008) and chitosan (this study). c mRNA levels for *Cyp71A13* and *Cyp71B15* were measured in 4-day Arabidopsis wild-type (Col-0) or *cerk1* mutant seedlings treated with water (control), a solution containing the solvent used to solubilise chitosan, and chitosan (150 µg/ml). Relative expression level, measured by qPCR, is shown. Relative expression levels are shown as Fold Change values (1 = control, time 0). Data are means of three replicates  $\pm$  SD

reaction catalyzed by CYP79B2 and CYP79B3 (Glawischnig et al. 2004). *At4g39950/CYP79B2* was induced fivefold by chitosan (Fig. 4a). In the camalexin pathway, IAOx represents the direct precursor of this compound (Glawischnig et al. 2004). The transcripts encoding *At2g30770* (*Cyp71A13*) and *At3g26830* (*Cyp71B15*, *PAD3*) showed a very high induction after chitosan treatment (Fig. 4a). Cyp71A13 catalyzes the first committed step in camalexin biosynthesis by dehydrating IAOx to indole-3-acetonitrile (IAN; Nafisi et al. 2007). *Cyp71B15/ PAD3* on the other hand, encodes an essential enzyme in the camalexin biosynthetic pathway (Zhou et al. 1999), which catalyzes the oxidative decarboxylation of dihydrocamalexic acid to camalexin (Schuhegger et al. 2006).

We also observed a fourfold induction of At3g52430 (*PAD4*) (Table S2), which is required for the bacterial, but not fungal, induction of camalexin (Jirage et al. 1999; Ferrari et al. 2003).

# Chitosan perception is independent of CERK1

A comparison of the transcription profiles of Arabidopsis plants treated with chitin (Wan et al. 2008) or with chitosan (this study), revealed that 33.7% of the genes modulated by chitosan are also regulated by chitin (Fig. 4b; Table S5). This was suggestive of limited overlap with the targets of chitin.

To evaluate whether the effects of chitosan were indeed largely independent from the chitin signalling pathway, we analysed the expression of *CYP71A13* and *Cyp71B15* in both wild type Arabidopsis and in the chitin-insensitive *cerk1* mutant (Miya et al. 2007). Chitosan treatment strongly induced these two genes in both the wild-type and *cerk1* (Fig. 4c).

Chitin triggers the regulation of 118 transcription factors (Libault et al. 2007). We identified 49 TF genes up-regulated (more than twofold) after chitosan treatment (Table S3). Only 22 of the 118 TFs regulated by chitin (Libault et al. 2007) were found to be responsive to chitosan (Table S3). These results further suggest that distinct pathways operate following chitin or chitosan treatments.

Some transcription factors are responsive to both chitin (Libault et al. 2007; Wan et al. 2008) and chitosan (Table S3) (e.g. WRKY18, WRKY40, WRKY46 and ERF2) while others are induced by chitosan (Table S3) but not by chitin, e.g. WRKY75 and WRKY45 (Libault et al. 2007; Wan et al. 2008). We tested the induction of these TFs in the *cerk1* mutant. The results indicated that (i) chitin-insensitive TFs are induced normally in the *cerk1* mutant and that (ii) also chitin-sensitive TFs are responsive to chitosan in *cerk1* (Fig. 5). The first point confirms that chitosan activates a CERK1-independent signalling pathway, while the second point indicates that a convergence downstream of CERK1 is possible.

#### Discussion

Chitosan induces defense responses in several plant species. Our findings demonstrated that chitosan protected



Fig. 5 Effects of chitosan on the expression of transcription factors. mRNA levels were measured in 4-day Arabidopsis wild-type (Col-0) or *cerk1* mutant seedlings treated with water (control), a solution containing the solvent used to solubilise chitosan, and chitosan (150  $\mu$ g/ml). Relative expression level, measured by qPCR, is shown. Relative expression levels are shown as Fold Change values (1 = control, time 0). Data are means of three replicates  $\pm$  SD

Arabidopsis leaves against the necrotrophic fungus B. cinerea (Fig. 1). This is because chitosan was perceived by the plant as a powerful elicitor, acting as a pathogenmimicking stimulus, as demonstrated by the activation of a large number of biotic-stress related genes (Fig. 2). The application of a chitosan solution may sensitize Arabidopsis leaves to respond more rapidly to pathogen attack. For example, camalexin biosynthetic genes were strongly transcriptionally up-regulated after a chitosan treatment (Fig. 4b, c). Some peroxidases were also up-regulated. Extracellular, secreted plant POXs can catalyze the generation of ROS coupled to defense-related compounds, such as salicylic acid (SA), aromatic monoamines (AMAs) and chitooligosaccharide (COSs) elicitors (Kawano 2003). We found that the effect of chitosan at the transcriptional level was mirrored by a corresponding increase in peroxidase activity, indicating that the effects at the transcriptional level are accompanied by an increase in the level of the corresponding proteins.

Whether chitosan is perceived by CERK1, the chitin receptor is under debate. CERK1 was found to bind weakly chitosan (Petutschnig et al. 2010), and this evidence corroborated the idea that CERK1 is involved in chitosandependent plant defense activation (Petutschnig et al. 2010). The cerk1-2 mutant is insensitive to partially de-acetylated chitosan in terms of ROS-burst and MAP kinase assays, suggesting that either CERK1 is able to recognize chitosan or that acetylated residues present in commercially available chitosan are recognized by CERK1 (Petutschnig et al. 2010). Iizasa et al. (2010) found very weak binding of chitosan to CERK1, a result suggestive of the existence of a CERK1-independent pathway for chitosan perception. Fully de-acetylated chitooligomers are indeed not effective and this is highly suggestive of the importance of GlcNAc residues for CERK1 binding (Petutschnig et al. 2010).

In this context, a comparison of the transcription profiles of Arabidopsis plants treated with either chitin (Wan et al. 2008) or with chitosan (this study), revealed that the overlap in modulated genes is relatively limited, with only 33.7% of the genes modulated by chitosan also regulated by chitin (Fig. 4b; Table S5).

More direct evidence for a CERK1-independent chitosan signalling arose from the use of the *cerk1* mutant. Transcription factors and the key-genes for camalexin biosynthesis (*CYP71A13* and *Cyp71B15*) are responsive to chitosan in the *cerk1* mutant, demonstrating that chitosan perception is independent of CERK1 and thus from the chitin signaling pathway.

The signaling pathways for chitin and chitosan thus appear to be independent, but may converge downstream of CERK1 (Fig. 6). Chitin perception by CERK1 activates transcription factors that induce the expression of genes



Fig. 6 Schematic representation of the chitosan signalling pathway and its relation with chitin signalling. Chitin is perceived by CERK1, while chitosan appears to trigger defense-related genes through a CERK1-independent pathway. The two pathways may converge at the level of TFs such as WRKY18, WRKY40 and WRKY46. Both pathways result in the induction of resistance to pathogens

directly involved in plant defence to pathogens, and which contribute to a resistant phenotype. Chitosan is perceived by a still unknown receptor, activating a downstream signaling pathway that largely converges with that induced by chitin. Some transcription factors are known to be induced by both chitin and chitosan (e.g. WRKY18, WRKY40, WRKY46) and demonstrate convergence of the two pathways. The identification of the chitosan receptor is of considerable interest and certainly deserves further studies.

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