Cerrena unicolor Laccases, Genes Expression and Regulation of Activity

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Abstract: A white rot fungus Cerrena unicolor has been identified as an important source of laccase, unfortunately regulation of this enzyme genes expression is poorly understood. Using 1D and 2D PAGE and LC-MS/MS, laccase isoenzymes were investigated in the liquid filtrate of C. unicolor culture. The level of expression of laccase genes was measured using qPCR. The elevated concentrations of copper and manganese in the medium caused greatest change in genes expression and three laccase transcripts were significantly affected after culture temperature was decreased from 28 to 4 °C or increased to 40 °C. The small differences in the PAGE band intensities of individual laccase proteins were also observed, indicating that given compound affect particular laccase’s transcript. Analyses of laccase-specific activity, at all tested conditions, showed the increased activities as compared to the control, suggesting that enzyme is regulated at the post-translational stage. We observed that the aspartic protease purified from C. unicolor, significantly stimulate laccase activity. Moreover, electrochemical analysis of protease-treated laccase sample had 5 times higher redox peaks. The obtained results indicate that laccases released by C. unicolor are regulated at transcriptional, translational, and at the post-translational steps of gene expression helping fungus adapt to the environmental changes.

Keywords: Cerrena unicolor; laccase; gene expression; proteomic; protease; PAGE; cyclic voltammetry

1. Introduction

Cerrena unicolor a white-rot basidiomycete, with parasitic or saprobic lifestyle, can attack several hardwood tree species, but also is proficient in decomposing dead softwoods’ logs. In order to decay a wooden tissue, fungus produce an array of hydrolytic and oxidative enzymes including: cellulases, xylanases, manganese peroxidase, lignin peroxidase, and the most extensively studied, laccase [1–3]. However, the coding potential of C. unicolor genome (https://mycocosm.jgi.doe.gov/Cerun2/Cerun2.home.html, accessed on 15 January 2021) suggests that the inventory of enzymes involved in wood degradation is far greater than the number of enzymes identified up to date. Our recent analysis of C. unicolor transcriptomes revealed about 300 transcripts coding for proteins putatively involved in sound wood degradation, of which one-fifth was differentially regulated depending...
on the type of wood substrate used [4]. Beside of lignolytic enzymes mentioned above, dye-decolorizing peroxidase, alcohol oxidase, and versatile peroxidase were identified, all are classified as lignin-metabolizing enzymes [4].

Attention has been drawn to biotechnological laccase production [1–3] and C. unicolor was identified as an organism capable of secretion of the high amounts of laccase in liquid culture [2,5]. Production of this enzyme was proven to be affected by aromatic acids [6], copper [7], cadmium [8], and temperature shock [9].

Over the past two decades, the number of reports describing applications of laccase in textiles industry [10], dye decolorization [11], wastes treatment [12], and in medicine [13] has noticeably increased. However, little is known about the physiology of C. unicolor and the role of laccase in its “lifestyle”. Analyses of genome and its annotation suggests existence of eight genes coding for laccase (Kyoto Encyclopedia of Genes and Genomes). However, in our previous work, up to 17 transcripts encoding putative laccases were found during C. unicolor growth in sawdust media [4]. Considering the importance of laccase [14] in wood degradation, it is vital to understand physiological role of C. unicolor laccases, particularly in lignin and xenobiotics degradation. Other possible biological functions of the enzyme, include involvement in fruit body formation [15] and response to environmental stress [16,17]. Recent findings suggest, that laccase primary role in living systems is protective function against adverse environmental factors including competitive or parasitic organisms and toxic compounds [17]. Little is also known about whether induction of laccase by metal ions and heat-shock occurs only at the stage of transcription (such regulation was described in Trametes velutina [18]) and which genes are involved. Fungal peroxidases and laccases have already been proven to be differentially regulated by the pH and temperature [19] or different media components [20]. Though, nutrient deficiency, copper, manganese, and aromatic compounds such as 2,5-xylidine and ferulic acid increase individual laccase gene transcripts [20,21]. Staszczyk and Jarosz-Wilkolazka [22] proved that inhibition of proteasomal activity in Trametes versicolor may stimulate laccase synthesis by cadmium ions. In this paper, we compared laccase genes expression with protein analysis and activity in C. unicolor at different growth conditions. We postulated earlier that laccase activity in this fungus may be regulated by proteases [23,24], therefore, purified endogenous aspartic-protease was incubated with C. unicolor laccases and the resulting digest was analyzed by biochemical and electrochemical methods. Results suggest that laccase synthesis by C. unicolor is regulated at transcriptional, translational, and post-transcriptional stages of the protein synthesis.

2. Materials and Methods

2.1. Growth Conditions

The strain FCL139 of Cerrena unicolor was obtained from the culture collection of the Regensburg University and maintained on 2% (w/v) malt agar slants. A piece of 0.5 cm diameter of agar hyphae per conical flask were used to inoculate Lindenberg and Holm medium [25] and the culture was grown in non-agitated conditions for 7 days at 28 °C. Mycelial mats were collected and homogenized in a disperser (IKA, Poland) and homogenate of mycelial suspension (2.5% v/v) was used as an inoculum for all studies. The fungus was grown in an orbital rotary shaker (INFORS, Bottmingen-Basel, Switzerland) at 200 rpm for 8 days at 28 °C in 100 mL Erlenmeyer flasks containing 40 mL of the Lindenberg and Holm medium. Each of the culture variants was performed in triplicates.

All stock solutions of tested compounds were prepared in water (CdCl₂, MnCl₂, and CuSO₄) or in ethanol (veratric and ferulic acids) and sterilized by filtration through Sterivex-GS filters (pore size, 0.22 µm; Millipore Corp.). Individual solutions were added to the fungal cultures on the sixth day of growth, and the growth was continued for the next 2 days. Final concentrations in the media were 1.0 mM for veratric and ferulic acids, and 10 µM for CdCl₂, MnCl₂, and CuSO₄. The final concentration of ethanol in the growth medium was always less than 0.5%, and an equivalent amount of ethanol was added to control cultures without the aromatic acid. The influence of low and high temperatures...
was also evaluated on the sixth day of growth, and the cultures were subjected to 4 or 40 °C for 2 h before analyses were performed [26]. The culture filtrates were collected using Miracloth (Sigma Chemical Co., St. Louis, MO, USA) and centrifuged (at 6000 × g, 4 °C) for 20 min in order to remove hyphal fragments. Resulting liquid was used for proteomic and the other analyses.

2.2. Laccase Activity and Protein Assay

Laccase activity was measured in triplicates, using an Infinite 200 Pro microplate reader (Tecan, Männedorf, Switzerland) in three separate biological replications with 25 μM syringaldazine (4-hydroxy, 3,5-dimetoxybenzaldehyde) (Aldrich, St. Louis, MO, USA) as a substrate, suspended in 0.1 mM citrate-phosphate buffer pH 5.3 [27]. The formation of oxidation product was recorded at 525 nm at 20 °C. Laccase activity was expressed as nkat/L. Protein concentration was determined using Coomassie Brilliant Blue G-250, dye-binding method [28].

Kinetics parameters (K_m, V_max) and electrochemical experiments for digested laccase were determined as described earlier [23].

2.3. Separation of Protein Samples by Native PAGE and SDS-PAGE

In order to separate and visualize the laccase isozymes in the culture native, 10% PAGE was used. The electrophoresis was conducted at 4 °C and 145 V. A 10-μg aliquot of the individual protein sample was loaded per gel lane. The laccase activity was visualized with 0.01% guaiacol (Sigma Chemical Co., St. Louis, MO, USA) in 0.1 M citrate-phosphate buffer (pH 5.3), at 20 °C. Simultaneously, 10% sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed as described by Laemmli [29]. After separation, proteins were visualized by silver [30] or Coomassie Brilliant Blue G-250 staining. Prestained protein ladder (Fermentas, Glen Burnie, MA, USA) was used as a molecular weight marker. A G:Box (Syngene, Frederick, MD, USA) gel documentation system was used for gel imaging.

2.4. Peptide Sequencing by LC-MS/MS

Secretome analysis was performed after 8 days of growth. The culture filtrate was concentrated 10 times using 3 kDa ultrafiltration membrane of Vivaspin 20 (Millipore, Darmstadt, Germany) and then, after measuring protein concentration, volume containing 100 ug of protein was lyophilized in a FreeZone 12, Freeze Dryer (Labconco, Kansas City, MO, USA). The individual dry sample containing 100 μg of protein was suspended in 100 μL of 100 mM ammonium bicarbonate and reduced with 5 mM tris[2-carboxyethyl]phosphine-HCl (TCEP) for 1 h at 60 °C, blocked with 10 mM methyl methanethiosulfonate (MMTS) for 10 min, and digested overnight with 10 ng/μL mass spectrometry-grade trypsin. Resulting peptides mixture was analyzed using UPLC (NanoACQUITY, Waters, Milford, MA, USA) coupled with an Orbitrap Velos mass spectrometer. The resulted output list of precursor and product ions was compared with protein database of C. unicolor (12,978 sequences: 5,371,935 residues) using the local MASCOT server (version 2.4.1). The C. unicolor v1.1 (Cerun2) genome assembly was downloaded from the U.S. Department of Energy, Joint Genome Institute (DOE JGI, http://jgi.doe.gov, accessed on 16 January 2016) and used as the reference for the peptide sequences mapping. The MS/MS analysis was performed in the Environmental Laboratory of Mass Spectrometry, Institute of Biochemistry and Biophysics of Polish Academy of Sciences in Warsaw, Poland. The protein score cut-off was set for 150.

2.5. RNA Isolation and RT-PCR

Six-day-old cultures of C. unicolor were treated as described earlier, and RNA was extracted from the mycelium after 24 h of induction. A control experiment without additions was also performed. Each sample in 2.0 mL Lysing Matrix A tubes (MP Biomedicals, Solon, OH, USA) was homogenized in a FastPrep-24 homogenizer (MP Biomedicals, Solon,
OH, USA) in 0.5 mL of lysis buffer (Qiagen, Hilden, Germany) for 2 × 20 s at the 5 m/s rotor speed. After centrifugation (10,000 × g, 4 °C, 8 min), a supernatant was used for the automated total RNA purification using an RNA Plus Mini Kit (Qiagen, Hilden, Germany) and a QIAcube robotic workstation (Qiagen, Hilden, Germany) according to the manufacturer’s protocol, which include a genomic DNA digestion step with RNase-Free DNase Set (Qiagen, Hilden, Germany). The obtained RNA was analyzed using a Qubit 2.0 fluorimeter and a Qubit RNA HS Assay Kit (Thermo Fisher Scientific, Waltham, MA, USA), as well as an Agilent 2100 Bioanalyzer and RNA 6000 Nano Kit (Agilent Technologies, Santa Clara, CA, USA). Single-stranded complementary DNA was synthesized from 1 μg of total RNA using a SuperScript® VILO cDNA Synthesis Kit (Invitrogen, Carlsbad, CA, USA), according to the manufacturer’s protocol.

A StepOne™ Real-Time PCR System (Applied Biosystems, Foster City, CA, USA) was used for the quantitative analysis of mRNA expression. The qPCR reaction was carried out with 1 μL of cDNA (1:20 diluted) in a 25 μL reaction volume using the SYBR Green JumpStart Taq ReadyMix (Sigma, St. Louis, MO, USA). For amplification of the laccase transcripts, a set of specific primers (Table S1) was used. Primes were designed with Lasergene v.8.0.2 software (DNASTAR Inc., Madison, WI, USA). The standard curves of serial 1:10 dilutions of DNA templates (Eamp = 10^{(-1/slope)}) [31] were used to calculated PCR efficiencies. Quantification of the mRNA was performed using the comparative Ct method (DDCt) with a reference gene coding for β-actin. Primers for the housekeeping β-actin gene were selected (Table S1) based on the available GenBank sequence. Amplification was carried out as follows: 40 cycles of 94 °C for 15 s and 60 °C for 1 min with an initial denaturation cycle of 94 °C for 2 min. All qPCRs were performed in a MicroAmp optical 96-well reaction plate with PE optical caps (Applied Biosystems, Foster City, CA, USA). For each experiment, a no-template reaction (NTC) was included as a negative control. All reactions were performed in triplicate.

The promoter regions sequences of C. unicolor 303 laccases were retrieved from the 1000 Fungal Genomes Project website (genome.jgi.doe.gov/programs/fungi/index.jsf, accessed on 16 June 2015). Nucleic acid sequences were analyzed using Lasergene v.8.0 software (DNASTAR, Inc., Madison, WI, USA). Database searches at the National Centre for Biotechnology Information (Bethesda, MD, USA) and the European Bioinformatics Institute (Hinxton, UK) were performed with the BLAST and FASTA.

2.6. 2-D gel Electrophoresis

Proteins from mycelium free culture fluid was concentrated by two ultrafiltration steps, first using Vivapsin 20, MWCO 3 kDa (PES; Millipore, Germany) and then Amicon Ultra-0.5 3K (Millipore, Darmstadt, Germany).

Each 5 μL aliquot of a concentrated protein sample was mixed with 125 μL of rehydration buffer containing 8 M urea, 2% CHAPS, 50 mM DTT, 0.2% (w/v) Bio-Lyte ampholyte solution (Bio-Rad, Hercules, CA, USA), and Bromophenol Blue (trace). The mixture was incubated for 30 min at 4 °C and protein sample in rehydration buffer (65 μg protein in 125 μL) was loaded onto a 7-cm IPG Ready Strip pH 3–6 (Bio-Rad, Hercules, CA, USA), covered with 2 mL of mineral oil, and then actively rehydrated for 12 h at 50 V and 20 °C, in a Protein IEF Cell (Bio-Rad, Hercules, CA, USA).

The IEF was carried out at 20 °C with a 50 μA per IPG strip limit. The settings for IEF were 250 V for 20 min (linear ramping), 4000 V for 2 h (linear ramping), and finally 4000 V for 10,000 Vh (rapid ramping). Immediately following IEF, the IPG strip was equilibrated in 2 mL of equilibration buffer I (0.375 M Tris-HCl pH 8.8, 6 M urea, 2% SDS, 20% (v/v) glycerol, and 2% DTT) and then in 2 mL of equilibration buffer II (the same equilibration buffer containing 2.5% iodoacetamide instead of DTT) for 15 min, each step on a shaker. Iodoacetamide and DTT were dissolved in equilibration buffers about 15 min before use.

Prior to second dimension separation, the IPG strips were rinsed in SDS running buffer (25 mM Tris base, 192 mM glycine, 0.1% (v/v) SDS) and then in the same buffer containing 0.0015% Bromophenol Blue. The IPG strips were placed on 12% gels (acrylamide-
bisacrylamide at a ratio of 29:1) and electrophoresed at 150 V for 2 h. As a molecular mass marker, the Thermo Scientific PageRuler Prestained Protein Ladder was used. After electrophoresis, the proteins were visualized by staining with Coomassie Brilliant Blue R-250 (0.05% Coomassie, 25% isopropanol, and 10% acetic acid) using microwave-assisted method according to [32]. Images of gels were taken with ChemiDoc™ MP Imaging System (Bio-Rad, Hercules, CA, USA).

2.7. Purification of Endogenous Aspartic-Like Protease from the C. unicolor Culture Fluid

At the first stage of purification, the post-culture fluid was centrifuged at 10,000 × g on a 6K15 (Sigma, Osterode am Harz, Germany) centrifuge for 15 min at 4 °C. Then, the supernatant was concentrated by ultrafiltration, using Pellicon 2 Mini holder with an Ultracel mini cartridge (cut-off 10 kDa) (Millipore, Bedford, MA, USA). The extracellular aspartic-like protease was purified in the two chromatographic steps using FPLC, EconoSystem (Bio-Rad, Richmond, VA). The concentrated liquid was applied on a DEAE-Sepharose (Fast Flow) (2.5 cm × 15 cm) pre-equilibrated with 50 mM acetate buffer pH 5.0 and elution of the bound proteins was conducted with a linear gradient (0–0.5 M NaCl) at flow rate of 1 mL/min. Zymographic and spectrophotometric determination of the protease and laccase activity in the obtained fractions allowed for identification of the aspartic-like protease activity in the wash fractions and selecting it for the further purification. In the second stage of purification, the same column pre-equilibrated with 50 mM Tris-HCl (pH=7.4) was used. After the elution with a step gradient (1 M NaCl), one protein peak was observed. The peak fractions were pulled and desalted using Sephadex G25 and then lyophilized on a FreeZone 18 system (Labconco, Kansas City, MO, USA).

2.8. Measurements of Protease Activity

The protease activity was assayed using 1% hemoglobin solution in 0.1 M citrate-phosphate buffer at 3.5 pH according to the Anson [33]. After incubation of 0.2 mL of sample for 60 min with 0.5 mL of substrate, the 2 mL 5% TCA was added. The precipitated undigested proteins were centrifuged out and peptide and amino acid residues remaining in solution were detected spectrophotometrically at 280 nm (1U of protease activity was equal to the amount of the enzyme able to 0.01 AU increase in 1 min/1 mg of protein). For the qualitative characterization of the appropriate protease inhibitors (pepstatin, E-64, phenanthroline, EDTA, PMSF) in the 0.1 M citrate-phosphate buffer at pH 3.5 and 5.4 or 100 mM Tris-HCl at pH 9.5 were used. After 30 min incubations with given inhibitor, the proteolytic activity of sample was measured. The protease influence on laccase proteins was determined also using 12% native PAGE separation. Electrophoretic separation of 20 µg of each sample deposited per lane was performed for 45 min at 145 V and 4 °C. Protease activity was visualized after the incubation of gels in 0.1 M citrate-phosphate buffer (pH 3.5), at 37 °C containing a 1% Coomassie Brilliant Blue (R250), Heussen and Dowdle [34]. Analysis of the gel scans was performed with a G:Box (Syngene, Frederick, MD, USA).

2.9. Laccase-Protease Experiments

The solution of the laccase (0.25 U/mg) in water was incubated in a protein ratio of 1:1 (v/v) with the solution of aspartic-like protease (0.15 U/mg) at 37 °C. In the protease-treated samples laccase activity was determined at 60 and 120 min of incubation as was described earlier [23], and resulting reaction mixtures were subjected to further kinetic and electrochemical analysis with appropriate controls.

3. Results

3.1. LC-MS/MS Analyses

The extracellular proteins of C. unicolor were analyzed by LC-MS/MS. Based on the presence of unique peptides, five laccase isoenzymes and one aspartic protease were identified (Table 1). The identified laccase enzymes had similar molecular weight but
differed in pI, which ranged from 4.44 to 5.92 pH. The calculated MW of all identified laccases was ca. 55 kDa, except one (protein ID 418196), whose mass seemed to be rather low (23 kDa) as for fungal laccase. Although, detailed analysis of gene coding for this protein (C. unicolor 303 genome) revealed repetitive sequence of unidentified nucleotides, which suggest that molecular weight of this laccase is higher than initially predicted.

Table 1. Laccases identified in the secretome of C. unicolor FCL139 by means of liquid chromatography–mass spectrometry (LC-MS/MS).

| Protein ID | Transcript ID | Score | Query Coverage (%) | MW (Da) | pI  |
|------------|---------------|-------|-------------------|---------|-----|
| Laccase    |               |       |                   |         |     |
| 193382     | XLOC_008690   | 8959  | 52                | 55139.50| 4.64|
| 390880     | XLOC_011744   | 6576  | 37                | 54248.29| 5.92|
| 418196     | XLOC_000669   | 3806  | 25                | 23445.25| 4.44|
| 408157     | XLOC_011551   | 720   | 9                 | 55484.45| 5.08|
| 364416     | XLOC_011286   | 206   | 13                | 54798.69| 5.20|
| Aspartic protease | | | | | |
| 404065     | XLOC_010406   | 218   | 9.5               | 56095.00| 4.66|

3.2. Analysis of Differentially Expressed Laccase Genes

The production of laccases proteins detected by LC-MS/MS in the C. unicolor culture filtrates were compared with qPCR analyses, which employed a set of specific primers targeting relevant mRNA transcripts (Table S1). The transcription level of individual genes was measured in cultures containing aromatic acids (ferulic or veratic), metal ions (cadmium, manganese, or copper) or after incubation at 4 and 40 °C. A classical approach was used comprising the delta–delta Ct method under a paired experimental design (treatment vs. control).

Only one gene coding for protein 364416 was found to be upregulated (5 folds) by all the three metal ions. The transcription of genes coding for laccases, 408157 and 418196 were slightly induced by copper and/or manganese ions. Temperatures affected laccase gene transcription in a less specific way (Figure 1b). The transcription of four genes coding for laccases was upregulated at 4 °C and downregulated at 40 °C. However, only transcript for laccase 390880 was repressed when fungus was grown in both temperatures. Ferulic acid upregulated the transcription of 193382 gene, whereas veratric acid stimulated only the expression of the 408157 laccase coding gene (Figure 1c). Summarizing, aromatic acids slightly downregulated the transcription of majority laccase coding genes. It seemed that gene coding for protein of 418196 gene was least influenced, whereas the expression of 364416 and 40157 genes were most frequently affected by investigated factors.

3.3. Laccase Enzymatic Activity and Electrophoretic Profiles

The qPCR analyses have shown some small changes in laccase gene expression after 24 h, therefore, we also assayed laccase activity 48 h after induction. All tested variants resulted in higher activities of laccase after 48 h in the fungal cultures (Figure 2). The results showed that the addition of manganese or veratic acid stimulated laccase activities after 48 h, up to 9 and 7 times, respectively, as compared to the control. Only slight increase in laccase activity was observed after incubation of C. unicolor culture at 40 °C as compared to 4 °C.
**Figure 1.** Effect of (A) Cu, Mn, and Cd ions, (B) temperature of 4 and 40 °C, and (C) ferulic (FER) and veratric (VER) acids on the expression of seven *C. unicolor* laccase genes. Error bars represent the standard deviations of the mean ± SD for three independent measurements (n = 3). LH – Lindeberg and Holm medium.

**Figure 2.** Effect of temperature, ferulic and veratric acids, Cu, Mn, and Cd ions on enzymatic activities of laccase after 24 and 48 h after induction of *C. unicolor*. Error bars represent the standard deviation of the mean ± SD for three independent measurements (n = 3).
In the visualized electropherograms, slight variations in the band intensities of individual laccases were noticed after culture incubation at 4 °C and addition of cadmium ions as well as in the presence of ferulic or veratric acids (Figure 3a), indicating that these compounds affected the individual laccase transcripts.

The differences in the intensities of the putative laccase bands were visible when extracellular proteins from *C. unicolor* were analyzed with SDS-PAGE (Figure 3b). In culture variants, 4 °C and veratric acid, a lower amount of protein band (denoted Lac5) was observed, whereas manganese was added to the *C. unicolor* culture the band corresponding to Lac3 was less intensive. Moreover, the intensity of the putative Lac2 band at 4 °C suggested its overproduction. In other culture variants, slight differences in the protein amounts can be observed, indicating that compounds tested, most likely affect the translations level of individual laccases.

### 3.4. 2-D Electrophoresis and LC-MS/MS

The extracellular proteins from the uninduced, 8-day old *C. unicolor* culture was concentrated and analyzed using 2D-PAGE. Most proteins focused in a pH range of 3–6 and had molecular masses ranging from 15 to 130 kDa (Figure 4). Protein spots with activity toward guaiacol were cut and submitted to LC-MS/MS sequencing. Based on the presence of unique peptides corresponding to the gene sequences in the available *C. unicolor* genome, the same five laccase isoenzymes were identified (Table S2) as identified in the culture liquid before. Nine distinct spots (L1–L9) were observed with laccase activity and each consisted of several laccase proteins.

Despite predicted molecular weights ranging from 35 to 55 kDa, the laccase protein spots were observed between 35 and 70 kDa. The computed theoretical pI values of laccases ranged from 4.3 to 5.9, whereas the experimental pI values were found in a range from 3.5 to 4.0. It should be noted that both secretomic analysis performed for the total protein from the culture fluid and LC-MS/MS analysis of protein spots derived from 2D gels indicated existence of 23 kDa protein with putative laccase properties. However, the additional resequencing of the gene coding for this laccase was performed and the results proved that our strain (*C. unicolor* FCL139) genome contains complete sequence of laccase instead of ambiguous nucleotides sequence, as was reported earlier for this species. Therefore, the proteomic results of that spots should indicate protein of MW 55 kDa instead of 23 kDa.
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3.5. Proteolytic Modifications of Laccase by C. unicolor Endogenous Protease

The steps described earlier for protease isolation allowed the separation of extracellular protein from C. unicolor culture with aspartic-like protease. The isolated protease showed high sensitivity to pepstatin (85.7% inhibition) which taken together with our LCMS/MS data suggest that it is aspartic-like protease (Table 1).

The purified laccase sample when mixed with purified protease showed increased activity to 146%, after 60 min incubation. The prolonged time (120 min) of incubation slightly elevated activity to 155%. The $K_m$ of digested laccase decreased in time from 0.54 to 0.16 mM, after 120 min of incubation (Table 2). Similar phenomena were observed for $V_{max}$ whose value dropped from 64.43 to 37.85 μM/min.

Table 2. Laccase kinetic constants after protease treatment.

|                | Time 0 min | Time 60 min | Time 120 min |
|----------------|------------|-------------|--------------|
| $K_m$ (mM)     | 0.54       | 0.19        | 0.16         |
| $V_{max}$ (μM/min) | 64.43     | 43.46       | 37.85        |

Analysis of laccase redox potential was performed by cyclic voltamperometry (Figure 5). Slow-scan voltammograms were recorded in presence of 0.2 mM ABTS pH 5 with laccase and proteolytic enzyme-coated GCE (glassy carbon electrode). When laccase and protease were incubated for 60 min a well-shaped anodic peak ($E_{p,a}$) with potential of 600 mV (vs Ag/AgCl) was observed. These peaks moved to 700 mV (vs Ag/AgCl) when enzymes were incubated for another 60 min (120 min in total). Similar difference was observed in case of cathodic peak ($E_{p,c}$) at 400 mV (120 min incubation) and 450 mV (60 min incubation). Protease-treated laccase generated five times higher peaks than in the control experiment. The current measured in the presence of protease indicated a faster rate of the catalytic reaction.
Surprisingly, only five laccases were detected in the culture medium, as demonstrated by LC-MS/MS analysis. Consequently, the genes encoding the identified laccases were further studied in case of cathodic peak (E vs Ag/AgCl) when enzymes were incubated for another 60 min (120 min in total). Similar difference was observed. These peaks moved to 700 mV (vs Ag/AgCl) when endogenous aspartic-like protease was added. The current measured in the presence of protease indicated a faster rate of incubation. Protease-treated laccase generated five times higher peaks than in the control experiment. The current measured in the presence of protease indicated a faster rate of incubation. Protease-treated laccase generated five times higher peaks than in the control experiment.

Considering multifunctional laccase nature, it would be interesting to assign the expression of specific genes of individual laccases to their different physiological functions. In this work, the liquid medium of Lindeberg and Holm was chosen for the production of high amounts of laccase [7]. This enzyme plays an important role in lignolysis, and in other fungal physiologically processes, such as a response to harsh environmental conditions [9,36]. It was suggested that in Polyporales high number of laccases isozymes is the result of multiplication of a single ancestral gene and their further specialization [37].

4. Discussion

A number of reports indicate that C. unicolor culture is capable of production of high amount of extracellular laccase [7,35]. This enzyme plays an important role in lignolysis, and in other fungal physiologically processes, such as a response to harsh environmental conditions [9,36]. As this fungus lives in the Northern hemisphere, where seasonal temperature fluctuates, low (4 °C) and high (40 °C) temperatures were chosen as extreme physical conditions. The obtained results demonstrated that laccase production is influenced by changing temperature in the fungus growth environment. Copper and cadmium ions have already been suggested to stimulate laccase production in C. unicolor cultures [7,8], which was also confirmed during this study. Different lignin derivatives were proposed to influence laccase synthesis in the genus Cerrena [6] and our research confirmed this phenomenon. Taking into consideration that laccase is lignin-degrading enzyme, it is surprising that not all laccase genes are upregulated by aromatic acids, but it seems correlated with our previous results when not all genes were influenced by different sawdust in the medium [4]. Our results support the concept that the laccase expression is the response to different environmental factors, occurs at both transcriptional and translational levels and enzyme is also regulated at post-translational stage. Since laccase synthesis is triggered by different chemical and physical factors, it is reasonable to expect that the promoter regions of respective genes containing various regulatory elements are responsible for regulation of expression. Although the genomic sequence of the C. unicolor FCL139 strain is not available, we searched the promoter regions of the genome of C. unicolor 303 to identify putative regulatory elements (Table S3). It was suggested that Heat Shock Elements (HSE) present in the laccase promoter may be involved in the response to temperature changes in the environment [39,40]. Although laccase gene, 390880 has the highest number of six
HSE in its promoter region, its transcription is hardly affected by temperature in *C. unicolor* FCL139. Similarly, the promoter region of the laccase 193382, as the only one among the examined, possesses a putative ACE1 element responsible for regulation by copper ions, while the expression of laccase 364416 without ACE1 was strongly upregulated by all the metal ions used. Further detailed analysis of *C. unicolor* laccase expression is needed to understand their genes regulation. There is a chance that its genes may be clustered with other genes coding for wood degrading enzymes and therefore expressed together. Moreover, still little is known about laccase expression beside promoter elements. It should be emphasized that no metal-responsive elements (MRE) were found in the *C. unicolor* laccase promoters, which is in contrast with the respective laccase gene 364416 induction observed in the presence of cadmium. In the *C. unicolor* FCL139, only one laccase promoter region was sequenced and analyzed [41]. Comparing the respective promoter regions in *C. unicolor* 303 and FCL139 strains, slight differences were noticed in the number of hypothetical ACE1 and metal regulatory elements (MRE). Considering the observed differences in the DNA sequence of the respective promoter regions between fungal strains, further studies of laccase genes transcriptional regulations are needed, which may explain their contribution to the level of laccase biosynthesis. Additionally, the complexity of the regulation of *Cerrena unicolor* laccase synthesis and secretion results from the fact that the process is presumably affected by endogenous proteases [23,24], which was proved in taxonomically related *Trametes versicolor* [42]. We observed that in the *C. unicolor* culture filtrate (free of mycelium) laccase isoenzymes number, Osinska-Jaroszuk et al. [43], was reduced from five to two when incubated both in room and 4 °C temperature for prolonged time (up to 30 days). The obtained results in the presented manuscript of laccase activity assays, electrophoretic profile of laccases, and LC-MS/MS analyses demonstrated differences in enzyme biosynthesis in response to the tested factors. However, comparing these data to the results of transcriptional analyses, some differences can be observed, suggesting further, e.g., translational regulation of laccase activities. Therefore, based on LC-MS/MS data, one endogenous aspartic-protease was purified and used to digest laccase. The obtained results confirmed our previous observations [23], it seems that in nature laccase, kinetic and electrochemical properties may undergo post-translational proteolytical processing and despite 2 h incubation time, laccase was not completely digested as it was in case of animal proteinase K [23]. Perhaps, it is the way that hyphae recycle amino acid from secreted proteins since wood is considered a poor nitrogen source. It should be noted that the *C. unicolor* cultures exposed to different visible light conditions show also strong induction of laccase and protease activities [24]. Therefore, it is possible that proteases secreted by *C. unicolor* not only play role in plant matter digestion but also are an important post-translational enzymes regulator/recycler.

5. Conclusions

Our previous results and findings in this paper confirm that *C. unicolor* laccase plays an important role in fungus adaptation to harsh environmental conditions, including presence of metal ions or xenobiotics and adaptation to adverse physical factors such as temperature or light. We postulate that the number of laccase genes facilitate the saprotrophic lifestyle of the fungus allowing it to decompose several kinds of wood and different isoenzymes might promote adaptation to changing chemical and physical growth conditions. Even if recent studies emphasize transcriptional regulation of laccase expression, still little is known about post-translational processing and secretion of this enzyme. Therefore, further studies of laccase post-translational processing and activation by native proteases is warranted.

**Supplementary Materials:** The following are available online at [https://www.mdpi.com/2218-273X/11/3/468/s1](https://www.mdpi.com/2218-273X/11/3/468/s1), **Table S1:** Primer sequences targeting individual laccase-coding genes used in qPCR and the length of amplification products, **Table S2:** Laccases identified in the secretome of *C. unicolor* FCL139 by means of 2D electrophoresis and liquid chromatography–mass spectrometry (LC-MS/MS), **Table S3:** Putative regulatory elements in the promoter regions of *Cerrena unicolor* 303 laccase genes.
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