Regulation of pectin methylesterase and polygalacturonate lyase activity during differentiation of infection structures in *Uromyces viciae-fabae*

Holger Deising, Alexander K. Frittrang, Stefan Kunz and Kurt Mendgen

The broad bean rust fungus *Uromyces viciae-fabae* differentiates infection structures up to the haustorial mother cell stage on thigmotropically inductive membranes in the absence of its host plant. Formation of pectin methylesterase (PME) and polygalacturonate lyase (PL), potentially involved in host cell wall degradation, was studied during infection structure differentiation by this biotrophic fungus. PME was first detectable when substomatal vesicles were formed and reached a maximum when infection hyphae and haustorial mother cells were differentiated. Four isoenzymes, exhibiting pl's of 8-2, 5-6, 5-2 and 4-5, were separated by chromatofocusing, and the kinetics of their synthesis and the $K_{s}$s of the three major isoenzymes were determined. The enzyme activity was formed independently of the presence of its substrate and its regulation was thus differentiation-specific. A single PL was induced when haustorial mother cells were formed and its synthesis appeared to be controlled by both the developmental stage of infection structures and the availability of its substrate. Polygalacturonate concentrations lower than 0-025 mg ml⁻¹ induced enzyme synthesis, and at 0-25 mg ml⁻¹ the induction process appeared to be saturated. Enzyme formation in the presence of 50 mM glucose, fructose or sucrose suggested that neither pectic enzyme was subject to catabolite repression. Significant proportions of PME (approx. 57%) and PL (approx. 76%) activity were located extracellularly in 24-h-old differentiated infection structures and could contribute to the establishment of the parasite. Physico-chemical and kinetic properties of the enzymes and associated alterations of the apoplastic pH of infected host plants appeared to be important factors in the success of infection and could explain the restriction of cell wall damage at the penetration site usually observed in interactions involving obligately biotrophic fungi.

**Keywords**: *Uromyces viciae-fabae*, cell-wall-degrading enzymes, differentiation specificity, infection structures, obligate biotrophy

INTRODUCTION

Cell-wall-degrading enzymes are thought to be of critical importance in the success of fungal plant pathogens (Cooper, 1984; Keon *et al.*, 1987). However, while in some host–pathogen interactions these enzymes are thought to be determinants of virulence (Köller *et al.*, 1982; Crawford & Kolattukudy, 1987; Cotty *et al.*, 1990; Wattad *et al.*, 1994), this issue has been disputed in other systems (Keen & Erwin, 1971; Howell, 1976; Cooper & Durrands, 1989; Scott-Craig *et al.*, 1990). Fungal pathogens differ with respect to host cell wall degradation and tissue maceration, and these differences are particularly evident when different groups of fungi, for example necrotrophs on the one hand and obligate biotrophs on the other, are compared (Cooper, 1984). Necrotrophs and saprophytes synthesize and secrete large quantities of cell-

*Abbreviations*: IWF, intercellular washing fluid; p.i., post-inoculation; PL, polygalacturonate lyase; PME, pectin methylesterase.
wall-degrading enzymes and have been extensively studied with respect to the regulation of these enzymes, with special emphasis given to pectic enzymes. In general, the presence of substrate results in enzyme induction. In contrast, low-molecular-mass carbon sources, such as mono- and disaccharides, function as repressors of enzyme induction (Crawford & Kolattukudy, 1987; Dean & Timberlake, 1989; Wattad et al., 1994; H. Deising, E. Rodriguez-Galvez & K. Mendgen, unpublished data). This type of regulation results in the formation of large quantities of enzymes in the host and extensive tissue maceration (Cooper, 1983).

In contrast, obligately biotrophic fungi such as rusts and powdery mildews cause minimal damage to the cell walls of their host plant, and this has raised the question of whether these fungi are capable of producing ‘true’ wall-degrading enzymes (see review by Mendgen & Deising, 1993). In uredospores and germings of *Uromyces viciae-fabae* which had differentiated appressoria, cellulases and ‘true’ pectic enzymes were not detectable (Cooper, 1984; Keon et al., 1987), and the authors speculated that this biotroph penetrates plant cell walls by means of glycanases and glycosidases which degrade neutral wall polymers.

However, if cell-wall-degrading enzymes were synthesized only after the fungus had invaded the leaf mesophyll, experiments performed with spores, germ tubes or appressoria would not allow their detection. Therefore, membranes providing a thigmotropic signal that causes differentiation of rust infection structures, including infection hyphae and haustorial mother cells (Deising et al., 1991), have been used to analyse enzyme formation in relation to morphogenesis. Using the same approach, Heier et al. (1993) showed that in *U. viciae-fabae* formation of cellulases begins when appressoria are differentiated. Frittrang et al. (1992) reported the partial purification and characterization of pectin methylesterases (PMEs) which, like cellulases of this fungus, are formed in a strictly differentiation-specific and substrate-independent manner.

In this paper we show that, in addition to PME, *U. viciae-fabae* produces a polygalacturonate lyase (PL). While the formation of both enzymes requires fungal morphogenesis, induction of PL occurs only in the presence of its substrate. Neither enzyme is subject to catabolite repression and significant quantities of both are secreted during infection structure differentiation. The kinetics of synthesis and physico-chemical properties of these pectic enzymes, and the associated changes in the apoplastic pH of infected plants are discussed with respect to highly localized wall degradation by rust fungi.

### METHODS

**Fungal culture and production of infection structures.** Uredospores of *U. viciae-fabae* isolate 12 were propagated on *Vicia faba* cv. Con Amore (Deising et al., 1991). Spores were either used directly after harvesting or stored at -70°C until needed.

Infection structure differentiation was induced by thigmotropic stimuli provided by scratched polyethylene membranes as described previously (Deising et al., 1991). Non-differentiated germlings were obtained 4 h post-inoculation (p.i.) from non-inducing substrates (membranes that had not been scratched). Each membrane (1870 cm²) was inoculated with 170 mg uredospores. Since PME activity is tightly controlled by fungal morphogenesis and not by the presence of substrate, membranes were sprayed with 4 ml of sterile distilled H₂O after inoculation. They were subsequently incubated at 19°C and 100% relative humidity. To induce PL, membranes were routinely sprayed with 4 ml sterile aqueous sodium polygalacturonate (1 mg ml⁻¹) (Sigma). To determine the dependence of enzyme induction on substrate concentration, sodium polygalacturonate concentrations ranging from 0 to 1 mg ml⁻¹ were applied. In order to analyse whether pectic enzymes of the broad bean rust fungus are catabolite-repressible, mono- and disaccharides were applied as 50 mM solutions either alone (PME) or in combination with 1 mg ml⁻¹ sodium polygalacturonate (PL). The pH of the solutions used for PL induction was adjusted to 5.4. To relate the pattern of expression of enzyme activities to different stages of fungal morphogenesis, structures were harvested at different times after inoculation of membranes. These time points were based on the kinetics of infection structure differentiation as published previously (Deising et al., 1991).

To demonstrate the fungal origin of PL, cycloheximide was applied at 0·1 or 10 µg ml⁻¹ to uredosporelings 8 h after inoculation and substrate application, and streptomycin sulfate or ampicillin (0·5 mg ml⁻¹ of each in combination with 0·1% polygalacturonate) (Melgarejo et al., 1985; Heier et al., 1993) were applied directly to inoculated membranes. Cycloheximide and ampicillin were obtained from Sigma, and streptomycin sulfate from Serva.

**Enzyme extraction and assay conditions.** PME was extracted and assayed as described previously (Frittrang et al., 1992), with minor modifications. Briefly, spores, germ tubes and infection structures were homogenized in 50 mM Tris/HCl buffer, pH 8·0, containing 150 mM NaCl. Homogenization of dormant uredospores was carried out using a pestle and mortar. A Potter-Elvehjem homogenizer (Kummer) was used to homogenize all other structures. Homogenization was carried out at 2·4°C. The homogenate was centrifuged (24000 g, 20 min, 2°C) and the clear supernatant was subjected to (NH₄)₂SO₄ precipitation (degree of saturation 0.8) and centrifuged as described above.

The pellet was resuspended in 25 mM Tris/HCl, pH 8·0, dialysed against 500 vols of the same buffer overnight, centrifuged (20000 g, 30 min, 2°C), and the supernatant stored at -70°C or used directly to determine PME activity.

The assay mixture, if not otherwise stated, consisted of 16 mg ml⁻¹ apple pectin type B (degree of esterification 72–79%) and 150 mM NaCl in 5 mM PIPES buffer, pH 7·0. For substrate specificity analyses, other apple pectins (type A1, degree of esterification 35–40%; type A2, degree of esterification ~65%) and pectin type N (degree of esterification ~8%) (all from Roth) were used at a final concentration of 3·2% (w/v) in the buffer described above. Enzyme activity was determined pH-metrically at 30°C (Frittrang et al., 1992).

For extraction of PL, homogenization was carried out in 50 mM Tris/HCl buffer, pH 8·5, containing 1 mM PMSE and 1% (v/v) methanol. The homogenate was adjusted to contain 500 mM NaCl, and after centrifugation (45000 g, 20 min) the supernatant was subjected to (NH₄)₂SO₄ precipitation (degree of saturation 0·9). After centrifugation (45000 g, 20 min), the pellet was dissolved in 10 mM Tris/HCl buffer, pH 9·5, and dialysed against 500 vols 1 mM Tris/HCl, pH 9·5, overnight. Protein precipitates were removed by centrifugation as described above.
and the supernatant was used for PL assay and protein determination. All operations were performed at 2-4 °C.

The PL assay mixture consisted of (final concentrations) 5 mg polygalacturonate ml⁻¹ and 1-41 mM CaCl₂ in 100 mM 2-amino-2-methyl-1,3-propanediol buffer, pH 10-3. The reaction was carried out at 30 °C, and the increase in A₅₆₅ was used as a measure of enzyme activity. Controls contained either equal amounts of heat-inactivated enzyme, or active enzyme plus EDTA at a final concentration of 5 mM.

**Column chromatography**

**Ion exchange chromatography.** To separate PME isoforms A and B, ion exchange chromatography was performed on a polybuffer exchanger (PBE) 94 (Pharmacia). The column (gel bed 9 ml, 18 cm long) was equilibrated with 25 mM Tris/HCl, pH 9-0. Crude extract (15 ml) containing a maximum of 470 mg protein was dialysed against equilibration buffer overnight and applied to the column. After collecting the unbound fraction containing PME isoform A and extensive washing with equilibration buffer (10 column vols), PME isoforms B and C were eluted with 1 M NaCl. The flow rate was 0-25 ml min⁻¹. Bound and unbound fractions were (NH₄)₂SO₄-precipitated and dialysed against the respective equilibration buffer used in chromatofocusing. All operations were carried out at 4 °C.

**Chromatofocusing**

Chromatofocusing was performed on a PBE94 column (Pharmacia) (gel bed 9 ml, 18 cm long). For chromatofocusing in the pH range 9-0-7-0, the column was equilibrated with 25 mM ethanolamine, pH 9-0. Unbound fractions from ion exchange chromatography columns (containing PME isoform A), dialysed against 100 vols equilibrium buffer, was applied to the column, and 5 ml equilibrium buffer was subsequently loaded. Proteins with PIS in the range 9-0-7-5 were eluted with 10% (v/v) polybuffer PB 96 (Pharmacia), adjusted to pH 7-0. The column was then washed with 1 M NaCl. For chromatofocusing in the pH range 7-0-4-5, the PBE94 column was equilibrated with 25 mM imidazole, pH 7-0. The bound fraction from the ion exchange chromatography column (containing PME isoforms B and C) was dialysed against the equilibration buffer and applied to the column as described above. Elution of proteins was performed using 10% (v/v) polybuffer PB74 (Pharmacia), pH 4-0. The flow rate was adjusted to 0-25 ml min⁻¹. Fractions of 2 ml were collected. Chromatofocusing was carried out at 4 °C.

The pH gradient was monitored using a type 405-S7 pH electrode (Ingold), and the A₅₆₅ of each fraction was determined. Prior to determination of PME activity, each fraction was dialysed overnight against 5 mM PIPES buffer, pH 7-0, containing 150 mM NaCl.

**Localization of PL.** To obtain extracellular proteins, polyethylene membranes with adhering 24-h-old differentiated infection structures were inverted and floated on 350 ml 50 mM piperazine/HCl buffer, pH 6-0 for 10 min. The washing fluid from 15 membranes was adjusted to contain 0-5 M NaCl, cleared by filtration (MN 615 filter paper, Macherey-Nagel), and concentrated to approximately 7 ml by ultrafiltration using an Amicon 8200 cell with an Amicon YM5 filter (62 mm) at 2-5 bar pressure.

The washed infection structures were homogenized in 50 mM piperazine/HCl buffer, pH 6-0, containing 0-1% (v/v) Triton X-100 and 0-5 M NaCl as described above. After centrifugation (45000 g, 20 min, 2 °C), the pellet was washed with the same buffer and centrifuged again. The supernatants, referred to as intracellular proteins, were combined and concentrated by ultrafiltration as described above.

After dialysis against 500 vols distilled H₂O overnight at 4 °C, extracellular washing fluid and intracellular proteins were centrifuged as described above and used for PL assay and protein determination. To estimate the fraction of damaged infection structures, the activity of the cytoplasmic marker enzyme NADH-malate dehydrogenase (EC 1.1.1.37) was measured. The optimized assay for fungal malate dehydrogenase consisted of 0-15 mM NADH and 2-5 mM oxaloacetate in 200 mM potassium phosphate buffer, pH 9-0. The assay was performed at 30 °C and the decrease in A₅₆₅ was used as a measure of enzyme activity.

**Changes in the apoplastic pH and electrical conductivity of V. faba leaves after rust infection.** Fourteen-day-old V. faba plants were inoculated by spraying with a uredospore suspension [ approx. 0-65 mg uredospores and 0-05 mg BSA (ml H₂O)⁻¹] and 8 d after inoculation heavily infected and uninfected control leaves were excised and washed twice in distilled H₂O at 0-2 °C. The leaves were infiltrated with distilled H₂O at 0-2 °C at ~ 800 mbar for 3 min, and the vacuum was slowly released over 3 min. The plant material was then blotted dry between paper towels and centrifuged at low speed (36 g, 20 min, 4 °C). The intercellular washing fluid (IWF) obtained was used to measure the apoplastic pH, electrical conductivity and extracellular malate dehydrogenase activity.

Vacuum-infiltration of the leaves was repeated with 50 mM NaCl followed by low speed centrifugation. The leaves were then ground under liquid nitrogen with a pestle and mortar, 1-5 ml 50 mM NaCl (g leaf fresh wt)⁻¹ added and the suspension stirred for 30 min at 4 °C. After centrifugation (45000 g, 20 min, 2 °C), the leaf extract was used to determine intracellular pH and malate dehydrogenase activity.

The assay mixture used to measure plant malate dehydrogenase activity contained 0-15 mM NADH and 2-5 mM oxaloacetate in 50 mM potassium phosphate buffer, pH 7-0. The assay was performed at 30 °C, and the decrease in A₅₆₅ was used as a measure of enzyme activity.

To determine the intracellular pH of rust infection structures, 24-h-old differentiated structures were homogenized in distilled H₂O as described above.

To measure the ion activity of the apoplast, 1-5 ml IWF was added to 13-5 ml distilled H₂O and the conductivity of this solution measured with a conductivity meter (WTW).

**Protein determination.** Protein concentration was determined in 50 mM NaOH using a commercially available protein assay kit (Bio-Rad) and a method based on that described by Bradford (1976). γ-Globulin served as the standard.

**RESULTS**

**Characterization of PME isoforms**

In fully differentiated 24-h-old infection structures of U. viciae-faba, pre-fractionation of crude extracts by ion exchange chromatography followed by chromatofocusing revealed the presence of four PME isoforms (Fig. 1). The PIs were 8-2 (isoform A) (Fig. 1a), 5-6 (B1) and 5-2 (B2) (Fig. 1b) and 4-5 (C). Use of PBE94 as the gel matrix and polybuffer 74 instead of a DEAE Si 500 column, and the
carrier ampholyte Servalyt 3-8 (Frittrang et al., 1992) allowed resolution of PME isoforms B (originally described by Frittrang et al., 1992) into two isoforms, designated B1 and B2. The activities of isoform B1 and B2 were each found to be 49% of the total recovered activity applied to the chromatofocusing column in the pH range 7.0-4.5. The remainder was due to isoform C which generally contributed approximately 2-5% of the activity applied to the column. Chromatofocusing was also used on a preparative scale to yield sufficient enzyme to determine substrate specificity and the $K_m$ values of PME isoforms A, B1 and B2. Isoform C, due to its low activity, was not further studied.

With pectins of different degrees of methylation, PME isoforms A, B1 and B2 exhibited significant differences in substrate specificity (Fig. 2). All isoforms showed very low activity with pectin N (degree of esterification 8%).

At a degree of esterification of 35-40%, isoforms B1 and B2 reached approximately 80% of their maximum activity while isoform A showed only about 10% maximum activity. At 65% methylation of the substrate, all isoforms exhibited maximum activity.

To determine the dependence of the activities of the PME isoforms on substrate concentration, increasing concentrations of pectin B (degree of esterification 72-79%) were used in the enzyme assay (Fig. 3). Isoforms B1 and B2 were both highly active in the presence of low pectin concentrations, whereas isoform A required high substrate concentrations for maximum activity. The $K_m$ values as determined by Eadie–Hofstee transformation of the kinetics data (Fig. 3, insert) showed significantly higher substrate affinity of isoforms B ($K_m$ = 2.32 ± 0.66 and 2.05 ± 0.77 mg ml$^{-1}$ for B1 and B2, respectively), as compared to isoform A ($K_m$ = 23.42 ± 10.1 mg ml$^{-1}$).

**Differentiation-specific PME formation**

The kinetics of the expression of PME were followed during differentiation of *U. viciae-fabae* infection structures for 24 h. PME activity was first detected at 8 h p.i. on inductive substrates, i.e. when substomatal vesicles were beginning to form (Fig. 4). From 10 to 12 h p.i. isoforms A, B1 and B2 increased dramatically. From 12 h p.i., isoforms B remained unaltered while isoform A increased until 20 h p.i. and remained at that level for the next 4 h. Isoform C contributed 2-5% of total activity. Isoforms B accounted for the majority of PME activity when the substomatal vesicles were differentiated (data not shown) but their relative activities decreased during further development. In contrast, from vesicle to haustorial mother cell differentiation, isoform A increased from approximately 25 to more than 70% of total activity, as measured in the presence of 10 mg pectin ml$^{-1}$ in the...
Regulation of pectic enzymes of *U. viciae-fabae*

![Graph](image1.png)

**Fig. 3.** Effect of substrate concentration on the activity of *U. viciae-fabae* PME isoforms and determination of *Km* values. Isoforms A (■), B1 (○) and B2 (▲) were separated by chromatofocusing and assayed in the presence of different concentrations of pectin type B (degree of esterification 72–79%). Eadie–Hofstee transformations were used to determine *Km* values of the individual isoforms (insert). Means of three independent experiments are shown.

![Graph](image2.png)

**Fig. 4.** Kinetics of formation of PME activity in differentiating uredosporelings of *U. viciae-fabae*. The activities of the separated PME isoforms A (■), B1+B2 (○) and C (▲) were determined using pectin type B (degree of esterification 72–79%). Arrows indicate the occurrence of fungal structures. gt, germ tube; app, appressorium; sv, substomatal vesicle; ih, infection hypha; hmc, haustorial mother cell. Representative results are shown.

![Graph](image3.png)

**Fig. 5.** Kinetics of formation of PL activity in differentiating uredosporelings of *U. viciae-fabae*. Sterile polygalacturonate solution (1 mg ml⁻¹) was applied 0 (■), 8 (○) or 18 (▲) h p.i.; distilled H₂O (0) was sprayed onto inoculated membranes 0 h p.i. as a control. Arrows indicate the occurrence of fungal structures (abbreviations as in Fig. 4). The experiment was done in triplicate and bars represent SE.

reaction mixture. Interestingly, PME was formed in the absence of potential inducers such as pectin or polygalacturonate acid since none of these substances had been added to differentiating infection structures.

**Characterization of PL induction**

When uredospores sprayed with sterile distilled H₂O differentiated to form infection structures, no PL activity was detected during the time frame covered by the experiment (Fig. 5). However, when a polygalacturonate solution was sprayed onto inductive surfaces inoculated with uredospores, PL activity became detectable 14 h p.i. and increased up to 24 h p.i. When polygalacturonate was applied 8 h p.i. to morphogenetic stages that, under natural conditions contact pectic substances of the plant mesophyll, the induction kinetics measured were almost identical. Polygalacturonate applied to 18-h-old differentiating structures resulted in significant PL activity after 6 h. It thus appears that the single PL isoform observed in *U. viciae-fabae* can be induced only in infection hyphae or haustorial mother cells.

The dependence of induction of PL activity on the polygalacturonate concentration applied to uredospores on inductive membranes is shown in Fig. 6. PL activity was induced by polygalacturonate concentrations lower than 0.025 mg ml⁻¹, and at approximately 0.125 mg ml⁻¹ the induction process appeared to be saturated with substrate.

Approximately 76% of the PL activity of the rust fungus
was found in the extracellular washing fluid (Table 1). The fact that only minor proportions of protein and the cytoplasmic enzyme malate dehydrogenase were detected extracellularly shows that the extracellular location of PL is not due to damage of infection structures.

**Effect of mono- or disaccharides on enzyme formation**

Different mono- or disaccharides occurring in the plant apoplast were not involved in the regulation of pectic enzymes of the broad bean rust fungus (Fig. 7). While glucose, fructose and sucrose did not significantly affect the development of PME activity, galacturonic acid monomers reduced activity by approximately 50% (Fig. 7a). The addition of 50 mM glucose, fructose or sucrose to the polygalacturonate solution needed to induce PL activity of *U. viciae-fabae* did not reduce the extractable enzyme activity by more than 50% (Fig. 7b).

**Characterization of the fungal environment**

Since PME and PL exhibit pH optima of 6.5–8.0 (PME; Frittrang *et al.*, 1992) and 10.3 (PL; H. Deising, unpublished data) it was of interest to determine alterations of apoplastic pH associated with rust infection. Table 2 shows that the pH in the apoplast of *U. viciae-fabae*-infected broad bean leaves is significantly higher than in control plants 8 d.p.i., and also that the intracellular pH of non-buffered homogenates of rust-infected leaves show pH values approximately 0.6 units above those of non-infected control plants. Homogenates of 24-h-old differentiated infection structures show pH values of 5.44 ± 0.03. The small proportion of malate dehydrogenase activity present in the apoplast [15.75 ± 0.612 and 10.68 ± 3.93 nkat (g fresh wt)^-1 in rust-infected and control leaves, as compared with 76.48 ± 7.61 and 53.90 ± 1.02 μkat (g fresh wt)^-1 in the respective homogenates] demonstrates the integrity of mesophyll cells. However, the electrical conductivity in IWFs of rust-infected leaves is significantly higher than that of healthy control samples.

**Exclusion of bacterial contamination**

Microscopic examination showed that no bacterial contaminants were present during the experiments. In addition, the protein biosynthesis inhibitor cycloheximide, which specifically blocks eukaryotic mRNA translation on 80S ribosomes, was added to solutions used to spray inoculated membranes. As shown in Fig. 8, PL induction was inhibited in a dose-dependent manner by cycloheximide. Furthermore, inhibitors such as streptomycin sulfate and ampicillin, which block prokaryotic and plastid protein synthesis, and bacterial cell wall synthesis, respectively, inhibited PL formation to only a minor extent.

**DISCUSSION**

Cell-wall-degrading enzymes are thought to be important factors in determining the outcome of interactions between plants and plant-pathogenic fungi (Mendgen & Deising, 1993). However, little is known about the regulation of these enzymes in obligately biotrophic fungi. In previous reports (Frittrang *et al.*, 1992; Heiler *et al.*, 1993) and in this paper we have demonstrated that the broad bean rust fungus forms cell-wall-degrading enzymes mainly during the time when infection hyphae and haustorial mother cells are differentiated. This

**Table 1. Localization of *U. viciae-fabae* PL and malate dehydrogenase activity, and of protein in 24-h-old differentiated infection structures**

<table>
<thead>
<tr>
<th>Location</th>
<th>PL (nkat per membrane)</th>
<th>Malate dehydrogenase (pkat per membrane)</th>
<th>Protein (mg per membrane)</th>
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<tbody>
<tr>
<td>Extracellular</td>
<td>203.3 ± 16.0</td>
<td>2.82 ± 0.02</td>
<td>0.145 ± 0.036</td>
</tr>
<tr>
<td>Intracellular</td>
<td>61.5 ± 16.7</td>
<td>20.47 ± 46.7</td>
<td>1.151 ± 0.199</td>
</tr>
<tr>
<td>Percentage extracellular</td>
<td>76.8</td>
<td>1.4</td>
<td>12.6</td>
</tr>
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</table>

![Graph](image.png)

**Fig. 6.** Effect of polygalacturonate concentration on PL formation in infection structures of *U. viciae-fabae*. Inductive membranes were inoculated with uredospores of the fungus and sprayed with solutions of different substrate concentrations. Infection structures were allowed to differentiate for 24 h. The experiment was carried out four times and bars represent SE.
require fungal morphogenesis but not the presence of substrate, and is not subject to catabolite repression. The acidic cellulases start to appear during appressorium development (Heiler et al., 1993), while the formation of PME isoforms B1 and B2 (pls 5·2 and 5·6) begins with differentiation of young infection hyphae, which are usually enlarged to substomatal vesicles (Mendgen & Deising, 1993; Fig. 4). Under the pH conditions of the broad bean leaf apoplast (approx. pH 4·5; Aloni et al., 1988), even these acidic enzymes would be bound to the plant cell wall. Comparison of the $K_m$s of the PMEs of $U$. viciae-fabae reveals that isoforms B1 and B2 have an approximately tenfold higher affinity for their substrate than isoform A (Fig. 3, insert). Similar $K_m$ values have been reported for PMEs from Botrytis cinerea (Marcus & Scheijter, 1983). Isoforms B1 and B2 also act on pectins of low degrees of methylation much more efficiently than isoform A (Fig. 2). Based on their $K_m$ values, the rust PME isoenzymes B1 and B2 could therefore efficiently de-esterify pectins with low degrees of methylation.

Immunocytochemical studies have shown that the polygalacturonate strands are not uniformly esterified across the cell wall. Whereas high degrees of methylation are found in cell wall layers facing the plasmalemma, low degrees of methylation have been demonstrated at the apoplastic side of the cell wall (Liners & Van Cutsem, 1991). The polygalacturonate strands de-esterified by PME isoforms B1 and B2 should be a good substrate for PL which is formed when haustorial mother cells are differentiated and the penetration process is initiated. PL is induced at a polygalacturonate concentration less than 0·025 mg ml$^{-1}$ (Fig. 6), and the half maximal reaction velocity of the affinity-purified enzyme has been measured at a polygalacturonate concentration of 0·160 mg ml$^{-1}$ (H. Deising, unpublished data). The low substrate concentration required for PL induction, the low $K_m$, the high percentage (approx. 76%) in an extracellular location (Table 1) and the high PI of the enzyme should result in efficient and localized degradation of the pectic layers of the host cell wall. These results provide an alternative explanation for the speculations (Keon et al., 1987) that cell-wall-degrading enzymes of obligate biotrophs might be bound to the fungal cell wall in order to minimize tissue damage. Since $U$. viciae-fabae PL requires highly de-esterified substrates (35% methylation results in 90% reduction of PL activity; H. Deising, unpublished data), PME is likely to play an important role in the preparation of the infection court. The mobility of wall-degrading enzymes, however, can be regarded as critical for the degree of tissue damage (Kotoujansky, 1987; Benhamou et al., 1991). As deduced from its extreme PI (10·5), the PL should be tightly bound to the negatively charged plant cell wall at the site of its secretion. At late stages of infection structure development, i.e. when infection hyphae and haustorial mother cells are formed, the neutral endocellulases (pls 7·1 and 7·3; Heiler et al., 1993) and the basic PME isozyme A (pl 8·2; Frittrang et al., 1992) are present and these enzymes, like PL, should be tightly bound to the plant cell wall at the site of penetration. For localized wall degradation, both a tight binding and a pH

**Fig. 7.** Effect of different sugars on the formation of PME (a) and PL (b) in infection structures of $U$. viciae-fabae. Solutions contained 50 mM glucose (Glc), fructose (Fru), sucrose (Suc) or mono-galacturonate (Ga) (a), and 50 mM Glc, Fru or Suc in combination with 0·1% polygalacturonate (PGA) (b). The sugars were applied to inductive membranes inoculated withuredospores and infection structures were harvested 24 h p.i. Activities are expressed as a percentage of distilled water (a) or PGA (b) controls. Means of three independent experiments are shown and bars represent SE.
**Table 2.** Effect of *U. viciae-fabae* infection on apparent apoplastic and intracellular pH (pH<sub>app</sub>), and on cell integrity (measured as percentage extracellular malate dehydrogenase activity and electrical conductivity) of *V. faba* leaves

<table>
<thead>
<tr>
<th><em>V. faba</em> tissue</th>
<th>Apoplastic pH&lt;sub&gt;app&lt;/sub&gt;</th>
<th>Intracellular pH&lt;sub&gt;app&lt;/sub&gt;</th>
<th>Extracellular malate dehydrogenase activity (%)</th>
<th>Electrical conductivity in IWF (μS cm&lt;sup&gt;-1&lt;/sup&gt;)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>U. viciae-fabae</em>-infected leaves</td>
<td>6.69 ± 0.10</td>
<td>5.63 ± 0.06</td>
<td>0.021 ± 0.002</td>
<td>15.1 ± 0.1</td>
</tr>
<tr>
<td>Non-infected leaves</td>
<td>6.31 ± 0.11</td>
<td>5.08 ± 0.04</td>
<td>0.020 ± 0.001</td>
<td>10.1 ± 0.6</td>
</tr>
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</table>

**Fig. 8.** Effect of cycloheximide, streptomycin sulfate or ampicillin on formation of PL activity of *U. viciae-fabae*. Inductive membranes inoculated with uredospores were sprayed with 0.1% polygalacturonate alone (control, A), or in combination with streptomycin sulfate (E) or ampicillin (D) (0.5 mg ml<sup>-1</sup> each). Cycloheximide [0.1 (B) or 10 (C) μg ml<sup>-1</sup>] was applied 8 h after application of polygalacturonate. Differentiated structures were harvested 24 h p.i. Means of three independent experiments are shown and bars represent SE.

Shift towards alkaline conditions would be important. Alkalization of the medium would activate the enzymes in the order of their pH optima. A number of studies have shown that fungi growing on polygalacturonate, but not on non-pectic carbon sources such as cellulose, shift the pH of the medium towards an alkaline pH (Hancock, 1966; Wijesundera et al., 1989; Bugbee, 1990). On artificial membranes however, the pH of the pectate solution used to spray uredospores or infection structures was not altered during the experiment, i.e. for 24 h (unpublished data).

In barley leaves infected with brown rust (*Puccinia hordei*), the pH of the apoplastic sap increased from 6.6 to 7.3 in controls to 7.5 in diseased leaves (Tetlow & Farrar, 1993). In rust-infected broad bean leaves, the pH had increased from 6.3 to 6.7 by 8 d after inoculation (Table 2). The method used to determine apoplastic pH alterations, i.e. infiltration of the leaf with distilled H<sub>2</sub>O followed by low speed centrifugation to collect IWF, may be criticized since localized changes which may occur in small areas around the haustorial mother cell cannot be detected. Because of this it is probable that more pronounced and highly localized increases in pH occur in restricted cell wall areas of the *V. faba* mesophyll. Since pH values increased by more than 0.6 units have also been found in homogenates of infected leaves, and since rust infection structures show intracellular pH values around 5.4, it appears that the host plant rather than the fungus is responsible for the increase in the apoplastic pH. Low extracellular malate dehydrogenase activity demonstrates cell integrity, but it is possible that losses of OH<sup>-</sup> ions could occur locally. The increased electrical conductivity found in IWF from rust-infected leaves supports this hypothesis. One might assume that a shift in pH to 9 or higher could activate the PL of the fungus. It may also result in the release of cellulases and PMEs from the cell wall exchangers. At sites where the highly methylated pectin at the inner side of the primary wall (Liners & Van Cutsem, 1991) would not be completely de-esterified by PME, wall degradation by PL would be restricted, due to remaining methyl groups esterified to the substrate. The interplay of physico-chemical properties of the pectic enzymes resulting in tight binding to the plant cell wall, and their requirements for pH and substrate properties may be the physico- and biochemical basis for limited tissue degradation by obligate biotrophs.

Other reports on pectic enzymes from biotrophs or hemibiotrophs, for example *Bremia lactucae* (Van Pelt-Heerschap & Smit-Bakker, 1993) and *Venturia inaequalis* (Valsangiacomo & Gessler, 1992) have been published recently. The only pectate-degrading enzyme produced by *V. inaequalis* is an exopolygalacturonase. An exo-degrading mode could prevent the formation of elicitor-active pectin fragments (Nothnagel et al., 1983; Bishop et al., 1984; Bruce & West, 1989; Mathieu et al., 1991; Tenhaken & Barz, 1991) and possibly the induction of plant defence mechanisms. The fact that polygalacturonase, which is thought to be highly destructive, is formed not only by necrotrophs but also by obligate biotrophs such as *B. lactucae* (Van Pelt-Heerschap & Smit-Bakker, 1993) and *Puccinia graminis* f. sp. *tritici* (Van Sumere et al., 1957) is not in conflict with the requirements of the concept of minimal tissue damage needed for, and typically observed in, interactions involving biotrophs.
since polygalacturonases are known which do not cause maceration of plant tissue (McClandon, 1979). The fact that some biotrophs or hemi-biotrophs form either lyases or hydrolases to degrade polygalacturonate may result from adaptation to the chemical composition of the cell walls of their host plant. An enzymic adaptation to the monocotyledonous primary wall has also been discussed for cereal pathogens such as *Rhizoctonia cerealis*, *Fusarium culmorum*, and *Pseudocercospora herpotrichoides* (Cooper et al., 1988).

In general, cell-wall-degrading enzymes of plant-pathogenic fungi have been shown to be subject to catabolite repression. For example, Horton & Keen (1966) demonstrated that the formation of endopolygalacturonase and cellulase of the onion pathogen *Pyreochacta terrestris* is repressed in the presence of 5 mM glucose. Likewise, formation and secretion of a number of cell-wall-degrading enzymes, for example polygalacturonase, pectate lyase, cellulase, β-glucosidase, α-galactosidase and β-xylidosidase of the root-rotting fungus *Pyreochacta hypo- persic* is repressed in the presence of 1-4 mM glucose, 1.2 mM fructose, and 0.72 mM inositol (Goodenough et al., 1976). Holz & Knox-Davies (1986) showed that the concentrations of sugars found in onion apoplasts are sufficient to repress the formation of cell-wall-degrading enzymes in *Fusarium oxysporum* f. sp. *cessae* and these authors discuss apoplastic sugar concentration as a factor in resistance to the pathogen. This type of regulation has been described not only for hydrolases and lyases, but also for PME of the polyphagous fungus *Botrytis cinerea* (Reignault et al., 1993), suggesting common regulatory mechanisms. If, however, the formation of cell-wall-degrading enzymes of obligate biotrophs such as rust fungi are catabolite repressible by apoplastic sugars (Holz & Knox-Davies, 1986), it is very unlikely that these enzymes are available at the time when the pathogen attempts penetration of the host cell wall. Thus, repression of these enzymes by catabolites would represent a disadvantage to the fungus. Heiler et al. (1993) have shown that cellulase activity of the broad bean rust fungus is not repressed even by glucose, fructose or sucrose concentrations as high as 50 mM, but rather appears to be increased. Likewise, the present study on PL and PME has shown that neither enzyme is repressed by the concentrations of these sugars found in broad bean apoplasts. Delrot et al. (1983) report that sucrose, which is the most prominent apoplastic low-molecular-mass sugar in *V. faba* leaves, shows nymchmolar changes from 1 to 4.5 mM. Hexoses such as glucose and fructose are present at lower concentrations.

A major concern in the study of pectic enzymes of rust fungi in artificial systems is bacterial contamination of the infection structures. To test possible synthesis of pectic enzymes by bacteria associated with uredospores, inhibitors blocking eukaryotic or prokaryotic protein biosynthesis or bacterial wall synthesis have been used. Similar studies have demonstrated the eukaryotic origin of PME (Frittrang et al., 1992) and cellulases (Heiler et al., 1993), and several lines of evidence indicate that the PL studied in this paper is not due to bacterial contaminants but is produced by the broad bean rust fungus. First, the protein biosynthesis inhibitor cycloheximide blocks formation of the enzyme in a concentration-dependent manner (Fig. 8). Since relatively high cycloheximide concentrations did not completely block enzyme formation, an inhibitor of bacterial cell wall synthesis and an inhibitor of protein synthesis specific for 70S ribosomes were also used. Even at high concentrations of ampicillin and streptomycin sulfate, PL induction was not significantly affected, indicating the eukaryotic origin of the enzyme. In addition, if bacterial contaminants did account for PL synthesis the application of pectate at 0, 8 and 18 h p.i. should result in the onset of enzyme formation at more or less comparable times after application. The pattern observed (Fig. 5) suggests that a combination of morphogenesis and the presence of substrate are required for enzyme synthesis by the rust fungus. Furthermore, low-molecular-mass sugars have been shown to be potent repressors of PL induction in bacteria (Kotoujansky, 1987), whereas in *U. viciae-fabae* sugar concentrations as high as 50 mM did not result in substantial repression of either PL or PME. Also, the application of polygalacturonase to a naturally occurring population of different bacterial species, as would be present on contaminated uredospores, should result in the formation of a mixture of polygalacturonases and PLs rather than a single lyase as observed in differentiated infection structures of *U. viciae-fabae*.

In conclusion, the data presented show that the broad bean rust fungus *U. viciae-fabae* produces pectic enzymes during infection structure differentiation. In contrast to saprophytes and highly destructive necrotrophs, the regulation of the quantities of the enzymes produced, the timing of their synthesis and their physico-chemical properties appear to result in limited cell wall degradation and possibly preparation of the infection court. Pressure applied by the haustorial mother cell may then be involved in supporting growth through the enzymically softened plant cell wall and in allowing the fungus to penetrate without disruption of the cell integrity needed for biotrophy.

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