Lignocellulolytic capability of endophytic phyllosticta sp
Saowanee Wikee, P. Chumnunti, A. Al Kanghae, Ekachai Chukeatirote, S. Lumyong, Craig Faulds

To cite this version:
Saowanee Wikee, P. Chumnunti, A. Al Kanghae, Ekachai Chukeatirote, S. Lumyong, et al.. Lignocellulolytic capability of endophytic phyllosticta sp. Journal of Bacteriology and Mycology, Austin Publishing Group, 2017, 4 (2). hal-01611437

HAL Id: hal-01611437
https://hal.archives-ouvertes.fr/hal-01611437
Submitted on 5 Oct 2017

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Lignocellulolytic Capability of Endophytic Phyllosticta sp.

Wikee S1, Chumnunti P2-3, Kanghae A*, Chukeatirote E4, Lumyong S5 and Faulds CB5*

1Department of Biology, Faculty of Science, Chiang Mai University, Thailand
2School of Science, Mae Fah Luang University, Chiang Rai, Thailand
3Institute of Excellence in Fungal Research, School of Science, Mae Fah Luang University, Chiang Rai, Thailand
4Aix Marseille Universite’, INRA, Biodiversite et Biotechnologie Fongiques (BBF), Marseille, France
*Corresponding author: Faulds CB, Aix Marseille Universite’, INRA, Biodiversite et Biotechnologie Fongiques (BBF), Marseille, France

Received: February 16, 2017; Accepted: March 22, 2017; Published: March 28, 2017

Abstract

The Dothideomycetes represent the largest fungal class within the phylum Ascomycota. It is an ubiquitous class of fungi whose members span a wide spectrum of lifestyles and host interactions [1-3]. Members of the Dothideomycetes can cause disease in every major crop [4]. Approximately 1,300 genera and 19,000 species have been identified either as endophytes, plant pathogens, or as saprophytes degrading plant biomass, thus threatening agriculture and food security throughout the world [5-8]. In addition to their mode of life, the Dothideomycetes are known for producing secondary metabolites grouped into four main categories based on their biosynthetic origin: polyketides, non-ribosomal peptides, terpenoids and tryptophan derivatives [9]. These secondary metabolites can be both toxic and beneficial to plants and humankind in applications such as agrochemicals, antibiotics, immunosuppressant’s, antiparasitics, antioxidants and anticancer agents [10]. New Dothideomycetes are still being discovered worldwide and a large number of these strains remained unexplored regarding their potential use in biotechnology [11-12].

Endophytes provide a broad variety of bioactive secondary metabolites with unique structures and so this class of fungi could be used as potential "nanofactories" producing a range of “green” alternatives to currently employed chemicals [13]. Although the ecological significance of endophytes is not completely clear, it is known that these fungi can exploit dead leaves immediately after their senescence and before they fall from the tree [14], and so could be also exploited for the production of enzymes acting on plant biomass.

A well-known representative of Dothideomycetes fungi in metabolite studies is Phyllosticta (with a Guignardia anamorph). Phyllosticta species are mostly plant pathogens of a broad range of hosts and they are responsible for numerous diseases including leaf spot and black spots to spot on fruits. For example, P. ampelicida species causes black rot disease on grapevines [15]. P. musarum species causes banana black rot disease on grapevines [16]. P. citricarpa is the cause of black spot on citrus and is regarded as a quarantine pest in Europe and the USA [17], and P. capitalensis while non-pathogenic, is found also on citrus usually isolated from black spot lesions and is known as an endophyte on an extensive range of host plants [6,18]. P. cardigena and P. ericae have been reported as saprophytes [19].

Phyllosticta spp. are commonly known to produce various kinds of secondary metabolites for example, Phyllosticta derivatives exhibiting growth-inhibitory activity in five cancer cell lines have been isolated from P. cirsii. Phytotoxins, phyllosinol, brefeldin, and PM-toxin are known as fungal pathogenic derivatives from Phyllosticta [20]. Recently, the phytotoxins guignarenones A-F and alaguignardic acid have been isolated which could stimulate the development of herbicides of natural origin [21-23]. In addition, antimicrobial activity active on growth inhibitor of Escherichia coli, Bacillus cereus, and Pseudomonas aeruginosa [24-25]. P. cirsii has been isolated from diseased leaves of Cirsium arvense and it evaluated as a potential biocontrol agent of this noxious perennial weed; furthermore, it produces different phytotoxic metabolites with potential herbicidal activity when grown in liquid cultures [20].

For any fungus which attempts to colonize a higher plant, whether it is endophytic, pathogenic, saprotrophic, biotrophic/necrotrophic, it must contend with the physical barriers of the host. There is a dearth of information however on the cell wall-acting enzymes produced by Phyllosticta sp. to help in the colonization of the leaves when compared to other endophytic and pathogenic fungi, such as Botryosphaeria sp. [26], Colletotrichum sp. [27], Fusarium sp. [28],

Keywords: Agro-industrial residues; Biological pretreatment; Dothideomycetes; Lignocellulosic biomass; Lignocellulytic enzyme; Phyllosticta
Table 1: Chemical composition of lignocellulosic biomass used as growth substrates.

<table>
<thead>
<tr>
<th>Substrates</th>
<th>Cellulose</th>
<th>Hemicellulose</th>
<th>Arabinose</th>
<th>Xylose</th>
<th>Galactose</th>
<th>Mannose</th>
<th>Glucose</th>
<th>Lignin</th>
<th>Protein</th>
<th>Fiber</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rice straw</td>
<td>32.5</td>
<td>28</td>
<td>3.3±0.2</td>
<td>20.7±0.2</td>
<td>1.2±0.2</td>
<td>0.5±0.2</td>
<td>41.7±2.2</td>
<td>0.3±0.3</td>
<td>19.6</td>
<td>7.9±2.4</td>
</tr>
<tr>
<td>Rice husk</td>
<td>25-35</td>
<td>18-21</td>
<td>1.7</td>
<td>14</td>
<td>1.6</td>
<td>0.3</td>
<td>33.4</td>
<td>0.1</td>
<td>26-31</td>
<td>2.9-3.6</td>
</tr>
<tr>
<td>Wheat straw</td>
<td>38</td>
<td>29</td>
<td>3.9±0.2</td>
<td>0.5±1.1</td>
<td>5.6-10.4</td>
<td>&lt;1</td>
<td>36.3±1.8</td>
<td>0.3</td>
<td>15</td>
<td>4</td>
</tr>
<tr>
<td>Miscanthus</td>
<td>43</td>
<td>24</td>
<td>1.1</td>
<td>14.9</td>
<td>0.3</td>
<td>0-14</td>
<td>38</td>
<td>0-1.6</td>
<td>19</td>
<td>3</td>
</tr>
<tr>
<td>Sorghum</td>
<td>34-44</td>
<td>-</td>
<td>-</td>
<td>24</td>
<td>-</td>
<td>4.9</td>
<td>-</td>
<td>18-20</td>
<td>9.7</td>
<td>36.6</td>
</tr>
<tr>
<td>Lavender straw</td>
<td>43±1.7</td>
<td>13±0.9</td>
<td>3.4±0.1</td>
<td>29±0.2</td>
<td>3</td>
<td>2.5</td>
<td>45.6</td>
<td>0.9</td>
<td>23±3.6</td>
<td>-</td>
</tr>
<tr>
<td>Lavender flower</td>
<td>22.4±2</td>
<td>12.6±2</td>
<td>7.3</td>
<td>17.9±0.1</td>
<td>6.2±0.2</td>
<td>3±0.1</td>
<td>36.2±0.4</td>
<td>1.2</td>
<td>23.6±2.6</td>
<td>6.3</td>
</tr>
</tbody>
</table>

Table 2: Radial growth rate of the fungal strains cultivated in different biomass substrates.

<table>
<thead>
<tr>
<th>Fungal strains (MFLUCC)</th>
<th>Radial growth rate (cm/day)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Miscanthus</td>
</tr>
<tr>
<td>10-0137</td>
<td>0.06</td>
</tr>
<tr>
<td>12-0015</td>
<td>0.34</td>
</tr>
<tr>
<td>12-0232</td>
<td>0.34</td>
</tr>
<tr>
<td>14-0233</td>
<td>0.34</td>
</tr>
</tbody>
</table>

Macrophomina phaseolina [29], Magnaporthes [30-31], smut fungi, such as Sporisorum scitamineum [32] and Ustilago maydis [33], and Stagonospora nodorum [34].

In this study, we examined the production of ligninolytic enzymes on seven previously characterized lignocellulosic agro-industrial residues by four Phyllosticta sp.; the pathogenic P. citrimaxima and three strains of the non-pathogenic endophyte P. capitalensis.

Materials and Methods

Collection of fungal cultures

Fungal cultures were obtained from Mae Fah Luang University Culture Collection (MFLUCC): the pathogen strains Phyllosticta citrimaxima MFLUCC10-0137 was isolated from Citrus maxima, P. capitalensis MFLUCC12-0015 and MFLUCC12-0232 were isolated from Euphorbia milii and Philodendron X ‘Xanadu’, respectively. An endophyte strain P. capitalensis MFLUCC14-0233 was isolated from Hevea brasiliensis [7]. Wheat straw and miscanthus were obtained from Vivescia (Reims, France), dried, and chopped (=4 mm). Lavender straw and flowers were obtained as the residues after steam distillation from la société Bontoux SA (Saint Aubansurl’Ouveze, Drôme, France). They were collected at the beginning of September 2013, air dried for 12 days, flowers and straw separated then the fractions knife milled. Rice straw, husks and sorghum were obtained from rice cultivation fields (Phan, Thailand).

Growth measurement

Four strains of Phyllosticta were monitored on agar plates containing 15mg/ml biomass for 20 days in the different biomass media as follow; lavender flower (LF), lavender straw (LS), miscanthus (MC), rice straw (RS), rice husk (RH), sorghum (SG), wheat straw (WS). Growth was established by measuring the diameter of the growing edge of the mycelium with time. The growth measurement was recorded at day 3, 5, 7, 14, and day 20.

Preparation of fungal supernatant for enzyme essays

Growth condition in Liquid State Fermentation (LSF): The cultures were grown over 10 days in the presence of the seven biomasses in liquid medium (20 g/L), and stored at 30°C 130 rpm. The culture supernatants from all treatments were collected on days 0, 3, 7, 10, and 12 of incubation and stored at -20°C until use. The culture supernatants were then concentrated by filtration using a 0.2-micron-pore-size (polysulfone membrane; Vivaspin; Sartorius, Germany), diafiltered, and concentrated (Vivaspin polysulfone membrane with a 10-kDa cutoff; Sartorius) in 50 mM acetate solution buffer, pH5, and stored at -20°C until use [35].

Enzyme activities

Preparation of enzyme assays

Growth condition in Liquid State Fermentation (LSF): The cultures were grown over 10 days in the presence of the seven biomasses in liquid medium (20 g/L), and stored at 30°C 130 rpm. The culture supernatants from all treatments were collected on days 0, 3, 7, 10, and 12 of incubation and stored at -20°C until use. The culture supernatants were then concentrated by filtration using a 0.2-micron-pore-size (polysulfone membrane; Vivaspin; Sartorius, Germany), diafiltered, and concentrated (Vivaspin polysulfone membrane with a 10-kDa cutoff; Sartorius) in 50 mM acetate solution buffer, pH5, and stored at -20°C until use [35].

Enzyme activities

The culture supernatants produced by the four Phyllosticta strains on different lignocellulosic substrates were assayed to determine the enzyme activities present. All chemicals were of the highest purity grade available and were purchased from Sigma-Aldrich (Saint-Quentin-Fallavier, France) if not stated otherwise. Cellulose degradation was assessed by the quantification of endo-1,4-β-D-glucanase (Azocellulose) (Megazyme, Ireland) and β-glucosidase (pNP-β-D-glucopyranoside [pGlu]) activities. Hemicellulose degradation was determined by measuring the activity of endo-1, 4-β-D-Xylanase on azo-birchwoodxylan (Megazyme, Ireland), acetyl esterase on pNP-acetate (pAc), and arabino1galactanase with 1% arabino1galactan [35,36]. Oxidative enzymes were assayed with ABTS (2,2’-azino-bis (3-ethylbenzthiazoline-6-sulfonic acid) [37] as a substrate for laccase activity, Manganese Peroxidase (MnP) by the oxidation of MnSO4 [38], and Lignin Peroxidase (LiP) was assayed using veratryl alcohol as substrate in the presence of H2O2 [39]. Pectinase activity was estimated by using apple pectin and citrus pectin. Cellulase activity was determined by measuring the amount of protein present in the culture supernatant was estimated using the Bradford assay with a BSA (Bovine serum albumin) standard curve [41]. Total reducing sugars were quantified using dinitrosalicylic acid method (DNS) [42]. All essays were performed in triplicate.
Results and Discussions

Growth profiles on the different lignocellulosic biomass

In this study, we focused on lignocellulosic residues from agro-agriculture in Thailand and France such as lavender flower, lavender straw, miscanthus, rice straw, rice husk, sorghum, and wheat straw. Their chemical composition is shown in Table 1. Growth of the four pure *Phyllosticta* strains was monitored on agar plates containing 15mg/ml biomass for 20 days. Growth was established by measuring the diameter of the growing edge of the mycelium over time (Table 2).

Colonies of the phytopathogen *P. citrimaxima* (MFLUCC 10-0137) were flat, initially white with abundant mycelium, gradually becoming greenish to dark green after 2–3 days growth, with white hyphae at the margin, eventually turning black. Very slow growing ($\bar{x} = 0.1$ cm/day) in all treatments except sorghum ($\bar{x} = 0.2$ cm/day). Between days 3-7, *Phyllosticta citrimaxima* was fast growing on all substrates, stabilizing after 7 days, except for growth on sorghum, where the diameter of growth reached 4.25 cm after 20 days. The colonies of all the endophyte *P. capitalensis* were flat, initially greenish to dark green after 2–3 days, eventually turning black. Fast growing (between =0.3 and 0.35 cm/day) on all substrates, with diameter of growth of *P. capitalensis* MFLUCC12-015 reaching 8.0 cm on rice husk on day 14. On day 3-7, *P. capitalensis* MFLUCC12-015 was fast growing in every growth conditions at day 2-10, but the growths were stable after 10 days. Between day 5-7, *P. capitalensis* MFLUCC12-0232 started to producing conidia on sorghum biomass agar. *P. capitalensis* (MFLUCC 14-0233), start producing conidia on sorghum biomass agar and pycnidia could be observed on biomass residue after day 5-7 of growth (Figure 1).

The rate of growth on lavender flowers and straw was initially faster for all species, except for *P. citrimaxima* MFLUCC10-0137, especially over the first 6 days. At day 3, the growing edge of *P. capitalensis* MFLUCC12-015 mycelium reached almost 3 cm, compared to growth on the other substrates, where it reached only 0.8-1.8 cm over the same time period (Figure 2). Growth tended to stabilize after 10 days. On lavender flower and lavender stem biomass agar, the agar at the margin of the leading edge of most fungal strains changed to a red color during 1-3 days of incubation and turn darker after 7-20 days, except for *P. citrimaxima*. Most of strains could produce pycnidia and conidia on biomass residue (Figure 3) except for *P. citrimaxima*. Essential oils from plants, such as lavender, oregano, sage and mint are generally conceived as inhibitory to fungal
growth [43]. This result suggests that in the residual biomass after oil extraction may contain compound(s) which are stimulatory to growth and thus opens the possibility of using lavender by-products for biomass treatment processes within a bio refinery concept.

Growth profiles are thought to identify differences between fungal species and strains which could be correlated to their genetic enzyme compliments and expression patterns. While two Dothideomycetes with different life styles show differences during growth on Dothistroma minimal media, C. fulvus and D. septosporum showed similar growth on simple sugars (25 mM), birchwood xylan, apple pectin and lignin (1% w/v) over 2-4 weeks, although the later grew slightly better on pectin, contradictory to its lower pectinolytic gene number but supported by the number of pectinolytic enzymes expressed during growth on their respective natural plant hosts [44]. This suggests that the expression regulation of gene involved in plant deconstruction is a more dominant factor than the number of the lignocellulosic genes in the genomes.

**Enzyme activities of the Phyllosticta sp. during growth on the lignocellulosic biomass**

The culture supernatants produced during growth of the four Phyllosticta strains on the 7 lignocellulosic residues were assayed to determine lignocellulose-degrading enzyme activities as well as the total amount of protein and total reducing sugars released into the media. All fungal supernatant from lavender media showed high amount of total protein; however, the activity levels were very low. All reducing sugar appeared to be consumed at the end of cultivation of *P. capitensis* MFLUCC12-0015 and MFLUCC12-0232 while the level of reducing sugars increased during 5-7 days of cultivation of *P. citrinum* MFLUCC10-0137 and *P. capitensis* MFLUCC14-0233. No cellulase, xylanase, peroxidase, esterase or arabinofuranosidase activity could be detected. Either the fungi do not produce these enzymes or the levels are too dilute in the sample extracts for detection, even after concentration.

**P. capitensis** MFLUCC14-0233 growths

Due to the more rapid growth of *P. capitensis* MFLUCC14-0233, the fungus was chosen to be cultured on Lavender Flower Medium (LFM), Lavender Straw Medium (LSM), Wheat Straw Medium (WSM) and potato Dextrose Broth (PDB). During the days of incubation, extracellular protein from LFM was determined to be 1.76±0.75 mg, 1.81±0.23 mg, and 12.89±1.35 mg at days 5, 7 and 10, respectively, while extracellular protein on LSM was determined to be 3.49±1.51 mg protein, 0.83±0.05 mg protein, 10.95±1.38 mg protein, and 5.76±1.19 mg protein at day 3, 5, 7, and 10, respectively. Extracellular protein was no longer detected at day 12. Additionally, extracellular protein in the supernatants from WSM and PDB grown cultures could also not be detected. A small amount of protein was detected in the supernatant of LSM at day 3, dropping by day 5 and reaching the highest concentration on day 7. On the other hand, protein in LFM could not be detected at day 3, increasing after day 5 and 7 then reached maximum protein concentration at day 10. Protein levels rapidly decreased to zero at day 12 during growth on both lavender media. Reducing sugar release, as glucose equivalents, was monitored at day 3, 5, 7, 10, and 12. The highest concentration of reducing sugar could be detected on day 3 for LFM (7.36±0.86 mM) and rapidly decreased after two days. The sugar concentration in the culture supernatant from LSM at day 3 was 3.86±0.44 mM, dropping to 1.93±0.69 mM at day 12. *P. capitensis* cultured on LSM and LFM for 12 days and selected enzyme activities were determined.

### Table 3: Enzyme activities were detected after recovered extracts and concentrated through a 10 kD membrane.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Lavender flower</th>
<th>Lavender straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arabinase</td>
<td>1.8</td>
<td>3.5</td>
</tr>
<tr>
<td>Protein (mg)</td>
<td>53,287</td>
<td>45,211</td>
</tr>
<tr>
<td>Activity (U/mg)</td>
<td>1.8</td>
<td>3.5</td>
</tr>
<tr>
<td>Xylanase</td>
<td>1.8</td>
<td>1.144</td>
</tr>
<tr>
<td>Protein (mg)</td>
<td>13,238</td>
<td>31,731</td>
</tr>
<tr>
<td>Activity (U/mg)</td>
<td>1.8</td>
<td>1.189</td>
</tr>
<tr>
<td>Laccase</td>
<td>1.8</td>
<td>1.189</td>
</tr>
<tr>
<td>Protein (mg)</td>
<td>336</td>
<td>1,144</td>
</tr>
<tr>
<td>Activity (U/mg)</td>
<td>1.8</td>
<td>1.189</td>
</tr>
<tr>
<td>Pectinase</td>
<td>1.8</td>
<td>31,731</td>
</tr>
<tr>
<td>Protein (mg)</td>
<td>13,238</td>
<td>1,144</td>
</tr>
<tr>
<td>Activity (U/mg)</td>
<td>1.8</td>
<td>1.189</td>
</tr>
</tbody>
</table>

**Figure 4:** Total protein concentration (a) and total reducing sugars (b) in the culture supernatants of MFLUCC14-0233 over a 12 days period.

**Figure 5:** The production of (a) cellulase and (b) Xylanase (c) Laccase (d) Arabinase activities produced by *Phyllosticta capitensis* MFLUCC14-0233 in lavender culture medium; The production of (e) Pectinase activity against apple pectin; (f) Pectinase activity against citrus pectin by *P. capitensis* during 12 days of incubation.
Pectinolytic activity

The maximum production of arabinasein LSM occurred after 3 days of incubation (904.22 U/ml) while arabinose activity in LFM could not be detected. However, the maximum activity of arabinase produced during growth on LFM was 944.18 U/ml at day 10. Pectinase activity in concentrated extracellular supernatant of *P. capitalensis* in LFM showed a maximum activity against apple pectin of 488.39 U/ml at day 7 and 465.47 U/ml at day 10. In comparison, activity against citrus pectin was lower, with the highest activity at day 7 in LFM (263.37 U/ml) then at day 10 for LSM (245.11 U/ml) (Figure 5).

Cellulolytic and xylanolytic activity

The highest of cellulase activity (27.10 U/ml) and xylanase activity (10.85 U/ml) could be observed in cultures grown in LFM at day 5 and 7, respectively. According to the result, cellulase activity on LFM or LSM were not significantly different from day 3 to 10 the activity range between 20-27 U/ml, especially, xylanase activity during cultivation time could be detected at approximately 7-10 U/ml. Due to the low number of cellulolytic and hemi-cellulolytic CAZymes (GH6, GH7, GH10, GH11, GH45, GH61, CE1, CE3 and CBM1) present in the Capnodiales order of Dothideomycetes compared to the Hysteriales and Pleosporales order, it has been suggested that the Capnodiales (sooty molds) do not extensively degrade cellulose or that they employ another strategy for degradation [4]. *Phyllosticta/Guignardia* sp. belongs to the Botryosphaeriaceae order of the Pleosporomycetidae. The causal agent of citrus black spot, *G. citricarpa*, produced significantly more amylase, endoglucanase, cellulase or that they employ another strategy for degradation [4]. *Phyllosticta/Guignardia* sp. belongs to the Botryosphaeriaceae order of the Pleosporomycetidae. The causal agent of citrus black spot, *G. citricarpa*, produced significantly more amylase, endoglucanase, cellulase than in LFM showed a maximum activity against apple pectin of 448.39 U/ml at day 7 and 465.47 U/ml at day 10. In comparison, activity against citrus pectin was lower, with the highest activity at day 7 in LFM (263.37 U/ml) then at day 10 for LSM (245.11 U/ml) (Figure 5).

Laccase and peroxidase activity

To confirm the presence of laccase, the laccase activity in the supernatant of *P. capitalensis* grown on lavender flower or lavender straw were assayed using 2,2’-azino-bis-3-ethylbenzthiazoline-6-sulphonic acid (ABTS) as substrate. The results show that the maximum laccase activity from *P. capitalensis* in LSM at day 10 was 35.68 U/ml (Figure 5). Furthermore, no lignin peroxidase activity in concentrated extracellular supernatant of *P. capitalensis* was observed during growth on LFM was 944.18 U/ml at day 10. Pectinase activity in concentrated extracellular supernatant of *P. capitalensis* in LFM showed a maximum activity against apple pectin of 488.39 U/ml at day 7 and 465.47 U/ml at day 10. In comparison, activity against citrus pectin was lower, with the highest activity at day 7 in LFM (263.37 U/ml) then at day 10 for LSM (245.11 U/ml) (Figure 5).

Laccase and peroxidase activity

To confirm the presence of laccase, the laccase activity in the supernatant of *P. capitalensis* grown on lavender flower or lavender straw were assayed using 2,2’-azino-bis-3-ethylbenzthiazoline-6-sulphonic acid (ABTS) as substrate. The results show that the maximum laccase activity from *P. capitalensis* in LSM at day 10 was 35.68 U/ml (Figure 5). Furthermore, no lignin peroxidase and manganese peroxidase activity was detected in the culture supernatants. This indicates that these strains possibly have different mechanisms to utilize the plant biomass as a carbon source during growth.

A co-localized block of genes, including oxidoreductases, is conserved in most members of the Dothideomycetes class, and upregulated during pathogenicity, although different Dothideomycetes have different strategies for growth on their plant substrates [4]. A laccase from the hemibiotrophic fungus *Colletotrichum orbiculare* has been shown to be involved in the production of mycelial melanin during plant infection [46]. This laccase had close homologues in *C. graminicola* and *C. higginsianum*, and different from the *Magnaporthe oryzae* laccase. Dothideomycetes have a high number of conserved nonribosomal peptides synthases which resemble those proteins responsible for siderophore biosynthesis and intracellular iron storage. This later role is important for cellular function as it prevents Fenton reactions from occurring and the concomitant accumulation of reactive oxygen species. *Schizophyllum commune*, a member of the order Agaricales, lacks the order Agaricales, lacks the genes encoding lignin-acting peroxidases [4], suggesting that it degrades lignin using alternative methods, possibly using the laccases identified in the genome [47]. Laccases, however, are absent from the white-rot basidiomycete *Phanerochaete chrysosporium*, a member of the order Polyporales [48], indicating that fungi employ a vast diversity of mechanisms to breakdown lignin and related compounds.

Two Dothideomycetes species which are phylogenetically close but have different lifestyles and hosts have recently had their genomes sequenced [44]. *Cladosporium fulvum* is a biotrophic infecting tomato while *Dothistroma septosporum* is a hemibiotrophic infecting pine. Both fungi colonize their host through germinating conidia on the leaf surface producing hyphae which enter the leaf through the stomata and colonizing the apoplastic space between the mesophyll cells. Their difference in lifestyle has led to the evolution of divergent sets of genes, but with conserved genes showing their common ancestry. *C. fulvum*, which is used as a model system for plant-microbe interaction as it causes leaf mold, has the genomic set of carbohydrate-degrading enzymes, including a large pectinolytic arsenal, but many of these genes are not expressed in planta. In contrast, *Botrytis cinera*, a necrotrophic pathogen of tomato, produces a high pectinolytic activity during the invasion of the soft pectin-rich plant tissues [49]. Thus, instead of degrading the plant cell wall, *C. flavus* is thought to facilitate the local modification of the middle lamella and primary cell walls to allow better penetration of the hyphae into these pectin-rich areas, similar to that described for the forest root colonizing ectomycorrhizal, such as *Laccaria bicolor* [50].

It is not necessarily true that the enzymes secreted by the *Phyllosticta* strains are involved in plant cell wall breakdown, but could be secreted in order to remodel the plant tissue and fungal wall to facilitate growth and to fend off competitors (fungal and bacterial), as in the case with the ectomycorrhizal *Laccaria bicolor* where a reduced repertoire of carbohydrate-acting enzymes remains, but this is sufficient to aid the fungal hyphae penetrate between the root cells by loosening contact between the epidermal cells [51]. Whenever the genome of these pathologically relevant *Phyllosticta* sp. are sequenced, this will enable us to compare these biochemical assays to annotated CAZymes and to begin to understand the mechanism in which *Phyllosticta*, and in particular *P. capitalensis*, employ to degrade plant biomass.

Conclusions

*Phyllosticta* produce low levels of lignocellulolytic enzyme during cultivation on agro-industrial residues. While the fungi grow well on these carbon sources, especially those derived from the lavender oil industry, there is low production of key degradative enzymes. Further genomic and proteomic analysis is required to determine the manner in which this group of fungi utilize plant biomass for growth.

Acknowledgement

This work was supported by the Franco-Thai PHC Program (Project Code: 31824W) which gives an opportunity for scientific cooperation between Thai and France scientists. We thanks Laurence Lesage-Meessen (INRA-BBF) for providing the lavender substrates used in this experiment.
References


44. De Wit PJGM, van der Burgt A, Okmen B, Stergiopoulos I, Abd-Elsalam KA, Aerts AL, et al. The genomes of the fungal plant pathogens Cladosporium fulvum and Dothistroma septosporum reveal adaptation to different hosts and lifestyles but also signatures of common ancestry. PLOS One 2012; 8.


