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Differential contribution of the two major polygalacturonases from Penicillium digitatum

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Abstract

The fungus *Pencillium digitatum* is the causal agent of the citrus green mould, the major postharvest diseases of citrus fruit. Lesions on the surface of infected fruits first appear as soft areas around the inoculation site, due to maceration of fruit. The macerating activity has been associated with pectinases secreted by the fungus during infection. In order to evaluate the contribution to virulence and macerating activity of the two major polygalacturonases (PGs) secreted by *P. digitatum*, we have obtained and characterized mutants lacking either pg1 or pg2, the genes encoding PG1 and PG2, respectively. Disease incidence of deletants in either gene was not different from that of the parental strain or ectopic transformants. However, disease progressed more slowly in deletants, especially in those lacking the pg2 gene. The lesions originated by the $\Delta pg2$ deletants were not as soft and the pH was not as acid as those originated by either the wild type strain or the ectopic transformants. Total PG activity in the macerated tissue was also lower in fruits infected with the $\Delta pg2$ deletants. Interestingly, the macerated tissue of oranges infected with $\Delta pg2$ deletants showed around 50% reduction in galacturonic acid content with respect to lesions caused by any other strain.

1. Introduction

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Green mould rot, caused by Penicillium digitatum, is the most common postharvest disease affecting citrus fruit in Spain (Tuset, 1987) and all production areas characterized by low summer rainfall (Eckert and Eaks, 1989). This pathogen may invade the fruit during the preharvest period through injuries occurred in the field, or/and in the packinghouses during storage and shelf-life periods. P. digitatum is a specialist pathogen that under natural conditions infects citrus fruit uniquely, although previous works demonstrated that it can infect overripe apple tissues (Buron-Moles et al., 2012; Vilanova et al., 2012; Vilanova et al., 2014). The use of synthetic fungicides has been the standard procedure to control this pathogen (Harding, 1972). However, these chemical treatments have several disadvantages, such as the persistence of the residues on the treated fruit, increase of the pathogen-resistant strains, as well as health and environmental problems (Bus, 1992). New approaches for designing new and safer control strategies would benefit from the knowledge of the molecular mechanisms underlying the pathogenesis of *P. digitatum*. P. digitatum is a necrotrophic wound pathogen that requires pre-existing injured fruit peel to penetrate the plant tissue (Kavanagh and Wood, 1967). Necrotrophs kill host cells by means of toxic molecules, which can be either host-specific, as tentoxin, or nonhost-specific toxins, as AM toxin, and lytic enzymes. However, the ultimate purpose of a necrotroph is not to kill its host, but to discompose the plant tissue and utilize the host-derived nutrients for its own growth (Zhang and van Kan, 2013). During infection, necrotrophic plant pathogens macerate the host tissue by secreting significant amounts of carbohydrate-active enzymes (CAZYmes) that contribute to the degradation of plant cell wall polymers to obtain the nutrients required for its development (Zhao et al., 2013). Among these CAZYmes special attention has been paid to those involved in pectin degradation. Pectin is the collective name for a complex of polysaccharides that constitute the

major carbohydrate type in the middle lamella (Jayani et al., 2005, Caffall and Mohnen, 2009). The most abundant type of pectin is homogalacturonan, a linear polymer of α-1,4-linked Dgalacturonic acid, which can be modified by acetylation and methyl-esterification. Other pectins include rhamnogalacturonan I and II, and xylogalacturonan. Enzymes involved in the degradation of the pectin backbone include polygalacturonases (PGs), pectate and pectin lyases (PLs), rhamnogalacturonases and rhamnogalacturonase lyases (recently reviewed by Ramoni and Seiboth (2016)). Depending upon the pattern of action (random or terminal) polygalacturonases (PGs) are termed as endo- or exo-enzymes, respectively. Endo- PGs are widely distributed among fungi, bacteria and many types of yeast whereas, in contrast, exo-PGs occur less frequently (Jayani et al., 2005). PGs play a critical role in pectin degradation by fungal pathogens and they hydrolyse the polygalacturonic acid chain across the oxygen bridge (Jayani et al., 2005). PG activity has been detected in decayed tissue and has been implicated as a virulence factor in several soft rot diseases (Reignault et al., 2008). In some pathogens, the disruption of PG genes reduced virulence, which suggests that this enzyme is a significant virulence factor in several plant-infecting fungi (Scott-Craig et al., 1990; Shieh et al., 1997). However, in several other cases, disruption of cell walldegrading enzymes caused only partial or no reduction in pathogenicity, suggesting that not all enzymes produced by the pathogen are required for pathogenicity (Scott-Craig et al., 1990). However, few studies on P. digitatum's cell walls degrading enzymes (CWDEs) encoding genes as virulence factors have been conducted so far (López-Pérez et al., 2015; Zhang et al., 2013a; Zhang et al., 2013b). Zhang and collaborators (2013b) have shown that a P. digitatum mutant lacking the polygalacturonase PG2 was able to infect citrus fruits, although it was less virulent than

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the parental strain. A similar phenotype has been described for *P. digitatum* mutants lacking the pectin lyase PL1, which showed reduced virulence (López-Pérez et al., 2015).

In a recent work, the importance of CWDEs in the virulence of *P. digitatum* was highlighted because they constituted the second most abundant group of genes in a library containing upregulated fungal genes during the infection of oranges (López-Pérez et al., 2015). The genome of *P. digitatum* is enriched in two families involved in pectin degradation, when compared to *P. chrysogenum*, a closely related but not pathogenic species. Thus, *P. digitatum* possesses eight polygalacturonases and rhamnnogalacturonases belonging to family GH28 and three pectin methylesterases belonging to family CE8 (Marcet-Houben et al., 2012).

In order to clarify the role of P. digitatum's PGs in pathogenicity, different strategies including physiological, biochemical and molecular investigations should be performed. In this work, we have compared the role in virulence of the two major PGs in P. digitatum by obtaining and characterizing in the same genetic background knockout mutants for the genes pg1 and pg2, to provide evidence that these genes play a different role during pathogenesis on orange fruit.

2. Materials and Methods

2.1. Fruits

'Valencia' orange fruits (*Citrus sinensis* L. Osbeck) were harvested from a commercial orchard in Tortosa (Catalonia, Spain) and processed the same day. Fruits were selected for uniform size, without physical injuries or apparent infections. Once the fruit arrived at the laboratory, they were surface-disinfected with a 10 % commercial bleach solution for 1 min, rinsed with tap water and allowed to dry at room temperature. Colour index, firmness, soluble solids and acidity were determined as quality parameters following standard procedures (Vilanova et al., 2013).

2.2. Fungal strains and culture conditions

Conidial suspensions from *Penicillium digitatum* Sacc. isolate Pd1 (CECT20795; (Marcet-Houben et al., 2012) were prepared by adding 5 mL of sterile water with 0.01 % (w/v) Tween-80 over the surface of seven- to 10-day-old cultures grown on potato dextrose agar medium (PDA; 200 mL boiled potato extract, 20 g dextrose, 20 g agar and 800 mL water) and rubbing the surface of the agar with a sterile glass rod. Conidia were counted in a haemocytometer and diluted to 10⁶ conidia mL⁻¹ inoculum concentration.

2.3. Construction of P. digitatum pg1 and pg2 disruption plasmids

P. digitatum Pd1 pg1 and pg2 genes correspond to NCBI gene entries PDIP_64460 and PDIP_19910, respectively. Further annotation of PDIP_64460 was required to match GenBank AB015286 sequence. Plasmids were constructed following the procedures described by López-Pérez et al. (2015). DNA fragments of 1.5-1.7 kb in length located upstream and downstream of both genes were amplified by PCR from P. digitatum Pd1 genomic DNA using primers pairs pg1-O1/pg1-O2 and pg2-O1/pg2-O2 (see Table 1 for primers sequences) for the upstream regions of pg1 and pg2, respectively, and pg1-A3/pg1-A4 and pg2-A3/pg2-A4 for the downstream regions. The amplified upstream, containing the first exon of the gene, and downstream regions from each gene were cloned flanking the hygromycin B resistance cassette in the vector pRF-HU2 (Frandsen et al., 2008), which was previously digested with Pac1 and Nt.BbvCI, following the USER friendly cloning technique (New Englands Biolabs, Beverly,MA, USA) to generate plasmids pRFDPG1 and pRFDPG2, respectively. An aliquot of each plasmid was used to transform E. coli DH5 quimio-competent cells. Kanamycin resistant colonies were screened for proper fusion of both upstream and downstream gene flanking fragments by PCR with primers RF1-RF6 and RF2-RF5

and verified by DNA sequencing. Selected plasmids were electroporated into *Agrobacterium* tumefaciens AGL-1 electrocompetent cells.

2.4. Agrobacterium tumefaciens-mediated transformation of Penicillium digitatum

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P. digitatum transformation was conducted as previously described (Marcet-Houben et al., 2012). Putative transformants were selected on PDA plates supplemented with hygromycin B (100 μg/ml) and cefotaxime (200 µg/ml). They were transferred to Eppendorf tubes containing PDA supplemented with hygromycin B (100 μg/ml) and cefotaxime (200 μg/ml) and incubated at 24 °C until sporulation. Conidia were inoculated into liquid GPY medium (glucose 40 g/l; peptone 5 g/l; yeast extract 5 g/l) supplemented with hygromycin (100 μg/ml) and incubated at 24 °C and 200 rpm for 2 days. DNA was extracted as described previously (López-Pérez et al., 2015). Insertion of the T-DNA in the transformants was verified by PCR with the primers HMBF1/HMBR1 (Table 1), which target the hygromycin B resistance gene. Integration by homologous recombination was analysed with primers pairs pg1-F7/pg1-R7 and pg2-F7/pg2-R7 for pg1 and pg2, respectively. The absence of the targeted gene in the deletants was further verified using the primers pg1-F8/pg1-R8 and pg2-F8/pg2-R8 (Table 1) for pg1 and pg2, respectively. Fig. 1A and 2A show a scheme with the relative position of the primers used in the characterization of the transformants. The number of T-DNA insertions present in selected monosporic transformants was determined by qPCR following the procedure described by Crespo-Sempere et al. (2013) using the primer pairs pg1-F9/pg1-R9 and pg2-F9/pg2-R9, which are located in the PCR-amplified upstream regions of pg1 and pg2, respectively. The P. digitatum gene encoding β-tubulin (GenBank accession number GU124566) was used as reference for normalization employing primers betatubPDIG1/betatubPDIG2 (Table 1). DNA from the wild-type Pd1 strain was used as a control.

PCR reactions were performed using a LightCycler 480 Real-Time apparatus (Roche, Manheim,

Germany) and the LightCycler 480 SYBR Green I Master kit (Roche) following the manufacturer's

recommendations.

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2.5. Fruit infections

Each orange was wounded by making two injuries with a nail (1 mm wide and 2 mm deep) in one side of fruit and was then inoculated with 15 μL aqueous conidia suspension of *P. digitatum* transformants or *P. digitatum* wild type strain. Oranges inoculated with sterile water were used as control treatment. After inoculation, oranges were stored at 20 °C and 85 % relative humidity (RH) for four days. Decay incidence and severity were measured. Four replicates per treatment were used and each replicate consisted of two inoculated wounds in five fruits.

2.6. pH

Mesocarp pH was determined by placing a micro-pH electrode directly into the wound (pH & Ion-Meter GLP 22 + Model 5033 pH electrode, Crison Instruments SA, Barcelona, Spain). Four replicates per treatment were used and each replicate consisted of two inoculated wounds in five

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2.7. Firmness measurements

Rot firmness was evaluated by measuring the maximal strength of compression of the infected lesion area using a TA-TX2 Texture Analyzer (Stable Micro Systems Ltd., Surrey, England). The resistance of the compression was measured using a cylinder probe with a round basis (P/0.75S) using the following conditions: pre-test speed (2 mm/s), test speed (0.3 mm/s), post-test speed (5.0 mm/s) and contact distance (2 mm) and results were expressed as Newtons (N). Two replicates per treatment were used and each replicate consisted of two inoculated wounds in five fruits.

2.8. Polygalacturonase activity assays

Two peel discs of 12 mm in diameter around the inoculation site and 4 mm deep were removed from 10 oranges using a cork borer. Twenty peel disks were so obtained, frozen immediately in liquid nitrogen, freeze-dried for 3 days, grounded to a fine powder in a coffee mill and stored at -80 °C until further analysis as described below. Twenty discs pooled from ten fruits were considered one replicate. Three replicates per treatment were used.

For the extraction of polygalacturonase (exo-PG; EC 3.2.1.67 and endo-PG; EC 3.2.1.15) activity, 100 mg of freeze-dried peel tissue from each replicate was homogenised (10 %, w/v) in extraction buffer prepared according to Lohani et al. (2004). PG activity was determined on the crude extracts as referenced in Ortiz et al. (2011). One unit (U) of PG activity was defined as the liberation of 1 µmol of GalUA min⁻¹ from citrus pectin (d.e. 70–75%), with galacturonic acid (GalUA) as a standard. Total protein content was determined with the Bradford (1976) method, with BSA as a standard. Results were given as specific activity over total protein (U mg-1 protein).

2.9. Analysis of organic acids

Two peel discs of 12 mm in diameter around the inoculation site and 4 mm deep were obtained in a similar manner to that used in the PG activity studies described above. Twenty discs from ten fruits were pooled and considered one replicate and four replicates per treatment were performed. Malic, ascorbic, oxalic, citric and fumaric acids were extracted and quantified using high performance liquid chromatography (HPLC), and gluconic and galacturonic acids using ultra-high-performance liquid chromatography—mass spectrometry (UHPLC—MS) system (Waters, Milford, USA) following the conditions described by Vilanova et al. (2014). Results were expressed as mg g-1 dry weight (DW).

2.10. Statistical analysis

Data regarding incidence and severity of fruit decay, quality parameters, pH, firmness, PG activity and organic acid levels were analysed for significant differences by analysis of variance (ANOVA) with the JMP 8 (SAS Institute Inc, NC, USA) statistical package. Statistical significance was defined as P<0.05; when the analysis was statistically significant, a Tukey test for separation of means was performed.

3. Results

3.1. P. digitatum pg1 and pg2 gene knockout mutants

We followed a gene deletion approach to study the role of the two major PGs from $P.\ digitatum$. The promoter and terminator regions of the genes pgI and g2 were PCR-amplified and cloned into the binary plasmid pRF-HU2 flanking the hygromycin B resistance cassette, originating plasmids pRFDPG1 and pRFDPG2, respectively. After $A.\ tumefaciens$ -mediated transformation of $P.\ digitatum$ Pd1, 50 and 96 transformants of pgI and pg2, respectively, were screened by PCR to detect the presence of the hygromcycin B resistance marker with primers HMBF1 and HMBR1 (Table1). All transformants were positive for the amplification of the expected 801 bp amplicon (Fig. 1B and 2B). We analysed the occurrence of double homologous recombination at the pgI and pg2 loci using primer pairs pg1-F7/pg1-R7 and pg2-F7/pg2-R7 for pgI and pg2, respectively. Eight pgI transformants and pg2 transformants showed a single amplicon of the expected size, 3.4 kb for pgI and pg2, respectively) in these transformants is indicative of gene replacement, thus originating pag2 and pag2 and pag2 null mutants. The remaining ectopic transformants showed two bands, the lower band corresponding to the original locus and the upper one corresponding to the T-DNA.

Further confirmation of gene replacement in $\Delta pg1$ and $\Delta pg2$ deletants was observed by the lack of amplification with gene-specific primers pg1-F8/pg1-R8 and pg2-F8/pg2-R8 for pg1 and pg2, respectively, (Fig. 1B and 2B). Amplification of a 700 or 650 bp band, for pg1 and pg2 respectively, with these primers was only observed in the wild type Pd1 strain and the ectopic transformants. The disrupted $\Delta pg1$ allele only contains a fragment of the pg1 gene encompassing the first 75 aa of the 367 aa of the PG1 protein, whereas the disrupted $\Delta pg1$ allele only contains the first 85 aa of the 378 aa. Hence, both disrupted Δpg mutants lack a functional gene.

Four deletant and two ectopic transformants for each gene were selected for determination of the number of T-DNA copies integrated in the genome by quantitative PCR using the wild type Pd1 strain as a control and the β -tubulin gene as the reference. All pg1 transformants contained only one copy of T-DNA. However, only two $\Delta pg2$ deletants and one ectopic transformant contained one copy of the T-DNA. The other two $\Delta pg2$ deletants contained two T-DNA copies and the second

and one ectopic transformant for each gene containing a single T-DNA integrated in the genome

were selected for further analysis. As shown in Fig. 1C and 2C growth and sporulation of selected

ectopic transformant contained three copies of the T-DNA integrated in the genome. Two deletants

deletant and ectopic transformants did not differ from the wild type strain.

3.2. Development of green mould caused by *P. digitatum pg* transformants

Fruit maturity stage was characterized by measuring colour index, firmness, soluble solids and acidity. Results for colour index and firmness on 'Valencia' oranges were 3.0 and 3.97 mm, respectively. Results for soluble solids and acidity were 11.0 % and 0.9 g L⁻¹ citric acid, respectively. In comparison to previous studies on 'Valencia' oranges (Vilanova et al., 2012,

Vilanova et al., 2013), our quality results indicated that oranges used in this study were at commercial maturity stage.

To evaluate the effect of both genes in virulence, oranges were inoculated with the wild type P. digitatum Pd1 and two Δpg mutants and one ectopic transformant for each gene. Our results showed that deletion of either pg1 or pg2 did not affect the decay incidence (Fig. 3A), but the lesion diameter in oranges inoculated with $\Delta pg1$ ($\Delta PG1$ -8 and $\Delta PG1$ -10) and $\Delta pg2$ ($\Delta PG2$ -0 and $\Delta PG2$ -13) was smaller than in those fruits inoculated the wild type strain after 4 days of inoculation (Fig. 3B). The average reduction in decay severity of the $\Delta PG1$ -8 and $\Delta PG1$ -10 was 31 and 33 %, respectively compared to the wild type strain, however no significant differences were found between its respective ectopic (EPG1-5) and the wild type strain. The average reduction in decay severity of the $\Delta PG2$ -0 and $\Delta PG2$ -13 was 47 and 51 %, respectively compared to the wild type strain. In addition, its respective ectopic mutant (EPG2-5) also showed lower lesion diameter than the wild type strain. In pathogenicity assays conducted with a lower inoculum dose (10⁴ conidia mL⁻¹ instead of 10⁶ conidia mL⁻¹) we observed the same results. Although there were no differences in disease incidence between the Δpg deletants and the wild type strain, decay severity was lower in fruits inoculated with the $\Delta pg2$ deletants (results not shown).

3.3. Changes in pH, firmness and polygalacturonase activity induced by *P. digitatum* mutants in orange decay

- The behaviour showed by the different mutants and the wild type strain in relation to pH, firmness and PG activity was evaluated at four days after inoculation (Fig. 4).
- *P. digitatum* wild type strain decreased orange peel pH from approximately 4.8 in control tissue (oranges inoculated with water) to approximately 3.0, representing around 38 % reduction (Fig.

4A). No significant differences were observed among the pH of the ΔpgI ectopic and null mutants (EPG1-5, Δ PG1-8 and Δ PG1-10) and the pH of the wild type strain. However, the pH value in fruits inoculated with the Δ PG2-0 and Δ PG2-13 deletants (3.2 and 3.1, respectively) was slightly higher than that measured in the wild type strain (3.0). Moreover, no significant differences were observed between the pH of EPG2-5 and the wild type strain.

Concerning firmness values, the *P. digitatum* wild type strain decreased the firmness of orange peel from approximately 6.32 N in control tissue to approximately 1.46 N, representing a reduction around 76 % (Fig. 4B). No significant differences in firmness were observed among the pgI ectopic and null mutants (EPG1-5, Δ PG1-8 and Δ PG1-10) and the wild type strain. However, firmness of

the \triangle PG2-0 and \triangle PG2-13 deletants (3.17 and 3.87 N, respectively) was markedly higher than that

measured in the wild type strain (1.46 N). No significant differences in firmness were observed

between the ectopic EPG2-5 and the wild type strain.

Large differences were found in PG activity levels between control tissue and the wild-type strain, activity values being approximately 20-fold higher in the latter (0.92 vs. 18.4 U mg⁻¹ protein, respectively) (Fig. 4C). No significant differences were observed among the PG activity of the pg1 ectopic (EPG1-5) and Δ PG1-8 null mutant and the wild type strain. However Δ PG1-10 showed lower PG activity (12.3 U mg⁻¹ protein) than the wild type strain (18.4 U mg⁻¹ protein). The lowest PG activity was found in Δ pg2 null mutants Δ PG2-0 and Δ PG2-13 (10.3 and 11.2 U mg⁻¹ protein, respectively). Activity levels in both Δ pg2 null mutants were significantly lower in comparison with their respective ectopic mutant (EPG2-5), which showed however similar levels in comparison with the wild-type strain.

3.4. Changes in organic acids induced by P. digitatum mutants in orange decay

Malic, ascorbic, oxalic, citric and fumaric acid level caused by the different mutants and the wild type strain was quantified after four days of inoculation. In general, no significant differences were observed among wild type strain and PG transformants (data not shown). Ascorbic acid level showed significant differences among control tissue (1.597 mg g⁻¹ DW) and that mutants with less lesion diameter: pg2 ectopic and null mutants also showed lower ascorbic acid levels (0.766, 0.866 and 0.516 mg g⁻¹ DW, respectively) than the wild type strain. Moreover, the lowest ascorbic acid level was detected in $\Delta PG2-13$ (0.516 mg g⁻¹ DW). Gluconic and galacturonic acid level caused by the different mutants and the wild type strain was quantified after four days of inoculation (Fig. 5) Gluconic acid level detected in control tissue was 18.666 mg g⁻¹ DW (Fig. 5A). However, no significant differences were observed among control tissue and pg1 and pg2 ectopic and null mutants and the wild type strain (in a range of 17.791 to 20.408 mg g⁻¹ DW). P. digitatum wild type strain increased galacturonic acid level from approximately 7.520 mg g⁻¹ DW in control tissue to approximately 108.087 mg g⁻¹ DW, being approximately 14-fold higher in the wild type (Fig. 5B). No significant differences were observed among the pg1 ectopic and null mutants (115.187, 118.718 and 110.794 mg g⁻¹ DW, respectively) and the wild type strain (108.087 mg g⁻¹ DW). However, the quantity observed in the \triangle PG2-0 and \triangle PG2-13 (55.645 and 54.727 mg g⁻¹ DW, respectively) was lower than that measured in the wild type strain (108.087 mg g⁻¹ DW) and the ectopic EPG2-5 mutant (135.541 mg g⁻¹ DW).

4. Discussion

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P. digitatum is a necrotrophic fungus that causes extensive maceration of the invaded citrus peel tissue, presumably due to the action of pectinases secreted by the pathogen during the infection process. A correlation between pectinases and orange peel maceration was observed in avirulent strains of P. digitatum obtained by UV irradiation (Garber et al., 1965). These mutants only caused necrotic lesions at the site of inoculation but did not cause rot. Pectinolytic activity was absent in the necrotic tissue but abundant in macerated tissue from diseased fruits. The analysis of P. digitatum pectinases obtained from culture filtrates showed that a secreted pectin lyase had macerating activity on orange rind tissue (Bush and Codoner, 1968). However, the macerating activity during lesion development was found to be associated with an exoPG purified from the macerate peel of oranges infected with *P. digitatum* (Barmore and Brown, 1979). We have previously identified in the genome of P. digitatum two PG encoding genes (pg1, PDIP 64460, and pg2, PDIP 19910, respectively) and two genes encoding pectin lyases (pnl1 and pnl2, PDIP 08080 and PDIP 57790) that showed a strong up-regulation during the infection of orange fruits (López-Pérez et al., 2015). P. digitatum mutants lacking either pg2 (Zhang et al., 2013b) or pnl1 (López-Pérez et al., 2015) showed reduced virulence compared to their wild type strains, although they were still able to develop infection on citrus fruit. In this work, we aimed to compare the role in virulence and maceration capability of the two major PGs, PG1 and PG2, from P. digitatum by using a functional genomics approach. To avoid strain specific effects, we used the same P. digitatum Pd1 strain for generating deletants of both pg1 and pg2. This is the same strain in which we have previously generated deletants lacking the pectin lyase PNL1 (López-Pérez et al., 2015). Moreover, the genome sequence of this strain is the species' reference genome at the NCBI's RefSeq database.

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Binary T-DNA plasmids containing the upstream and downstream regions of both genes flanking a hygromycin B resistance cassette were constructed and introduced into P. digitatum via Agrobacterium-mediated transformation. Two independent deletants and one ectopic transformant containing just one copy of T-DNA for each gene were selected. All of them grew and sporulated as the wild type strain on PDA medium (Figure 1 and 2). Pathogenicity assays showed that both $\Delta pg1$ and $\Delta pg2$ deletants had the same disease incidence on oranges as the wild type strain or the ectopic transformants. However, disease severity progressed more slowly in the Δpg deletants, specially in the two $\Delta pg2$ deletants, which showed an average reduction around 50% in lesion diameter with respect to the wild type strain. A similar reduction in disease severity was already observed in a $\Delta pg2$ deletant obtained in a different P. digitatum strain (Zhang et al., 2013b). Previous works conducted in Botrytis cinerea-infected tomatoes by Kars et al. (2005) showed that mutants in the *Bcpg2* gene had a >50 % reduction in virulence, meanwhile strains with a mutation in the Bcpg1 gene only reduced its virulence by 25 % (ten Have et al., 1998). The role of both P. digitatum PGs to disassemble the orange cell wall seems to be critical for the full virulence of P. digitatum; however, the disruption of one pectinase gene appears not to be enough to fully reduce the virulence of this pathogen, probably due to the presence of multiple pectinases in the genome of P. digitatum (Marcet-Houben et al., 2012). The presence and abundance of pectins into the wall matrix is considered to regulate the wall extensibility, and different pectin domains crosslink to each other via calcium and boron bonds. Additional cell wall-related enzymes such as pectin methylesterases can modify these connections, and therefore increase the susceptibility of pectins to depolymerisation by PGs and pectate lyases within the wall (Caffall and Mohnen, 2009). Furthermore, the accessibility of these enzymes to their pectin-backbone substrate is modulated by

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the activity of cell wall hydrolases acting on galactosyl- and arabinosyl-rich pectin side-chains, which thus regulate cell wall porosity (Goulao and Oliveira, 2008).

We noticed that the lesions in fruits inoculated with both $\Delta pg2$ deletants were not as soft as those observed in fruits inoculated with the wild type strain or any other transformants. Firmness values of $\Delta pg2$ deletants were significant higher than those obtained with the wild type and the other transformants (Fig 3B). This prompted us to analyse in more detail the characteristics of the macerated tissue in the lesions originated by the different strains. PG activity in the $\Delta pg2$ deletants showed a tendency to decrease in relation to wild type and the other transformants and this was clearly related with higher decayed tissue firmness and lower galacturonic acid production. The accumulation of galacturonic acid is a consequence of the complete pectin degradation by PG activity as reported Barmore and Brown (1979). In the case of *B. cinerea*, up to 13 endoPG isozymes have been described during the progress of the infection (van der Cruyssen et al., 1994). All BcPGs resulted true endopolygalacturonases, however, they showed different modes of action. PGA hydrolysis by PG1, PG2 and PG4 produced an accumulation of oligomers with DP < 7. However, PGA hydrolysis by PG3 and PG6 produced an accumulation of monomers and dimmers (Kars et al. 2005).

It is known that the timing and degree of *in planta* gene expression of the endoPG family differs depending on the host tissue, the degree of fruit ripening, the infection stage and the temperature. Besides pathogen PG activities, the peel also contains other cell wall degrading enzymes that contribute to softening of the tissue. Wubben et al. (2000) attributed the different expression patterns to four mechanisms: basal expression, induction by pectic monomers, repression of the glucose and ambient pH modulation. In our study, the deletion of *pg2* resulted in decayed tissue with higher firmness, lower PG activity and a 2-fold decrease in galacturonic acid level in relation

to the lesions originated by the wild type strain. These results demonstrated that $\Delta pg2$ strains had a reduced capacity for pectin decomposition and, hence, less amount of galacturonic acid was observed, and less nutrients were available for the fungus development. Restriction of growth may not be solely due to restriction of nutrients but also by physical limitation of hyphal growth through cells or more importantly between cells and air space. The monosaccharide D-galacturonic acid seems an important component for the nutrition of P. digitatum because the $\Delta pg2$ transformants showed less infection capacity, determined as the lesion diameter, which correlates with the production of lower amounts of galacturonic acid in the orange peel, indicating that the gene pg2 is a virulence factor. Taking into account that the albedo of citrus fruit is very rich in pectin, the capacity to degrade pectin by P. digitatum strain is critical to achieve a successful colonization of the host. Also in P. digitatum, Zhang et al. (2013b) found that one endoPG gene (Pdpg2) and one pectin lyase gene (*Pdpnl1*) were upregulated during citrus fruit infection in the wild type while not in the P. digitatum $\Delta pacC$ mutant. The PacC transcription factor is the terminal component of the pH signalling pathway. These authors hypothesize that PacC regulates the expression of some genes that are required for the degradation of pectin in the citrus peel, such as polygalacturonases and pectin lyases. The significant pH decrease observed in orange tissue infected by the different P. digitatum strains

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The significant pH decrease observed in orange tissue infected by the different P. digitatum strains agrees with previous reports by other authors (Prusky et al., 2004; Zhang et al., 2013b; Vilanova et al., 2014). From our results, the optima pH required by both PG from P. digitatum was at least between 3.0-3.2. Maceration tissue was clearly correlated with a decrease in pH, independently of the lesion diameter. No significant differences among wild type and both $\Delta pg1$ were observed, but there was a significant difference with the pH of the tissue infected by both $\Delta pg2$ deletants. The results obtained in this work confirm previous work that showed that galacturonic acid was not

responsible for the pH decrease observed in infected orange tissue (Vilanova et al., 2014); however, the difference in pH level between $\Delta pg1$ and $\Delta pg2$ (around 0.2 pH units) could be related to galacturonic acid content.

Different approaches to evaluate the role of CWDEs in pathogenesis have been conducted in other pathogens such as *Alternaria citris* (Akimutsi et al., 2004). They showed that the PG is essential for degradation of the plant cell wall components, mainly pectin, and for citrus fruit colonization and pathogenesis.

In conclusion, by obtaining gene knockout mutants, we have shown that the two major PGs, PG1 and PG2, produced by *P. digitatum* during infection of citrus fruit are required for full virulence. These two proteins seem to be dispensable to establish infection but they play a role in the colonization of the orange peel. Moreover, although these two proteins are endopolygalacturonases they show distinct enzymatic properties *in vivo*. Hence, PG2 plays a major role than PG1 in tissue softening, pH reduction and galacturonic acid production.

Acknowledgements

The technical assistance of Ana Izquierdo is gratefully acknowledged. LG-C's research is funded in part by the Spanish Ministry of Economy, Industry and Competitivness (AGL2011-30519-C03-01 and AGL2014-55802-R) and the Generalitat Valenciana (PROMETEOII/2014/027). ML-P was supported by a "Formación de Personal Investigador" scholarship (BES-2006-12983). Authors want to thanks the technical assistance of S. Dashevskaya and the financial support by AGL2011-30519-CO3-03 from the "Ministerio de Economía y Competitividad" (MINECO, Spain), and the CERCA Programme/Generalitat de Catalunya.

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Figure legends

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- Fig. 1. Analysis of *Penicillium digitatum pg1* transformants. (A) Diagram of the wild-type locus
- and the pg1 replacement with the Hyg^R selectable marker from pRFDPG1 by homologous
- recombination to generate the ΔpgI mutant. Primers used in the construction of plasmid pRFDPG1

and those used for the analysis of the transformants are shown. (B) Polymerase chain reaction (PCR) analysis of the wild-type Pd1 strain, an ectopic (Epg1-5) and two knockout (Δ pg1-8 and Δ pg1-10) transformants. (C) Growth of the wild type *P. digitatum* Pd1, an ectopic (Epg1-5) and two knockout Δ pg1 mutants (Δ pg1-8 and Δ pg1-10) after 7 days of incubation at 24 °C on PDA medium.

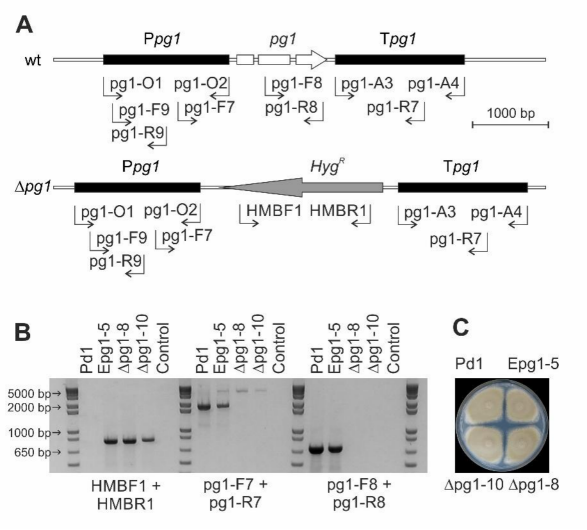
- **Fig. 2.** Analysis of *Penicillium digitatum pg2* transformants. (A) Diagram of the wild-type locus and the pg2 replacement with the Hyg^R selectable marker from pRFDPG2 by homologous recombination to generate the $\Delta pg2$ mutant. Primers used in the construction of plasmid pRFDPG2 and those used for the analysis of the transformants are shown. (B) Polymerase chain reaction (PCR) analysis of the wild-type Pd1 strain, an ectopic (Epg2-5) and two knockout ($\Delta pg2$ -0 and $\Delta pg2$ -13) transformants. (C) Growth of the wild type *P. digitatum* Pd1, an ectopic (Epg2-5) and two knockout $\Delta pg2$ mutants ($\Delta pg2$ -0 and $\Delta pg2$ -13) after 7 days of incubation at 24 °C on PDA medium.
- **Fig. 3.** Disease incidence (A) and lesion diameter (B) in 'Valencia' oranges inoculated with different *P. digitatum pg1* transformants at 10^6 conidia mL⁻¹ and stored at 20 °C and 85 % RH for 4 days. Wild type *P. digitatum* (Pd1), two ectopic mutants (EPG) and four knockout mutants (ΔPG) were analyzed. Each column represents the mean of four replicates and each replicate consisted of five fruits with two wounds per fruit. Samples with different letters are significantly different according to Tukey test (P<0.05).
- **Fig. 4.** pH (A), firmness (B) and polygalacturonase (PG) activity in 'Valencia' oranges inoculated with different *P. digitatum pg1* transformants at 10⁶ conidia mL⁻¹ and water as control treatment and stored at 20 °C and 85 % RH for 4 days. Wild type *P. digitatum* (Pd1), two ectopic mutants

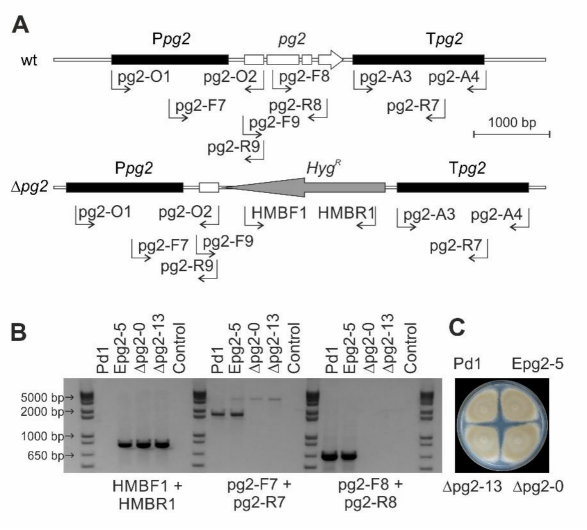
(EPG) and four knockout mutants (Δ PG) were analyzed. For pH, each column represented the mean of four replicates and for firmness, each column represented the mean of two replicates. In both cases, each replicate consisted of five fruits with two wounds per fruit. For polygalaturonase (PG) activity, each column represented the mean of three replicates and each replicate consisted of ten fruits with two wounds per fruit. Samples with different letters are significantly different according to Tukey test (P<0.05).

Fig. 5. Gluconic (A) and galacturonic (B) acid levels in 'Valencia' oranges inoculated with *P. digitatum pg1* transformants at 10^6 conidia mL⁻¹ and water as control treatment and stored at 20 °C and 85 % RH for 4 days. Wild type *P. digitatum* (Pd1), two ectopic mutants (EPG) and four knockout mutants (ΔPG) were analyzed. Each column represents the mean of four replicates and each replicate consisted of ten fruits with two wounds per fruit. Samples with different letters are significantly different according to Tukey test (P<0.05).

Table 1. Primers used in this study

Name	Sequence (5' to 3')	Purpose
pg1-O1	GGTCTTAAUGCCCCACTGGTCGATCCTAACCTTCCA	Amplification of the upstream region of <i>pg1</i>
pg1-O2	GGCATTAAUTGGGGGTTGACGCTTGCATAACAGAGC	Amplification of the upstream region of <i>pg1</i>
pg1-A3	GGACTTAAUGCCAGCGATCAAATGGTGAACACCAAAC	Amplification of the downstream region of pg1
pg1-A4	GGGTTTAAUAAGCGTCTGCGTGGTGGTGCAGT	Amplification of the downstream region of pg1
pg2-O1	GGTCTTAAUTGCGTGGTCTGTGGGGTGGTCGTTT	Amplification of the upstream region of pg2
pg2-O2	GGCATTAAUTGGGTGCCGGTGTTCAATCCAGTCA	Amplification of the upstream region of pg2
pg2-A3	GGACTTAAUTTTGACTCCTTGCTGGCCGGGCTTG	Amplification of the downstream region of pg2
pg2-A4	GGGTTTAAUTCCGCTCGTGAACAGGAGCACGTTG	Amplification of the downstream region of $pg2$
RF-1	AAATTTTGTGCTCACCGCCTGGAC	Analysis of plasmid constructs
RF-2	TCTCCTTGCATGCACCATTCCTTG	Analysis of plasmid constructs
RF-5	GTTTGCAGGGCCATAGAC	Analysis of plasmid constructs
RF-6	ACGCCAGGGTTTTCCCAGTC	Analysis of plasmid constructs
HMBF1	CTGTCGAGAAGTTTCTGATCG	Amplification of the hygromycin B resistance marker
HMBR1	CTGATAGAGTTGGTCAAGACC	Amplification of the hygromycin B resistance marker
pg1-F7	AAGCTCGATGGAATAGCTT	Detection of double homologous recombination at the pg1 locus
pg1-R7	CCCAGTAAAAGGACATGC	Detection of double homologous recombination at the pg1 locus
pg1-F8	AAAGAAGAAGCCCAAGTTCT	Detection of pg1
pg1-R8	AGCTACCGTTACCGCAGAGA	Detection of pg1
pg2-F7	ATGCTATTGGTTCTTTCCTC	Detection of double homologous recombination at the pg2 locus
pg2-R7	TCCCTCCGTAAACTAAACAA	Detection of double homologous recombination at the $pg2$ locus
pg2-F8	TCGATGGCGCTAAGGAGCTTACT	Detection of pg2
pg2-R8	CTCGGCACACAGAATGTA	Detection of pg2
pg1-F9	CGGACGGAGTAGATCTCACAACT	Determination of T-DNA copy number in <i>pg1</i> transformants
pg1-R9	CCTGCGCTAACATCCTCATGAAAC	Determination of T-DNA copy number in pg1 transformants
pg2-F9	CCTCGTGGTGCTTGTACCTTCTC	Determination of T-DNA copy number in $pg2$ transformants
pg2-R9	TCAGGGTAATGGTCGAGCAAGC	Determination of T-DNA copy number in $pg2$ transformants
betatubPDIG1	CGATGGCGATGGACAGTAAGTTT	Determination of T-DNA copy number in pg1 and pg2 transformants
betatubPDIG2	TTGGTTCGTGGTCGTTGTACTCA	Determination of T-DNA copy number in pg1 and pg2 transformants





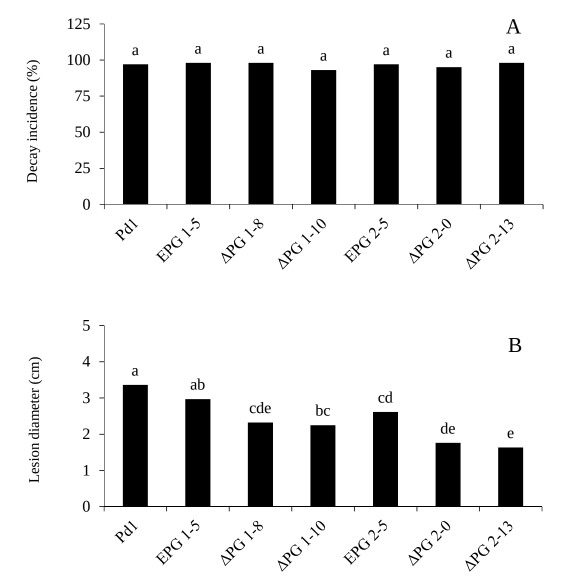
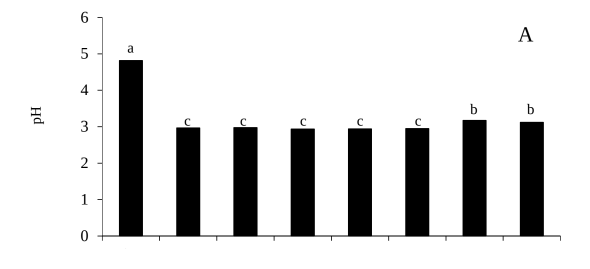
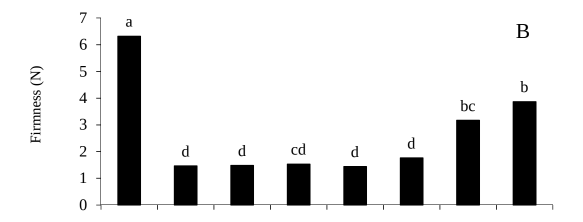


Fig. 3





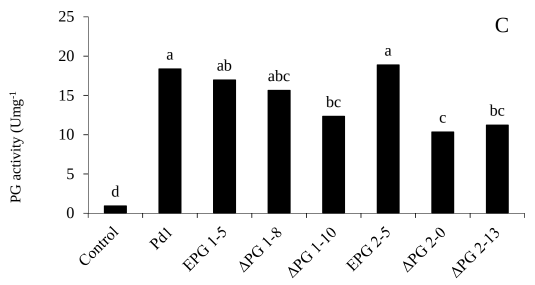


Fig 4

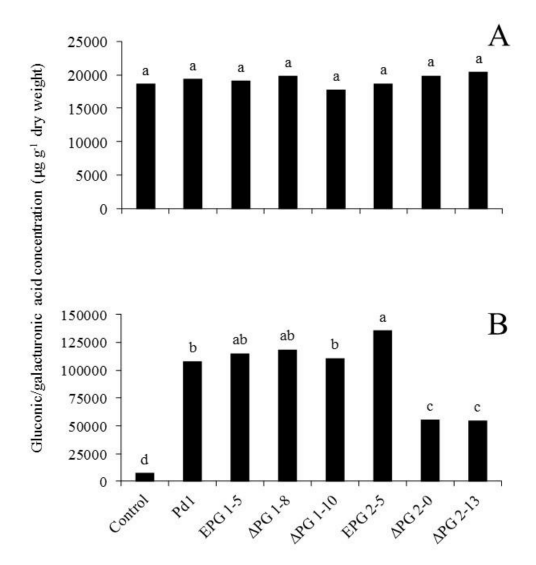


Fig. 5

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