Article

Phenols Removal from Hemicelluloses Pre-Hydrolysate by Laccase to Improve Butanol Production

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Abstract: Phenolic compounds are important inhibitors of the microorganisms used in the Acetone-Butanol-Ethanol (ABE) fermentation. The degradation of phenolic compounds in a wood pre-hydrolysate, a potential substrate for the production of ABE, was studied in this article. First, physicochemical methods for detoxification such as nanofiltration and flocculation were applied and the best combination was selected. With a flocculated sample, the concentration of phenolic compounds decreases from 1.20 to 0.28 g/L with the addition of a solid laccase at optimum conditions, which is below the phenolic compounds limit of inhibition. This results in an increase in butanol production, more than double, compared to a pre-hydrolysate non-treated with laccase enzymes.

Keywords: phenolic compounds; detoxification; butanol; pre-hydrolysate; laccase enzymes

1. Introduction

The acetone-butanol-ethanol (ABE) fermentation was developed in the late 19th and the early 20th century and is one of the first fermentation processes adapted to a large scale. The main reason for this was the growing need of acetone for munitions factories during the two World Wars [1]. Today, the production of butanol and its derivatives is attractive for international markets [2]. In addition, butanol is a high-energy fuel, less corrosive than ethanol, and its derivatives are used for the production of latex, plasticizers and coatings [3]. Although ABE fermentation with Clostridium acetobutylicum has been well known, improvements are still being proposed to decrease the cost of substrate and the subsequent operating costs. In a previous work conducted by Ajao et al. [4,5], the detoxification of a wood pre-hydrolysate, obtained from a dissolving Kraft pulp mill, was conducted by filtration and flocculation, prior to ethanol production by fermentation.

In the current study, the pre-hydrolysate was used as a substrate to grow C. acetobutylicum and to produce butanol. During the pre-hydrolysis step, phenolic compounds which are lignin residues are generated [6,7]. They can damage the structure of the cell membrane and the ability of microorganisms to absorb sugars, which makes them very toxic for the ABE fermentation [8]. The reduction of the phenolic compounds concentration by flocculation was significant [4]; however, it was not sufficient to improve fermentation yield. The conventional treatment of the pre-hydrolysate cannot reduce the concentration of phenolic compounds below the inhibition limit, as set by Mechmech et al. [9].

In this work, the degradation of the phenolic compounds using solid and liquid laccase enzymes was investigated to complement the physicochemical treatment. Laccases are copper-containing enzymes and can oxidize several types of phenolic compounds [10–12]. The enzymatic degradation of
phenolic compounds was investigated on aqueous solutions inspired by wastewater treatment [13–15]. These studies have been done with laccases or peroxydases, oxidative enzymes similar to the laccases. The method can significantly improve the degradation of phenolic compounds, thus reaching the thresholds that do not allow inhibition. There are two important aspects to understand the relevance of this work: the world population is concerned about the environment protection and the revalorization of resources can be an additional income for pulp and paper mills. The detoxification of lignocellulosic biomass by laccases was investigated to increase the quality of the substrate in ethanol production [16]. In the current study, it is shown that the degradation of the phenolic compounds prior to fermentation using laccases can increase the production of butanol from a wood pre-hydrolysate and improve the cost-effectiveness of the process. In North America, where the pulp and paper industry is under transformation, looking to implement innovative processes, it is important to maximize the reuse of resources in a biorefinery context [17].

2. Materials and Methods

2.1. Microorganism, Culture Maintenance and Inoculum Preparation

The culture preparation was performed as described by Mechmech et al. [9]. *C. acetobutylicum* ATCC 824 was obtained from the American Type Culture Collection (ATCC) and was cultured in sterilized Reinforced Clostridium Medium (RCM), its composition being (in g/L): tryptose 10, beef extract 10, glucose 5, yeast extract 3, soluble starch 0.5, sodium chloride 5, L-cysteine-HCl 0.5. The culture was kept under anaerobic conditions at 37 °C for 18–22 h, with shaking at 110 rpm, until an Optical Density (OD$_{600}$) of 1.9–2 was obtained. Glycerol 50% v/v was added to the bacterial culture to obtain a final concentration of 25% and the stock culture was immediately frozen at −80 °C until use. For complete anaerobic conditions, a drop of sodium sulfide nonahydrate was added in a cryogenic tube.

The inoculum for ABE fermentation was prepared in a RCM medium. The medium was first boiled and then purged with a gas mixture of 80% N$_2$ and 20% CO$_2$ to remove oxygen from the culture media. The culture was inoculated to a proportion of 1/2000 with stock culture. Culture conditions were identical to those of the stock culture. Inoculum was ready to be inoculated at an OD$_{600}$ between 0.6 and 0.9.

2.2. Fermentation Medium

The fermentations experiments were carried out in a complex fermentation medium. The medium was composed of 60 g/L of xylose and resazurin 0.001% base. Before sterilization, the base was first boiled and then purged with a gas mixture of 80% N$_2$ and 20% CO$_2$ during 5 min. After sterilization, a purged sterile solution of 200 g/L of yeast extract was added in a ratio of 1/40. A filtered mixture with KH$_2$PO$_4$ 50 g/L, K$_2$HPO$_4$ 50 g/L, ammonium acetate 220 g/L, para-aminobenzoic acid 0.1 g/L, thiamin 0.1 g/L, biotin 0.001 g/L, MgSO$_4$·7H$_2$O 20 g/L, MnSO$_4$·H$_2$O 1 g/L, FeSO$_4$·7H$_2$O1 g/L and NaCl 1 g/L was added in a ratio of 1/100. In some experiments, a phenolic compound was added to the xylose solution to maintain a specific concentration. Screw capped Schott bottles of 250 mL were filled with 200 mL of complete culture media and used for anaerobic fermentation. The culture media for the hydrolysate test consists of 195 mL of treated hydrolysate and 5 mL of yeast extract. Xylose (60 g/L) was added in order to compare with the control solution. Before inoculation with 10 mL of inoculums, the bottles were slightly open in anaerobic jars containing Gas Pak envelopes (BD Gas Pak™EZ Anaerobe Container System, Franklin Lakes, NJ, USA) with indicators (BD BBL™ Dry Anaerobic Indicator Strips, Franklin Lakes, NJ, USA) for 48 h to create perfect anaerobic conditions in the mixtures. If necessary, a solution containing 5% p/v of Na$_2$S·9H$_2$O was added in a ratio of 1/400 to eliminate the traces of oxygen. After inoculation, the cultures were incubated at 37 °C, 110 rpm and with a pH control. During fermentation, 5 mL samples were periodically withdrawn to analyze OD$_{600}$, residual xylose and alcohols. All fermentation experiments were performed in duplicate.
2.3. Analytical Methods

Spectrometer (Pharmacia Biotech Novaspec® II, Piscataway, NJ, USA) was used to monitor the growth of *C. acetobutylicum* and to determine the total phenols concentration with the Folin–Ciocalteu reagent method [18].

Gas chromatograph (GC 7890A, Agilent Technologies, Santa Clara, CA, USA) with an OV 624 capillary column and a flame ionization detector (H₂ flow rate: 30 mL/min; air flow rate: 2.23 mL/min) was used to measure butanol, acetone and ethanol concentrations in the fermentation medium after the inhibition test [9].

High performance liquid chromatography (HPLC Agilent Technology, Germany) was used to determine the concentration of vanillin, catechol, syringaldehyde gallic acid and simple sugars. To measure phenols concentration, separation was made using a mixture of 15% acetonitrile and 85% phosphoric acid 10 mM on a Nucleosil C18 (150 X 4.6 mm) column with a diode array detector (DAD) at 313 nm and 280 nm. Dilution for phenol analysis was performed in the same solvent to reach a maximum concentration of 500 ppm. To measure the simple sugar concentration, a refractive index detector and an EC Nucleodur RP-NH₂ (250 mm × 4.6 mm, 5 µm) were used with a mixture of 75% acetonitrile and 25% deionized water as a solvent. A temperature of 40 °C and a flow rate of 1.5 mL/min were applied for better separation [9]. Dilution for simple sugar analysis was performed in 50% acetonitrile and 50% deionized water to reach a maximum concentration of 10 g/L.

2.4. Degradation with Laccase Enzymes

For degradation assays, two types of laccase enzymes provided by industrial partners were used: a solid (or dehydrated) laccase and a liquid laccase. All degradation tests were performed in a test tube (14 mm diameter) with a 5 mL mixture composed of syringaldehyde, vanillin, catechol and gallic acid with the same concentrations. Total phenolic compounds concentrations of 2 g/L, 4 g/L and 6 g/L were used. After pH adjustment with 4M NaOH, a fixed dose of laccase was added and the test tube was incubated for 7h at the appropriate temperature and rotation at 180 rpm. When a large volume of flocculated hydrolysates was detoxified by laccase, an erlenmeyer was used with a ratio erlenmeyer volume/flocculated hydrolysate volume of 2.5. A preliminary screening was done to determine the optimal temperature and pH conditions for the subsequent experiments. The samples were analyzed once; however, occasional duplicate analyses were conducted to check the method used. Since the fermentation and enzyme degradation tests were performed, respectively, in duplicate and in triplicate, irregular results were easily detected and corrected.

2.5. Preparation and Treatment of Hydrolysates

Detoxification approaches described by Mechmech et al. [9] were first applied before the degradation of the phenolic compounds with laccase. To extract the hemicelluloses, a mixture of 60% aspen and 40% maple wood chips was treated with hot water and steam in a pilot digester at FPInnovations (Pointe-Claire, Québec, Canada). That pre-hydrolysate was forwarded to the Centre National en Électrochimie et en Technologies Environnementales (CNETE). Two different methods were applied to detoxify the pre-hydrolysate. First, pre-hydrolysate was filtered through the organic membrane NF270 (Molecular weight cut-off 200–400 Da), then hydrolyzed with 1.5% w/w sulfuric acid at 121 °C for 60 min to increase the monomeric sugars concentration. After these treatments, it was coagulated/flocculated. The pH was raised to 6.5 and ferric sulfate (Fe₂(SO₄)₃) with a ratio 1 g Fe/1 g phenolic compound was added. Flocculation experiments were carried out in jar tests with an agitation of 150 rpm for 10 min., then 50 rpm for 30 min. In a second step, the same operations were performed, but without applying the nanomembrane filtration.
3. Results and Discussion

3.1. Optimization of the Degradation Conditions

The screening of the solid and liquid laccase enzymes indicates that their efficiency is strongly affected by temperature and pH. The results with a 2 g/L synthetic solution containing syringaldehyde, vanillin, catechol and gallic acid in equal quantity have shown that the degradation of the phenolic compounds by laccases is optimal at a pH 8 and a temperature of 50 °C (Figures 1 and 2).

![Figure 1. Degradation of phenolic compounds by laccase enzymes at different pH.](image)

![Figure 2. Degradation of phenolic compounds by laccase enzymes at different temperatures.](image)

The optimal dose of the solid laccase is 100 mg of enzyme/g of phenolic compounds. The results are similar when a dosage of 200 mg/g is used; however, for economic reasons, it is preferable to use 100 mg enzyme/g of phenolic compounds (Figure 3). At optimal pH, temperature and enzyme dose, the degradation of the phenolic compounds by the solid laccase is 79%. The most relevant parameters for this type of laccase are the pH, the dose of laccase and the initial concentration of phenolic compounds ($p < 0.01$). Moreover, there is an interaction between the initial concentration of phenolic compounds and the dose of laccase; therefore, the dose of enzyme should be adjusted according to the initial concentration of phenolic compounds. An interaction regression coefficient of $-0.24$ between...
the dose of laccase and the initial concentration of phenolic compounds and a correlation value of 0.83 were calculated.

The optimal dose for the liquid laccase is 5 mL enzyme/g of phenolic compounds (Figure 3). At optimal conditions, the degradation of the phenolic compounds by liquid laccase is 33%. For economic reasons, it is suggested to use 3 mL/g of phenolic compounds. When 3 mL of laccase enzyme are used, a degradation percentage of 29% is reached. It is important to mention that for the liquid laccase, the impact of the pH on the degradation efficiency is not significant \((p > 0.05)\) and for practical reasons an initial pH of 8 was used. On the other hand, the temperature, the dose of laccase and the initial concentration of phenolic compounds are critical for the efficiency of the liquid laccase. A statistical analysis was performed and the results have shown a non-significant effect of pH and an important effect of the dose of laccase, the initial concentration of the phenolic compounds and the temperature. The regression coefficients for the three parameters were, respectively, 0.56, 0.55 and \(-0.49\), with a \(p\)-value < 0.01.

According to the results on the degradation of phenolic compounds with laccases on various stages of the conventional detoxification presented in Section 3.4, the best results were obtained for a sample with pH 8. It is important to point out that there is a link between the degradation time and the dose of laccase and the initial concentration of phenolic compounds and a correlation value of 0.83 were calculated.
the added dose of laccase \((p < 0.01)\). In fact, the more the laccase dose was high, faster the degradation was. The control sample without laccase demonstrated a degradation of the phenolic compounds of 20\% in time (Figure 3). Therefore, a part of the degradation was performed naturally and not by the laccases. The corresponding equation for the naturally occurring phenolic degradation in time was: \( y = 0.011x^2 - 0.12x + 1.8 \) with a correlation value of 0.77. The degradation mainly occurs during the first hour, after that there was a stabilization of the phenolic compounds concentration.

### 3.2. Influence of the Individual Phenolic Compounds

The degradation of the phenolics is influenced by the type of individual phenolic compounds in the mixture. When the solid laccase was used, gallic acid was always degraded in the first two hours of incubation, followed by catechol which was entirely degraded too, then syringaldehyde and vanillin which were partially degraded at the end of incubation (Figure 4). Although it is not shown in the figure, it seems that there is a link with the temperature. When the temperature is lower (40 and 30 °C), the degradation of gallic acid and catechol is accelerated, while the other phenolic compounds are not degraded. This implies a change in the conformation of the enzyme to make it suitable for the degradation of syringaldehyde and vanillin at high temperature. The conformation of the enzyme can explain that preference. Usually, when the number of alcohol group increases, the speed of degradation increases too. The laccases directly affect the alcohol groups by oxidation of a reducing substrate and the formation of a free radical [10]. The accessibility of these groups can also influence their oxidation and affect the degradation order. The nature of the individual phenolic compounds is therefore a critical factor for the degradation.

![Figure 4](image_url)

**Figure 4.** Degradation of the individual phenolic compounds using the solid laccase enzyme as a function of time.

The liquid laccase was efficient for gallic acid and syringaldehyde degradation, but had a limited effect on vanillin and catechol, as shown in Figure 5. This confirms the importance of the type of phenolic compounds on the degradation by laccases. Gallic acid is degraded first, then syringaldehyde with 74\% and 68\% degradation, respectively. Only 35\% and 29\% of the catechol and vanillin, respectively, are degraded by the liquid laccase. These results are lower, compared to the solid laccase.
To determine the capacity of laccase to degrade a concentrated mixture of phenolic compounds, their degradation at different initial concentrations was studied. At the same conditions and proportional dosage, there is a better degradation at the lowest initial concentration (Figure 6). With initial concentration of 2 g/L of phenolic compounds, the degradation is 79%, at 4 g/L it drops to 55% and at 6 g/L only 25% of the phenolic compounds are degraded by the solid laccase. It is important to point out that the increase of the dosage up to 200 mg/g of phenolic compounds can partially offset the increase of the initial concentration. Indeed, with 200 mg/g of phenolic compounds, the degradation reaches 38% with an initial phenolic compounds concentration of 6 g/L. It is 13% more than with only 100 mg/g of phenolic compounds. There is a link between the dosage and the efficiency at high concentrations of phenolic compounds. For the liquid laccase, an initial concentration of 2 g/L of phenolic compounds allows a degradation of 25% (Figure 6). However, with 4 g/L, only 15% of the phenolic compounds are degraded and with 6 g/L, 14%. Regarding the liquid laccase, the difference between the results was due to an increase in the concentration of syringaldehyde, catechol and vanillin. It was difficult to degrade these compounds by the liquid laccase and the residual amount was higher than expected. For the solid laccase, a similar phenomenon occurs. Regardless of the initial concentration (2, 4 or 6 g/L), gallic acid will always be degraded entirely in 2 h. The catechol degradation is around 100%, although it is slightly less effective when the initial concentration of the phenolic compounds increases to 6 g/L. The residual phenolics are vanillin and syringaldehyde. It seems to be difficult for the solid laccase to degrade those phenolic compounds at higher concentrations.
3.4. Hydrolysate Detoxification by Laccase

Finally, the efficiency of the laccase enzymes on pre-hydrolysate samples pretreated with different detoxification methods were tested. As shown in Figure 7, the best degradation of the phenolic compounds occurs in the flocculated non-filtrated hydrolysate, and by the solid laccase, at its optimum conditions. In fact, this is the only case where laccase sufficiently degrades phenolic compounds to a concentration below the minimum level of inhibition determined by Mechmech et al. [9]. The level of degradation can reach a leftover of phenolic compounds of 0.28 g/L, under the limit of inhibition of ABE fermentation of 1.1 g/L in flocculated media. However, none of the other samples can reach this limit or the limit of 0.3 g/L for the non-flocculated sample. Theoretically, the percentage of degradation is expected to be similar from one sample to another, but in practice, the composition of each sample may vary significantly. For example, phenolic compounds proportion can be different between flocculated samples and untreated samples. The results show a significant difference in the degradation of the phenolic compounds for each intermediate stage of detoxification. However, there is a link between the enzyme type and the efficiency of detoxification by laccase at each intermediate stage (Figure 7). The liquid laccase seems to be more efficient when used before flocculation, with a degradation percentage of 42% for the untreated hydrolysate and 36%, for the filtered hydrolysate. Two hypotheses are envisaged. First, it is possible that the flocculation removes mainly phenolic compounds that would also be degraded by the liquid laccase, thus reducing the total amount of initial phenolic compounds to be degraded. Then, the residual ferric sulfate used for flocculation can form a complex with the laccase, thus making the enzyme less effective or inactive. The solid laccase is very efficient on flocculated hydrolysate and less efficient on untreated hydrolysate. The hydrolysate composition is complex and may be the source of the variation in the effectiveness of the laccase enzymes.
In this work, it has been shown that laccase enzymes efficiently degrade phenolic compounds in wood hydrolysates. At an optimum temperature of 50 °C, pH of 8 and enzyme dose of 100 mg/g of phenolic compounds, the degradation of the phenolic compounds reaches 77%. The use of laccase for wood hydrolysate detoxification reduces the phenolic compounds concentration to 0.28 g/L, far below the limit of inhibition. The hydrolysate detoxification combining flocculation and laccase enzymes enhances the efficiency of the ABE fermentation, by decreasing the concentration of the phenolic compounds under the limit of inhibition. In the case of the liquid laccase, no butanol production occurs. The reason for this outcome should be investigated, but a negative interaction between the bacteria and the enzyme, or its degradation products is suspected.

4. Conclusions

In this work, it has been shown that laccase enzymes efficiently degrade phenolic compounds in wood hydrolysates. At an optimum temperature of 50 °C, pH of 8 and enzyme dose of 100 mg/g of phenolic compounds, the degradation of the phenolic compounds reaches 77%. The use of laccase for wood hydrolysate detoxification reduces the phenolic compounds concentration to 0.28 g/L, far below the limit of inhibition. The hydrolysate detoxification combining flocculation and laccase enzymes prior to fermentation increases the amount of butanol produced.

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Conflicts of Interest: The authors declare no conflict of interest.
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