

Oligogalacturonide production upon *Arabidopsis* thaliana—Botrytis cinerea interaction

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Despite an ever-increasing interest for the use of pectin-derived oligogalacturonides (OGs) as biological control agents in agriculture, very little information exists-mainly for technical reasons-on the nature and activity of the OGs that accumulate during pathogen infection. Here we developed a sensitive OG profiling method, which revealed unsuspected features of the OGs generated during infection of Arabidopsis thaliana with the fungus Botrytis cinerea. Indeed, in contrast to previous reports, most OGs were acetyl- and methylesterified, and 80% of them were produced by fungal pectin lyases, not by polygalacturonases. Polygalacturonase products did not accumulate as larger size OGs but were converted into oxidized GalA dimers. Finally, the comparison of the OGs and transcriptomes of leaves infected with B. cinerea mutants with reduced pectinolytic activity but with decreased or increased virulence, respectively, identified candidate OG elicitors. In conclusion, OG analysis provides insights into the enzymatic arms race between plant and pathogen and facilitates the identification of defense elicitors.

oligogalacturonides \mid plant-pathogen interaction \mid Arabidopsis thaliana \mid Botrytis cinerea \mid pectin lyase

The cell wall forms the first line of defense in the interaction of plants with their microbial environment. Cell walls consist of complex polysaccharide networks, which combine multiple functional properties such as strength (to resist turgor pressure), extensibility (to allow growth), as well as protection against microbial attack. Such protection involves so called "basal immunity," which is triggered upon recognition of pathogen-associated molecular patterns but also of plant-derived molecules associated with infection, referred to as damage-associated molecular patterns (DAMPs). The latter include cell wall-derived oligosaccharides with elicitor activity (also called oligosaccharins) (1) that are produced during infection with microbes and, in particular, necrotrophic bacteria and fungi, which feed on cell walls and have large arsenals of cell wall-degrading enzymes (2).

A major source of cell wall-derived DAMPs are pectins. The pectic polymer, homogalacturonan (HG) for example, represents in Arabidopsis 20% of the wall of growing cells. HG is a linear polymer of α-1-4-linked galacturonic acid (GalA) residues secreted in a methylesterified (on C6) and often acetylesterified (on C2 and/or C3) form (3), which can be enzymatically deacetylesterified by pectin acetylesterases and demethylesterified by pectin methylesterases (PMEs) in muro (4). Demethylesterification exposes the polymer to degradation by polygalacturonases (PGs) and pectate lyases (PLs, that cleave unmethylesterified HG by β-elimination, thus generating an unsaturated bond at the nonreducing end), which can be of plant or fungal origin. HG demethylesterification plays a key role in the control of cell wall rheology underlying plant growth (5). In addition, HG turnover generates oligogalacturonides (OGs), some of which act as DAMPs or, in certain conditions, may have a signaling role in development. Indeed, treatment with unmethylesterified OGs of degree of polymerization (DP) 10 to

15 elicits strong immune responses but can also promote tomato fruit ripening by triggering ethylene production (6–12). In addition, unmethylesterified GalA trimers or tetramers can promote immunity and may have a role in skotomorphogenesis (13–15). Despite considerable interest for OG signaling in plant immunity and the commercial use of OGs as natural defense stimulators, very little is known about the OGs that are actually produced during fungal infection. So far, only 1 study reported on such in vivo-produced OGs (16). This study revealed the structure of a few pectin fragments in tomato fruit infected with the necrotrophic fungus *Botrytis cinerea* that causes gray mold disease. The production of these OGs implied the activity of PGs, PMEs and, surprisingly, pectin lyases (PNLs), fungal enzymes that cleave methylesterified HG through a β -elimination mechanism. The study did not address potential elicitor activities of these OGs.

Here we developed a sensitive analytical method based on the separation of oligosaccharides combined with accurate determination of their sizes and acetylesterification and methylesterification patterns using MS/MS. This method reveals the global OG composition of plant samples. Surprisingly, the analysis of the OGs during different stages of *B. cinerea* infection of *Arabidopsis thaliana* leaves failed to detect the accumulation of the accumulation of the unmethylesterified DP10 to -15 OGs that were previously shown to have elicitor activity. Instead, despite the abundance of PGs in *B. cinerea*, 80% of the OGs were the product of fungal PNLs, 1 of which was characterized in vitro and shown to cleave highly methylesterified

Significance

Oligogalacturonides (OGs), oligomers of α -1,4-linked galacturonic acid generated during plant infection by necrotrophic fungi, have been proposed to trigger plant basal immunity when they are applied exogenously to plant tissues. However, very little information exists on the structure and activity of the OGs that really accumulate during pathogen infection. This report reveals features of OGs generated during infection of Arabidopsis thaliana with the fungus Botrytis cinerea and identifies potential candidate OG elicitors.

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Data deposition: Affymetrix Microarray data have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo/ (accession no. GSE120933).

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HG. We also found that most nonmethylesterified PG products were oxidized in muro, probably by recently described plant oxidases (17). Finally, the comparison of the OGs generated by a collection of B. cinerea mutants affected in pectinolysis led to the identification of specific PNL-produced OGs, the accumulation pattern of which was associated with reduced virulence and a specific gene-expression pattern in planta. These findings illustrate the evolutionary enzymatic arms race between plant and fungus, where the 2 partners, respectively, aim for detection and hiding of pathogen-associated oligosaccharide cues.

Development of a Sensitive Separative Method for OG Characterization. In a first experiment, OGs were analyzed in the hydrolysis products of commercial citrus pectins (degree of methylesterification [DM] 60) (18) digested for 20 h with Aspergillus aculeatus PG. OGs were separated using hydrophilic interaction liquid chromatography (HILIC) (19-21), and characterized using high-resolution mass spectrometry (HR-MS) in negative mode. Although a number of methylesterified and nonmethylesterified GalA oligomers originating from the HG backbone could be distinguished in the HILIC elution pattern (SI Appendix, Fig. S1A), non- or poorly methylesterified OGs were insufficiently separated in broad peaks, resulting in low MS sensitivity. To improve the separation power,

we developed a high-performance size-exclusion chromatography (HP-SEC)-MS-based method, which yielded sharper peaks (a few

seconds), resulting in a higher signal-to-noise ratio relative to the

HILIC separation (Fig. 1A and SI Appendix, Fig. S1B). This method

allowed the DP and methylesterification pattern of each OG to be determined (up to DP16) (SI Appendix, Fig. S1C) based upon accurate mass annotation, isotopic pattern, and MS/MS analysis (SI Appendix, Fig. S2 and Table S1).

It was surprising to see that the retention time was not only determined by the DP but also by the methylesterification status of the OG (Fig. 1A and SI Appendix, Table S1), methylesterified OGs eluting later (corresponding to a smaller hydrodynamic volume) than their unmethylesterified counterparts. In contrast, the presence of the bulkier acetylester groups did not affect the retention time as shown by the analysis of OGs produced from highly acetylesterified sugar beet pectins (Fig. 1B and SI Appendix, Table S1). This indicates that methylesterification, unlike acetylesterification, can reduce the hydrodynamic volume of the OGs. The structural basis for this effect was revealed by the analysis of OGs that did not follow this rule (Fig. 1C and SI Appendix, Fig. S3 and Table S1), such as GalA₃Me₂, which coeluted with, not after, GalA₃Me (GalA-GalAMe-GalA) and GalA₄Me₃, which coeluted with GalA₄Me₂ (GalA-GalAMe-GalAMe-GalA). Comparison of the structures (Fig. 1C) showed that OGs with 2 adjacent unmethylesterified GalAs systematically eluted earlier than OGs of the same size lacking this feature. The increased hydrodynamic volume can therefore be attributed to the electrostatic repulsion between 2 adjacent carboxylic functions, making the OG more extended (Fig. 1C).

OG Analysis to Characterize Pectinolytic Activities of B. cinerea. The B. cinerea genome encodes a large array of putative pectinolytic enzymes: 13 PGs, 5 PMEs, 2 PLs, and 5 PNLs. Knockout mutants

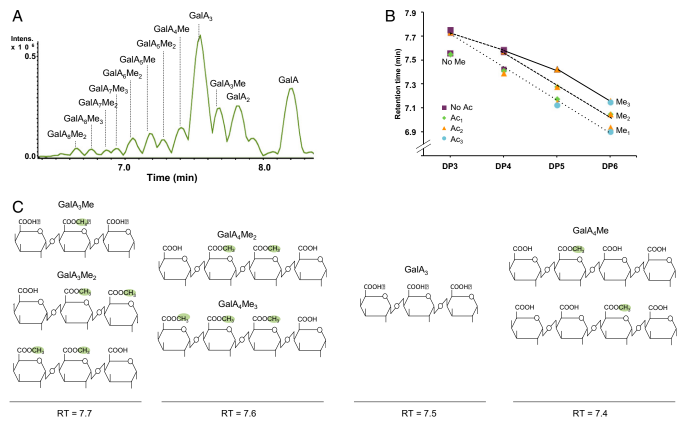


Fig. 1. A sensitive and HP-SEC method for separating complex mixes of OGs. (A) Base peak chromatogram obtained by LC-MS analysis in negative ionization mode of various methylesterified OGs obtained following digestion of citrus pectins by commercial PG from A. aculeatus. (B) Impact of the presence of methyland acetylesters on the retention time (RT) of OGs. OGs were obtained from sugar beet pectins digested by the commercial A. aculeatus PG. (C) Comparison of the structures of methylesterified OGs with a DP from 3 to 4 released by A. aculeatus PG and their respective RT in minutes. Positions of methyl groups are indicated according to the HG structure (3). OGs are named GalAxMevAcz. Subscript numbers indicate the DP and the number of methyl- and acetylester groups, respectively. Ac, acetylester group; Intens., signal intensity; Me, methylester group. Green color indicates the position of methylester groups.

were obtained for 6 PGs (22) (BcPG1 to -6) and 2 PMEs (BcPME1 and -2) (23). Their analysis showed that BcPG1 and -2 are necessary for full virulence on various plant species (24, 25), while BcPME1 and -2 are dispensable for normal virulence, at least on tomato and grapevine leaves or pear fruit (23). The activities of the 5 PGs (BcPG1 to -4, and -6) were previously characterized on polygalacturonic acid (25). Here, we used a more complex methylesterified substrate (citrus pectins, DM60) to study in more detail the activity of 2 PGs, BcPG2 and -3, representative of single-attack (nonprocessive) and multiple-attack (processive) enzymes, respectively. The former cleaves only once after formation of the enzyme–substrate complex and releases OGs of varying DPs, whereas the latter attacks the substrate multiple times and releases GalA and short OGs already from the start of the reaction (SI Appendix, Fig. S4).

We first purified both PGs from recombinant Pichia pastoris liquid cultures and incubated the enzymes with citrus pectins. After 20 h of digestion, both PGs produced mainly GalA, nonmethylesterified dimers (GalA₂), and trimers (GalA₃) (Fig. 2A), as well as small amounts of methylesterified OGs. As expected, whereas the nonprocessive enzyme BcPG2 generated mainly GalA₂ and GalA₃, the processive BcPG3 produced primarily GalA and GalA2. Such processive PG activity requires fully demethylesterified HG stretches, which are generated by (processive) plant PMEs. The presence of methylesterified OGs (up to 3 methylester groups in GalA₅Me₃ or GalA₆Me₃), however, indicates that both PGs can also cleave partially demethylesterified stretches. We next studied the combined activity of BcPME1 and BcPG2 or BcPG3. BcPME1 enhanced almost 5 times the production of GalA, GalA₂, and GalA₃ by BcPG2 and completely suppressed the accumulation of methylesterified OGs (Fig. 2A and SI Appendix, Fig. S4). In contrast, BcPME1 had a much more limited impact on the overall BcPG3 activity but promoted a shift from GalA + GalA₂ toward GalA₂ + GalA₃ for the most abundant OGs. Interestingly, some higher DP OGs, including GalA₆Me₃, resisted further digestion (Fig. 2A and SI Appendix, Fig. S4), suggesting that this OG is a poor substrate for both BcPME1 and BcPG3. Together, these results confirm that nonprocessive BcPG2 requires—and BcPG3 has a preference for—substrates of at least DP4 (25), but also show that their substrates can be partially methylesterified. In addition, they show that nonprocessive BcPME1 greatly enhances the number of cleavage sites for nonprocessive PG activities but, as expected, does not enhance the processive PG activity of BcPG3. Finally, they show that BcPME1 demethylesterifies both the HG polymer and OGs, even the trimers, except for GalA₆Me₃, which, for an unknown reason, is more recalcitrant to the enzyme.

We next used OG analysis to study the global *B. cinerea* pectinolytic activity. To this end, citrus pectins were incubated with *B. cinerea* spores and the OGs were analyzed over time. After 6 h of incubation, only a very small amount of OGs was detected (Fig. 2*B*). After 16 h, primarily GalA₂ and GalA₃ but also methylesterified OGs from DP4 to -7 had accumulated. Since purified BcPG2 and BcPG3 did not produce such methylesterified oligomers in large amounts, the production of these OGs most likely involved other pectinolytic enzymes. At 24 h, the relative amount of GalA₂ had increased and that of methylesterified OGs from DP4 to -7 had decreased. This confirms that *B. cinerea* PMEs and PGs can use OGs as a substrate.

OGS Released upon *B. cinerea***–***A. thaliana* **Interaction.** So far, we have shown the activity of *B. cinerea* and 3 of its purified enzymes on citrus pectins, which is a complex but inert substrate. We next investigated how the interaction with living plant cells affects the pectinolytic activity of the fungus. Previous studies have shown that plants produce defensive PG-inhibiting proteins (PGIPs) that attenuate the activity of fungal PGs (26). This is thought to promote the accumulation of large unmethylesterified OGs with elicitor activity (27), as observed for instance in *A. thaliana* upon coexpression of a PGIP with a fungal PG. However, very little is known on the OGs that are actually produced during the interaction of *B. cinerea* with its host.

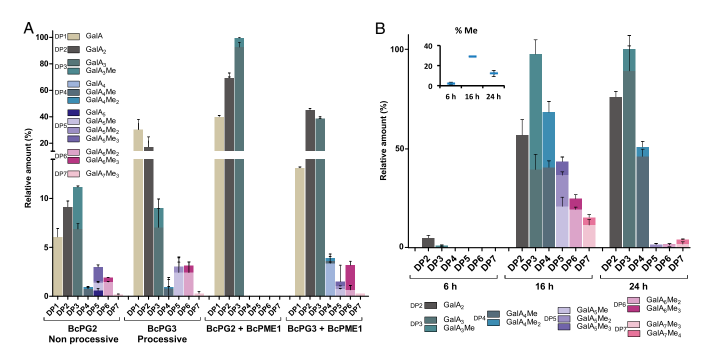


Fig. 2. OG analysis to characterize pectinolytic activities of *B. cinerea* on commercial citrus peel pectins. (*A*) Characterization of OGs by HP-SEC-MS released by BcPG2 and BcPG3 in the presence or absence of BcPME1 after 20 h of incubation with citrus pectins. Data are means \pm SD; n = 3. (*B*) Kinetics of OGs produced by WT *B. cinerea* strain over time from citrus pectins and evolution of the OG methylesterification rate over time. Data are means \pm SD; n = 3. OGs are named GalA_xMe_y. Subscript numbers indicate the degree of polymerization and the number of methylester groups.

Here we studied the OGs of A. thaliana leaves incubated with B. cinerea spores. The incubation was done in liquid medium since this facilitates the collection of large quantities of OGs. Under these conditions, the first plant symptoms can be observed between 9 and 12 h postinfection (hpi) (SI Appendix, Fig. S5). We chose 20 hpi to collect the medium, when significant maceration of the tissue was observed. After concentration, the medium was analyzed by HP-SEC. Whereas no OGs were detected in the control medium (SI Appendix, Fig. S6A), a complex OG mixture was observed upon infection with the spore suspension, which could be analyzed using in vitro-released OGs as standards. Surprisingly, the most abundant OGs detected in spore suspension at 20 hpi differed by 18 mass units (H₂O) from OGs released from citrus or sugar beet pectins (SI Appendix, Table S2). This difference was attributed to the loss of a water molecule between C4 and C5 of the nonreducing end GalA residue during the β-elimination reaction catalyzed by PNL (28). These OGs were named GalA_xMe_yAc_z-H₂O. We conclude that during infection, only 20% of the accumulated OGs were actually produced by PGs, the remaining 80% being the product of PNLs (Fig. 3A). This is in striking contrast with the OGs released by B. cinerea grown on citrus pectins, in which no PNL product was found (Fig. 2B).

Among the PNL products, 20 different OGs of DPs ranging from 3 to 10 were identified (Fig. 3B). The main products displayed a DP of 4 and 5, the GalA₄MeAc-H₂O being the most abundant, representing 25% of the OGs. To determine the precise position of the acetyl- and methylester groups on the GalA₄MeAc-H₂O, we analyzed the fragmentation pattern (SI Appendix, Fig. S7). The acetylester was only found on the A- and C-fragments, corresponding to the nonreducing end, but never on the Z-fragments, which correspond to the reducing end. Moreover, the ring cleavage product ${}^{0,2}A_1$ at 157.0051 m/z proves that the acetylesterification occurs on the C3 of the GalA at the nonreducing end. It is also worth noting that all of the PNL-generated OGs contained at least 1 GalAMe for 2 GalAs, some of them also being acetylesterified.

The PG-derived OGs were mainly of DP2 and -3. Surprisingly, DP3 OGs were all methylesterified and the bulk of the DP2 OGs was not GalA₂ but an ion at m/z 385, differing by 16 mass units (oxygen) from GalA₂ (369) (Fig. 3C and SI Appendix, Table S2). The MS/MS fragmentation pattern of this ion showed 2 major fragments corresponding to a uronic acid linked to a galactaric acid, which is a C1-oxidized uronic acid (Fig. 3D). To determine the origin of this oxidized GalA₂ (GalA₂ox), we performed a kinetic study of GalA₃, GalA₂, and GalA₂ox from 12 to 18 hpi. Whereas the amount of all 3 OGs increased from 12 to 15 hpi, GalA₃ and GalA₂ progressively disappeared in favor of GalA₂ox from 15 to 18 hpi (Fig. 3E). We assume that GalA₃ and GalA₂ are oxidized by a plant oxidase, as described recently (17). We also detected other oligosaccharides, the structure and origin of which were not investigated here.

The large proportion of PNL-derived OGs and the absence of long unmethylesterified OGs in infected leaves was surprising. To rule out that the OG profile was an artifact of the nonphysiological liquid culture system, we also analyzed the OGs released in spore suspension drops incubated for 24 h on detached A. thaliana leaves. Besides higher GalA₂ox levels, this shows OG structures that were similar to those detected in the culture medium of immersed leaves, albeit with differences in relative abundance of the OGs (SI Appendix, Fig. S6 A and B). These quantitative differences could reflect at least in part the less-advanced infection stage relative to that of immersed leaves. Then we incubated cell wall from A. thaliana leaves with B. cinerea spores and analyzed OGs produced over time. We showed that they all result from a PNL activity and not from a PG activity, suggesting that the accumulation of a large proportion of PNL-derived products during infection was due to A. thaliana pectin structure. It is worth noting that the main shortest OG was a DP6 and not a DP4, as observed during infection (SI Appendix, Fig. S8), suggesting that some plant enzyme activities might contribute to OG production upon infection.

To rule out that large OGs remained attached to the cell wall, we extracted residual OGs from the cell wall with a chelating agent. In these conditions only a major GalA20x peak was detectable (SI Appendix, Fig. S6C,) confirming that long and unmethylesterified OGs did not accumulate during infection. Finally, to determine why long and unmethylesterified OGs were not accumulated, we incubated OGs from DP3 to -15 with B. cinerea pregerminated spores and analyzed OGs over time. Surprisingly, as soon as the germinated fungus is in contact with these OGs, they were instantaneously hydrolyzed into shorter OGs (DP2 to -5) and to GalA, suggesting that GalA₆ to GalA₁₅ are too easily hydrolysable substrates for a B. cinerea arsenal to be accumulated during plant infection (Fig. 3F).

We next produced semipurified OG preparations from WTinfected leaves, enriched for GalA₄MeAc-H₂O (Fig. 3G and SI Appendix, Fig. S9) and which completely lacked the previously identified demethylesterified OG elicitors DP10 to -15 and DP3. GalA₄MeAc-H₂O represents 48% of total semipurified OGs from WT-infected leaves. The concentration of GalA₄MeAc-H₂O was determined by assuming that GalA3 and GalA4MeAc-H2O response factors are equal. Incubation of A. thaliana leaves with OGs from infected leaves containing 5 µM of GalA₄MeAc-H₂O induced, within 3 h, a much higher expression of a defense reporter gene (pAtPGIP1::GUS [β-glucuronidase]) than in leaves and in hypocotyls incubated with GalA₃ (200 µM) preparation (Fig. 3 H and I). However, after a longer time of incubation (6 h), OG3 preparation induced a similar expression of AtPGIP1 in etiolated hypocotyls except in vessels (Fig. 3H). This result supports the hypothesis that the plant may rapidly trigger its defense response upon recognition of PNL products,

A Fungal PNL Contributes to OG Production during Infection. We assumed that the PNL activity detected came from B. cinerea, since so far only PL and no PNL activity had been observed in plants (4). We therefore selected, from published transcriptome profiles (29), the most abundant transcript, encoding a putative PNL in germinating B. cinerea conidia (BcPNL1) to express the enzyme in P. pastoris. The enzyme, purified from the culture medium (SI Appendix, Fig. S10), showed PNL activity on various commercial substrates with a sharp optimum at pH 8 (Fig. 4A) and a preference for highly methylesterified pectins (Fig. 4B). To assess the potential role of this PNL in the production of unsaturated OGs during infection, we analyzed the OGs produced by this enzyme from A. thaliana leaf cell walls by HP-SEC-MS. We detected OGs from DP4, all methylesterified and some also acetylesterified (Fig. 4C and SI Appendix, Fig. S11). Among these OGs, 15 were also found in B. cinereainfected A. thaliana leaves (Fig. 4C and SI Appendix, Fig. S11), while the other OGs were more methylesterified. Together, these results are consistent with the involvement of BcPNL1, in particular in the initial steps of OG production during *B. cinerea* infection.

Relative Contribution of the Main PG and PME Activities in Pectin Degradation during B. cinerea Growth. To gain better insights into OG production during A. thaliana leaf infection and to understand the role of these OGs in plant defense activation, we took advantage of B. cinerea mutants altered for each 1 of the 6 PGs and the double-mutant Bcpme1/2 (22, 23). Unfortunately, we failed so far to produce a BcPNL1 mutant. gRT-PCR analysis showed that all of the PG and PME genes were expressed in WT B. cinerea grown in the presence of citrus pectins (SI Appendix, Fig. S12) after 6 and 16 h of incubation. We next analyzed the size of citrus pectins, using SEC, after 6-h incubation with the WT strain or each of the *Bcpg* and *Bcpme* mutants. Interestingly, multiple-angle light scattering, refractive index detection, and viscosimetry analyses (SI Appendix, Fig. S13) showed a larger size of digested pectins for all of the mutants relative to the WT strain, indicating that all of the investigated enzymes contributed to pectin digestion.

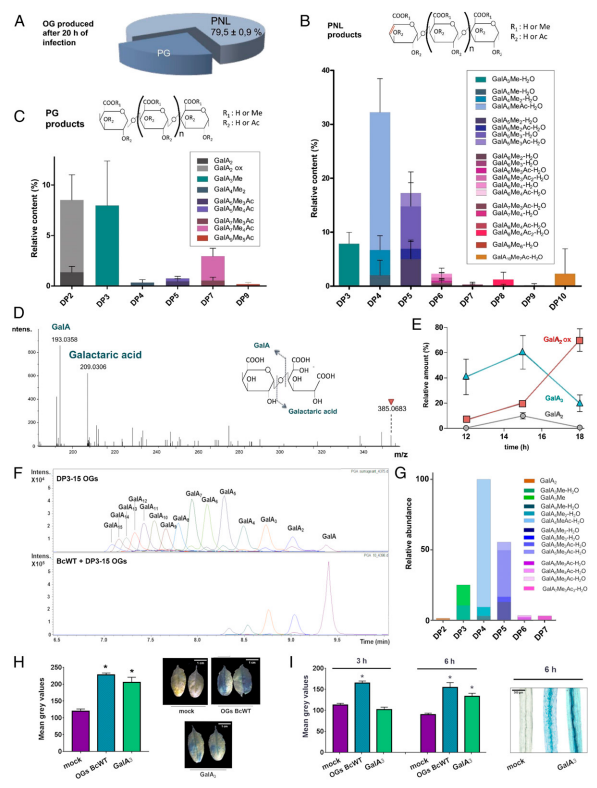


Fig. 3. OGs released upon *A. thaliana–B. cinerea* interaction. (*A*) Proportion of pectin lyase products among total OG produced after 20-h infection of *A. thaliana* leaves by *B. cinerea*. Data are means \pm SD; n=3. (*B* and *C*) Characterization of OGs produced after 20-h infection by HP-SEC-MS. (*B*) PNL products; (*C*) PG products. Data are means \pm SD; n=3. (*D*) MS² fragmentation pattern of oxydized GalA₂ oligomer (m/z 385.0683) eluting at 7.8 min in Fig. 1*A* using HP-SEC with online MS. (*E*) Kinetics of GalA₃, GalA₂, and oxydized GalA₂ upon infection. Data are means \pm SD; n=3. (*F*) DP10 to -15 OGs degradation by *B. cinerea*. OGs (2 mg/mL) were mixed with germinated *B. cinerea* spores and the reaction immediately stopped by addition of a volume of ethanol before analysis by HP-SEC-MS. (*G*) Composition of semipurified OGs from leaves infected by WT *B. cinerea* strain. (*H*) GUS quantification and staining of *A. thaliana* leaves expressing the promoter *PGIP1::GUS* fusion after the infiltration of GalA₃ (200 μM) and semipurified OGs containing 5 μM of GalA₄MeAc-H₂O. (*I*) GUS quantification and staining of *A. thaliana* dark-grown hypocotyls expressing the promoter *PGIP1::GUS* fusion after the infiltration of GalA₃ and semipurified OGs from leaves infected by WT *B. cinerea* strain. OGs are named GalA_xMe_yAc_z. Subscript numbers indicate the degree of polymerization and the number of methyl- and acetylester groups, respectively. Asterisks directly above bars denote statistical significance compared with mock treatment (**P* < 0.05).

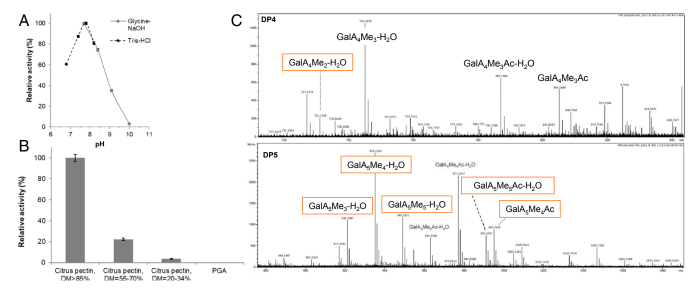


Fig. 4. BcPNL1 is active on highly methylesterified pectins and at alkaline pH. (A) PNL activity was measured at 235 nm at different pH, with glycine-NaOH and Tris-HCl buffers. Data represent the mean ± SE of 3 replicates. (B) PNL activity was measured at 235 nm using substrates with various DMs. Data represent the mean ± SE of 3 replicates. (C) OGs of DP4 and -5 released by BcPNL1 from A. thaliana leaf cell wall. Boxed OGs are found in infection OGs.

We also analyzed the OGs released after 16 and 24 h of B. cinerea mutant growth on citrus pectins. Principal component analysis (PCA) showed that after 16 h of growth Bcpg1 and Bcpme1/2 samples were separated from others (SI Appendix, Fig. \$144), and that at 24-hpi Bcpme1/2 samples formed an independent group (SI Appendix, Fig. S14B). We scored the results for log₂ fold-change compared to the WT strain (Fig. 5). After 16 h, Bcpg1, Bcpme1/2, and to a lesser extent Bcpg2 were affected in OG production while no differences were observed for Bcpg3 to -6. Interestingly, Bcpme1/2 was the only mutant affected in GalA₂ production and showed higher levels of methylesterified OGs of DP6 and -7. After 24 h, *Bcpme1/2* accumulated, relative to the WT strain, even more methylesterified OGs of DP5 to -7 and less GalA₃ and GalA₄Me. In addition, all of the mutants, except Bcpg1, accumulated less GalA₄Me than the WT strain, suggesting the involvement of all these enzymes in the production of this OG (Fig. 5). Interestingly, the large difference of the OGs released by Bcpg1 relative to that released by the WT at 16 h had completely disappeared at 24 h. This indicates that BcPG1 was critical for the rate of OG accumulation, not for OG accumulation per se.

Bcpg1 and Bcpme1/2 Are Affected in OG Accumulation during Infection of A. thaliana. We next investigated whether pectinolytic B. cinerea mutants were affected in OG accumulation also during infection of A. thaliana leaves. We focused on Bcpg1 and Bcpme1/2 since these mutants showed, in addition to compromised degradation of citrus pectins, the most perturbed OG profiles. A. thaliana leaves were infected with mutant and WT B. cinerea strains in liquid culture and the in vivo OGs were analyzed. HP-SEC-MS/MS analysis confirmed that OG production was affected in the 2 B. cinerea mutants also during infection of A. thaliana leaves (Fig. 6A). Indeed, for the WT strain, the relative OG composition evolved from 12 to 18 hpi. At 12 hpi, the OG size profile showed 2 peaks corresponding to large (>7) and small (3 to 5) DPs, respectively. The amounts of large DP OGs did not change during the time course. The amount of small OGs, however, strongly increased between 12 and 15 hpi and leveled off at 18 hpi. At this stage, a new peak corresponding to DP2 OGs (GalA₂ox) had appeared (Fig. 6A). In the mutants, the OG profile was strongly altered. First, whereas, like for the WT strain, the total amount of OGs increased over time, the relative OG composition did not evolve, suggesting that the enzymes encoded by the mutated genes were required for a specific step in the conversion of the OGs. For Bcpme1/2, more high-DP OGs accumulated relative to the WT strain, as observed on citrus pectins. In addition, DP4 levels were strongly reduced, DP5 to -10 accumulated and no DP2 accumulated at 18 hpi. BcPME activity therefore appears to be critical for the turnover of the large OGs, the production of DP4 OGs, and the conversion of OG into GalA20x. For Bcpg1, hardly any high DP OGs were

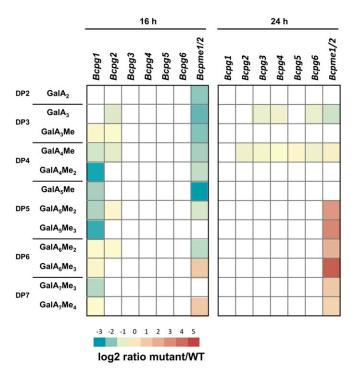


Fig. 5. Bcpme1/2 and Bcpg1 to Bcpg6 strains are defective in citrus pectin digestion and Bcpg1, Bcpg2, and Bcpme1/2 are impaired in OG production. OG production of Bcpme1/2 and Bcpg1 to Bcpg6 mutants compared to WT strain after 16 and 24 h of citrus pectin digestion. OGs are named GalA_xMe_{v.} Subscript numbers indicate the DP and the number of methylester groups, respectively.

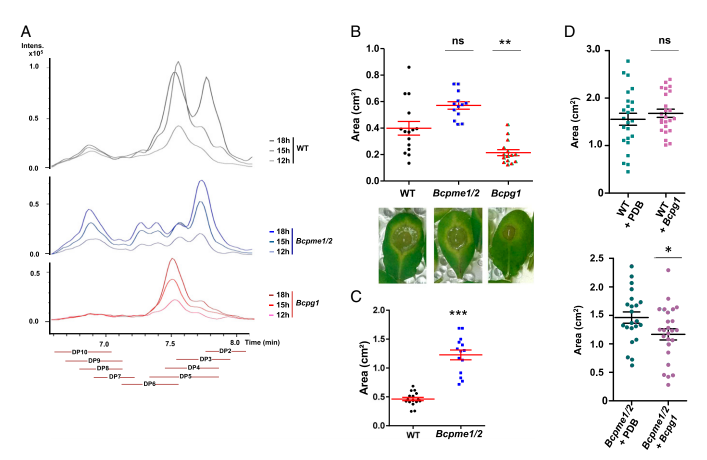


Fig. 6. Opposing effects of Bcpg1 and Bcpme1/2 mutations in virulence. (A) Extracted ion chromatograms obtained by HP-SEC-MS analysis in negative ionization mode of OGs endogenously produced during infection of A. thaliana leaves infected by WT B. cinerea strain and Bcpg1, Bcpme1/2 mutants over time. (B and C) Infection assays of WT B. cinerea and mutant strains on 5-wk-old (B) or 6-wk-old (B) rosette leaves of A. thaliana. Statistical data of lesion size 72 hpi. Values are means \pm SEM (B = 15); **P < 0.01 and ***P < 0.001 by Student test. (D) B. cinerea coinfection assay on rosette leaves of A. thaliana. Leaves were infected with WT B. cinerea strain or Bcpme1/2 followed 24 h later by Bcpg1 or a mock control (PDB medium). Lesion size of the infected leaves were measured 72 h later. Values are means \pm SEM (B = 25); *B < 0.05 by Student test; ns: nonsignificant.

detected, again as observed on citrus pectins, and most OGs were of DP4, with very minor amounts of DP2 or DP3. BcPG1 activity therefore contributes to the generation of the large DP OGs but also to the turnover of DP4 OGs.

Opposing Effects of *Bcpg1* **and** *Bcpme1/2* **Mutations on Virulence.** *B. cinerea* infection assays showed low virulence for *Bcpg1* (24) and a comparable (after 72 hpi on young leaves) or a higher virulence (on older leaves) for *Bcpme1/2* (23), relative to the WT (Fig. 6 *B* and *C* and *SI Appendix*, Fig. S15).

Since both mutants are compromised in pectin turnover, the results indicate that the virulence of the fungus does not only reflect its capacity to degrade pectin (which, on first sight, might explain the reduced virulence of *Bcpg1*) but also its ability to prevent elicitation of the defense system of the plant. To test the latter possibility, we infected leaves with *Bcpme1/2* followed, 24 h later, by *Bcpg1* or a mock treatment (Fig. 6D). Interestingly, the presence of *Bcpg1* reduced the virulence of *Bcpme1/2*, relative to the mock control. No such effect was observed when *Bcpme1/2* was replaced by the WT strain in the experiment. These results led us to hypothesize that the increased virulence of *Bcpme1/2*, despite its impaired pectinolytic capacity, reflects opposing activities of the OGs in leaves infected with *Bcpme1/2* or *Bcpg1/WT* respectively.

OG Analysis Released by *B. cinerea* Mutants Identifies New Defense Elicitor and Suppressor Candidates. To identify differentially accumulating OGs that might explain such differences in virulence, we

compared the OGs of A. thaliana leaves infected for 20 h (20 hpi) in liquid culture with Bcpme1/2 or WT strains. PCA results revealed clustering of replicate samples and a clear separation between data from treatment by Bcpme1/2 and WT strains, despite the higher variability among replicates in Bcpme1/2- relative to WT-infected conditions (SI Appendix, Fig. S16A). PCA on 72 OGs separated GalA20x and GalA4MeAc-H2O from the other OGs according to their relative abundance in the different samples (SI Appendix, Fig. S16B). Hierarchical cluster analysis on these OGs, after exclusion of GalA₂ox from the comparison because of its high intragenotype variability (Fig. 7A and SI Appendix, Fig. S17), showed that GalA₅Me₂Ac-H₂O/GalA₆Me₄-H₂O and GalA₄MeAc-H₂O/GalA₃Me were, respectively, under- and overrepresented in leaves infected with the WT strain, as compared to Bcpme1/2. This suggests that demethylesterification by BcPME1/2 might be a critical step in the conversion of methylesterified DP5 and -6 OGs into shorter OG, presumably also involving an exo-PG activity. To investigate potential correlations between the enrichment in some OG and virulence, we first quantified the methylesterified DP4, -5, and -6 OGs in Bcpg1-, WT-, and Bcpme1/2-infected leaves (Fig. 7B). Interestingly, low methylesterified DP4 OG accumulated more in leaves infected with Bcpg1 relative to the WT strain at 18 hpi, despite the reduced overall OG content in this mutant, and was barely detectable in Bcpme1/2. Opposingly, high methylesterified DP6 OG accumulated more in leaves infected with Bcpme1/2 relative to the WT and Bcpg1 strains.

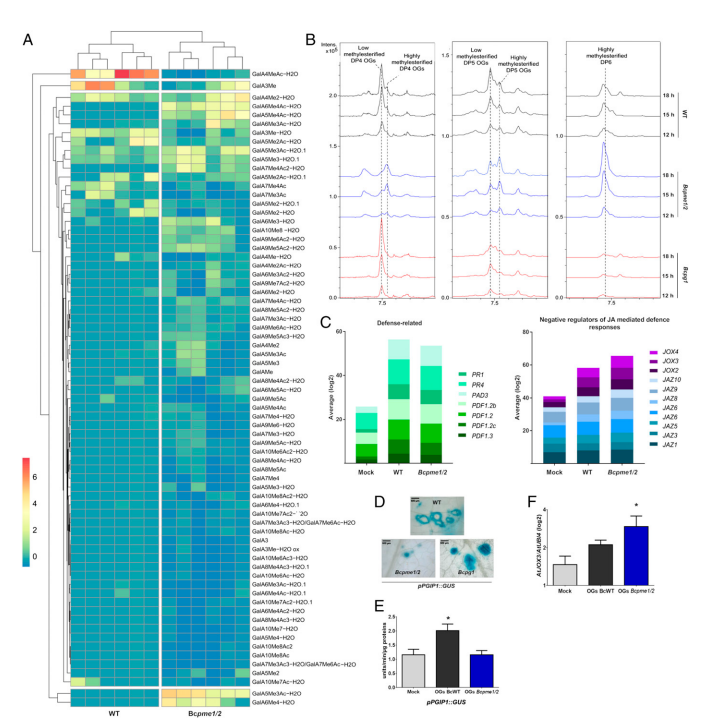


Fig. 7. The DP4/DP5 ratio of OGs produced by *Bcpg1*, *Bcpme1/2*, and WT *B. cinerea* strains correlates with their respective virulence. (*A*) Heatmap of all of the OGs (71 rows) for 12 samples (columns). Annotation labels refer to OG structure. Rows are centered; unit variance scaling is applied to rows. Both rows and columns are clustered using McQuitty distance and maximum linkage. (*B*) Extracted ion chromatograms for DP4, -5, and -6 OGs produced by *Bcpg1*, *Bcpme1/2*, and WT strains over time during infection of *A. thaliana* leaves. (*C*) Quantification of transcripts from JA and defense-related genes in noninfected leaves (mock) and leaves infected by WT and *Bcpme1/2* strains. Values of normalized transcript quantities from genes differentially expressed by WT or *Bcpme1/2* strains compared to noninfected leaves (log₂, false-discovery rate F-test, *P* value < 0.01) are shown in Dataset S1. (*D*) GUS staining of *A. thaliana* leaves expressing the promoter *PGIP1::GUS* fusion after the infection of *Bcpme1/2*, *Bcpg1*, and WT strains. (*E*) GUS activity quantified by fluorescence in leaves from transgenic *A. thaliana* plants expressing the *GUS* reporter gene under the control of *AtPGIP1* promoter. Leaves were infiltrated by semipurified OGs from leaves either infected by *B. cinerea* WT strain or *Bcpme1/2*. Values are means ± SEM (*n* = 6); **P* < 0.05 by Mann–Whitney *U* test. OGs are named GalA_wMe_yAc₂. Subscript numbers indicate the DP and the number of methyl- and acetylester groups, respectively.

The negative correlation between short OG levels and fungal virulence might indicate an elicitor specific activity for these OGs as opposed to longer-than-DP4 OGs that might have a defense-suppressor activity. The timing of appearance of these different

OGs may also be of importance for the virulence of the different strains. To clarify the role of these different OGs, we compared the transcriptomes of leaves infected for 24 h with WT or *Bcpme1/2*, when symptoms were already visible. Comparison of fungus-(WT or

Bcpme1/2) with mock-infected leaves revealed a total of 7,395 differentially expressed A. thaliana genes (Dataset S1), many of which were annotated as "defense related," involved in "protein processing in endoplasmic reticulum" or in "phenylpropanoid biosynthesis" (Fig. 7C and SI Appendix, Fig. S18). Among those, the pool of transcripts encoding negative regulators of JA signaling [JOX proteins hydroxylate and inactivate JA (30, 31) and JAZ proteins are transcriptional repressors of JA-regulated genes] (32) was slightly upregulated in leaves infected with Bcpme1/2 relative to the WT strain (Fig. 7C), suggesting that the Bcpme1/2 strain might suppress the plant defense mechanism. This did not simply reflect a more advanced infection stage of Bcpme1/2, since transcript levels for other defense-related genes were comparable, if not slightly lower, in Bcpme1/2 vs. WT-infected leaves. As mirrored in the enhanced and reduced induction of a defense reporter gene (pPGIP1::GUS) in response to infection with Bcpg1 and Bcpme1/2, respectively, relative to the WT strain (Fig. 7D), this suggests that, among the immune responses elicited by the fungus, JA-dependent defenses are induced more efficiently by the WT strain than by Bcpme1/2.

Finally, to test if low methylesterified DP4 OGs contribute to the plant defense elicitation and if highly methylesterified longer-than DP4 OGs contribute to the circumvention of plant defense mechanisms, we partially purified a mix of methylesterified OGs produced by BcWT and Bcpme1/2 (see SI Appendix, Fig. S19 for OG composition). First, we infiltrated leaves of transgenic A. thaliana plants transformed with the GUS reporter gene under the control of the AtPGIP1 promoter that is induced by trimers and long OG (DP10 to -15). Our results showed that at 24 h postinfiltration, AtPGIP1 is twice more activated by GalA₄MeAc-H₂Oenriched extract than with OGs produced by Bcpme1/2 (Fig. 7E). Second, we infiltrated A. thaliana leaves with these different OG mixes and monitored the expression of the JOX3 gene, which is involved in JA degradation. JOX3 was activated 3 h after the infiltration of OG purified from Bcpme1/2-infected leaves, enriched in highly methylesterified DP5 and -6 OGs, suggesting that these OGs may contribute to circumvent plant defense (Fig. 7F).

In conclusion, OG analysis revealed the presence of complex mixtures of OGs that are different from the previously described ones, and the nature and accumulation may be critical for the effectiveness of the immune response against *B. cinerea*, at least in part by affecting defense gene expression patterns *in planta*.

Discussion

This study showed that HP-SEC provides a high-resolution separation method, which is based on the DP, degree of acetylation, DM, and even the methylesterification pattern of the OGs and superior to commonly used HILIC-based methods (19–21). The OGs obtained with this method allowed the study of the mode of action of specific pectin-modifying enzymes, in vivo pectinolytic activity, OGs produced during plant–microbe interactions, and the identification of new cell wall-derived DAMP candidates.

The complexity of the OGs released upon in vivo plant-fungus interactions appears to reflect, to a large extent, the battleground of fungal pectinolytic activities and plant-defense strategies. For example, it was striking to see that unmethylesterified OGs of DP10 to -15 were absent in *B. cinerea*-infected citrus pectins or *A*. thaliana leaves while they have been detected in noninoculated leaves and in A. thaliana upon coexpression of a PGIP with a fungal PG (27, 33). They have, until now, been considered important DAMPs for necrotrophic pathogens and are frequently used in the study of defense responses (6–12). In B. cinerea grown on citrus pectins, the fungal enzymatic machinery rapidly converted PG products into GalA, GalA₂, and GalA₃, whereas higher DP OGs accumulated only in a methylesterified form. Similarly, germinated spore enzymes instantly convert long OGs (DP5 to -15) into shorter OGs. In B. cinerea-infected leaves, most PG products were converted into GalA₃Me and GalA₂ox. The production of the latter OGs presumably involved plant-produced OG oxidases

(OGOX) (17, 34, 35) of the berberine bridge enzyme family. Oxidized OGs are more recalcitrant to the hydrolysis of fungal enzymes and, interestingly, plants overexpressing OGOX1 are more resistant to B. cinerea. In our experimental settings, GalA₂ox accumulated at late infection stages (between 15 and 18 hpi) concomitantly with the reduction of GalA₃ and GalA₂ levels, which suggests that OG oxidase activity is induced only once the invader is detected. The exact role for GalA₂ox, however, remains to be demonstrated. Oxidation was shown to decrease the elicitor effect of OGs (17), in which case the enzyme activity may actually contribute to preventing OG-induced cell death during the necrotroph infection. Alternatively, the molecule itself or the H₂O₂ that is produced during the oxidation reaction may have signaling activity. Another strategy of the fungus to limit defense responses may be the suppression of unmethylesterified GalA₃ (another defense-response elicitor) (14) in B. cinerea-infected leaves relative to infected citrus pectins. This implies the existence, in B. cinerea-infected leaves, of an efficient PG activity able to hydrolyze such trimers, which might be BcPG3 (23).

This study showed that the overall OG profiles generated by *B. cinerea* grown on citrus pectins and on *A. thaliana* leaf cell wall were strikingly different. The former was exclusively the result of the hydrolytic activity of PGs, whereas the latter were for 80% the product of fungal PNL activity. The greatly reduced PG activity in the presence of *A. thaliana* cell wall would be the result of highly methylesterified and acetylated pectins.

One interesting application of the OG analysis in this study was the screening for novel OG DAMPs. The comparison of OGs of leaves infected with the WT strain or the mutants *Bcpg1* and *Bcpme1/2*, which both showed reduced pectinolytic activity but decreased and increased virulence, respectively, showed a negative correlation between virulence and the presence and timing of appearance of short (DP3 to -4) low methylesterified OGs during infection. Semipurified OG fraction from infected leaves by the WT strain was able to induce defense-gene expression, indicating the presence in the mixture of OGs with DAMP activity, which are different from the previously reported unmethylesterified GalA₁₀₋₁₅ or GalA₃. Similarly, long and highly methyl- and acetylesterified OGs were able to induce the *JOX3* gene, which is involved in JA degradation and suggests that these OGs are potential defense suppressors.

The final demonstration that some OGs have specific activities, like the induction or the suppression of defense responses, will await the in vitro production and purification of these OGs. This will be greatly facilitated by the identification in this work of the fungal enzymes that contribute to the generation of this OG. Indeed BcPNL1 (and perhaps 1 or more of the other 4 B. cinerea PNLs) generates, from A. thaliana cell walls, methylesterified OGs > DP4 with an unsaturated, often acetylesterified, GalA residue at the nonreducing end. The absence of GalA₄MeAc-H₂O in Bcpme1/2 indicates the involvement of 1 or both PMEs in the demethylesterification of a precursor molecule. This may be GalA₅Me₃Ac-H₂O, which selectively accumulated during infection by this mutant. It is conceivable that the PMEs remove 2 methyl groups from this molecule while leaving 1 methyl at position 3. A PG activity could then remove 1 GalA residue from the reducing end to yield GalA₄MeAc-H₂O. Finally, the accumulation of GalA₄MeAc-H₂O in *Bcpg1* indicates that BcPG1 is required for the turnover of the molecule, in particular during early infection

In conclusion, this study provides an extension of the omics toolbox with a method for the global analysis of the OGs that are produced during plant–pathogen interactions. This method will allow the study of other cell wall-derived signaling molecules involved in development and in abiotic stress responses, to investigate the evolution of enzymatic interactions between plants and their microbial environment, and may lead to the development of new strategies for the protection of crops against pathogens (36).

Materials and Methods

For full and detailed methods, please see SI Appendix, Materials and Methods.

Plant Material and Fungal Strains. A. thaliana WT Wassilewskija (Ws-0) plants were grown in soil in a growth chamber at 22 °C, 70% humidity, under irradiance of 100 μmol·m⁻²·s⁻¹ with a photoperiod of 8-h light/16-h dark. Leaves or pectins were incubated with different strains of B. cinerea named B05.10 Bcpq1-6 and Bcpme1/2.

OG Characterization and Quantification. Samples were analyzed by using chromatographic separation on an ACQUITY UPLC Protein BEH SEC Column and MS-detection performed in negative mode.

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