Growth-Promoting Effects of Press Water from the Mechanical Drying of Douglas-Fir Wood Chips on Lignocellulolytic Fungi

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Research Article

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Abstract

Background:

The mechanical drying of wood chips is an innovative method that improves the heating value of sawmill by-products in an energy-efficient continuous process. The liquid that comes out of the wood chips as press water (PW), however, contains a variety of undissolved as well as dissolved organic substances. The disposal of the PW as wastewater would generate additional costs due to its high organic load, offsetting the benefits in energy costs associated with the enhanced heating value of the wood chips. Our research explored if the organic load in PW could be utilized to boost the growth of cellulolytic filamentous fungi. Hence, using the industrially relevant Ascomycete *Trichoderma reesei* RUT-C30 as well as several Basidiomycete wood-rotting fungi, we examined the potential of press water obtained from Douglas-fir wood chips to be used in growth and enzyme production media.

Results:

The addition of PW supernatant to liquid cultures of *T. reesei* RUT-C30 resulted in a significant enhancement of the endoglucanase and endoxylanase activities with a substantially shortened lag-phase. Supplementation with PW allowed to replace several mineral ions (Fe$^{2+}$, Mn$^{2+}$, Zn$^{2+}$, Ca$^{2+}$, Mg$^{2+}$, K$^+$) from the liquid media without negative effects on the enzymatic activities. Concentrations of PW above 50% showed no adverse effects regarding the achievable endoglucanase activity but affected the endoxylanase activity to some extent. Exploring the growth-enhancing potential of several individual PW components after chemical analysis revealed that the observed lag-phase reduction of *T. reesei* RUT-C30 was not caused by the dissolved sugars and ions, nor the wood solids in the PW, suggesting that other, so far non-identified, compounds are responsible. However, also the growth rate of several basidiomycetes was significantly enhanced by the supplementation of PW to the agar medium. Moreover, their cultivation in liquid cultures reduced the turbidity of the PW substantially.

Conclusions:

PW was identified as a suitable media supplement for lignocellulolytic fungi, including the cellulase and xylanase producer *T. reesei* RUT-C30 and several wood-degrading basidiomycetes. The possibility to replace several minerals, trace elements and an equal volume of fresh water in liquid media with PW and the ability of fungal mycelia to filter out the suspended solids is a promising way to combine biological wastewater treatment with value-adding biotechnological applications.

Introduction

Sawmills generate large quantities of by-products, such as bark, sawdust, and wood chips. The average yield of sawn timber from round wood in sawmills has been reported to be around 60%, which means around 40% of the round wood ends up as a by-product [1–3]. The total production of wood chips, particles and residues in Germany reached an estimated 10.97 million m$^3$ in 2019 [4]. Wood chips are
often sold to thermal power stations or to the pulp and paper industry. Nonetheless, a large portion of these by-products is generally used for internal energy production [2]. With average moisture contents of 40 to 50 %-wet basis (wb) after production, microbial activity is facilitated during storage, which can increase the risk for spontaneous combustion due to heat accumulation inside the piles [5] as well as to considerable dry matter and energy loss [6–8]. Technical drying is therefore necessary to achieve suitable moisture contents that both prevent microbial activity and guarantee high fuel quality [9, 10].

Conventional drying methods based on the evaporation of moisture mostly use thermal energy from dedicated combustion of biogenic or fossil fuels [11, 12]. Regardless of the type of thermal dryer, energy efficiency and drying rate are critical issues associated with thermal drying methods [13]. Mechanical dewatering, on the other hand, is a process that uses high pressure [14]. The increased pressure within the wood will force the free water in the cell lumen to find multiple ways out, e.g. through the pits or cracks in the cell walls [15, 16]. The energy required in compression and thus for expulsion of the water is much lower than the energy required to vaporize the same water using thermal energy [17]. The combination of mechanical squeezing and thermal drying can therefore lead to substantial energy savings compared exclusively thermal drying [18, 19].

Large quantities of press water (PW) are released during the mechanical dewatering of wood chips. Ultimately, the generated volume of PW will depend on the quality and quantity of the wood chips being processed at sawmills. Since the PW originates from the free water in the lumen of the wood cells, it will contain chemical wood constituents, such as dissolved minerals, sugars, or other extractives. These substances could have negative effects for aquatic ecosystems if the disposal of press water is not performed adequately [20, 21]. Among the PW of different wood species, e.g. Pine, Fir, Spruce, Poplar, and Beech, Douglas-fir showed the highest phenol index, chemical and biochemical oxygen demand [22]. Moreover the chemical oxygen demand (COD), which describes the amount of oxygen required to oxidize the organic material, of the tested PW was above 10,000 mg L\(^{-1}\) with an acidic pH, and none of the tested PWs showed a 100 % degradability of the water-soluble organic substances (Zahn-Wellens-Test) with Douglas-fir PW having the least degradability with only 83 % [22]. Therefore, similar to the effluents of pulp and paper mills, PW must be treated before being released into water bodies or even before entering the local wastewater treatment plants [23, 24]. Depending on the local regulations for wastewater disposal, the economic costs of PW treatment could offset the benefits associated with the reduced energy costs of mechanical drying compared to an exclusively thermal drying [18, 19]. Hence, alternative applications should be examined.

Instead of viewing it as a wastewater, the organically loaded PW could act as a low-cost substrate for enzyme production in fungal cultivations, thus becoming a by-product of the mechanical drying and not a waste. The ability to degrade lignocellulose is widespread in fungi of the Ascomycota and Basidiomycota phyla [25]. The ascomycete filamentous fungus *Trichoderma reesei* is a major producer of carbohydrate active enzymes (CAZymes), which are used for the conversion of plant biomass to sustainable fuels and chemicals [26–28]. The high cost of enzyme production as an important factor for the economic feasibility of biorefineries, especially for the production of biofuels, has driven the development of several
hypercellulolytic mutant strains [29–31]. One of these strains, RUT-C30, has been extensively studied in academic research [32]. Additionally, wood-decaying basidiomycetes, which are typically classified as white rots or brown rots, have evolved alongside their plant hosts and developed distinct strategies for the degradation of lignin along cellulose and hemicelluloses [33–35]. White rots produce multiple lignin-degrading peroxidases, which allow them to degrade lignin, aside from other cell wall components like cellulose and hemicelluloses [36, 37]. On the other hand, brown rots leave lignin largely intact due to a lack of lignin-degrading peroxidases [36, 38]. These lignin-degrading enzymes make basidiomycetes interesting for industrial and biotechnological applications [39].

Generally, media for fungal enzyme production require high amounts of carbon and nitrogen sources. Aside from that, fungal media require considerable amounts of other macro nutrients, like sulfate, phosphate, K\(^+\), Ca\(^{2+}\) and Mg\(^{2+}\). Furthermore, several micro elements, like Fe\(^{2+}\), Mn\(^{2+}\), Zn\(^{2+}\), Co\(^{2+}\), are required. While necessary in much lower concentrations, these are nevertheless very important for the production of enzymes and other cofactors [40, 41].

The carbon source, which optimally serves both as an energy source and inducer molecule, accounts for a substantial portion of the cultivation costs. Therefore, the identification of abundant low cost residues, which display good rheological properties, are non-toxic, and induce cellulase production, have led to the utilization of various side products, like lactose and sugarcane bagasse [42, 43]. To this end, the PW could become an interesting side stream of the sawmill industry, which could have a huge potential to be used as media supplement in the production of cellulolytic enzymes.

For this study, we chose *Pseudotsuga menziesii* (Douglas-fir) as the PW source, since it displayed the highest organic load among other wood species [22]. Furthermore, Douglas-fir is the most widespread non-native tree species in Germany, covering 2% of the total forest area, with an expected increase in relevance in the coming years due to its desirable wood properties, small vulnerability to summer drought, and volume growth per hectare that exceeds that of native European tree species [44–46].

The aim of the present study was to analyze the potential of Douglas-fir-derived PW as a low-cost media supplement in the cultivation of wood degrading filamentous fungi. Since little is known about the PW composition, we initially investigated relevant physico-chemical characteristics of the PW, like dissolved ions, sugars, amount of solids and composition of the solids. We then tested the production of cellulases and xylanases with *T. reesei* RUT-C30 during growth on PW-supplemented media. Furthermore, the ability of several basidiomycetes to grow on PW-containing media and to clarify the aqueous medium containing suspended PW-solids was investigated (Fig. 1).

**Results**

**Chemical oxygen demand**
The chemical oxygen demand (COD) of PW was analyzed to evaluate its organic load and the overall environmental impact. The results show that pure PW has a rather high organic load with a maximum of 10,750 mg L\(^{-1}\). The removal of macro solids from the PW by centrifugation lowered the COD to 8,800 mg L\(^{-1}\). Ultimately, the high COD values of the PW are indicative for a high concentration of dissolved organic substances that are problematic in wastewater treatment but could be beneficial for fungal cultivations.

**Physico-chemical analysis**

The analyzed PW was a heterogeneous suspension containing solids of different sizes and shapes (Fig. 2a). Even after centrifugation the liquid fraction remained turbid and colored, suggesting a high quantity of dissolved substances and suspended particles (Fig. 2b). Furthermore, the rather low pH of 4.42 indicates the presence of acidic organic substances.

**Phenotype of *T. reesei* RUT-C30 on PW agar**

To characterize the growth phenotype of *T. reesei* RUT-C30 on different media including or excluding PW, agar plates were inoculated with the same concentration of conidia and cultivated under identical conditions. Three types of Mandels-Andreotti (MA) salt-based solid media were prepared: plain MA agar without any carbon source, MA agar with 0.2 g L\(^{-1}\) glucose, and MA agar with PW supernatant but without additional glucose. To observe the growth of *T. reesei* also on PW alone, one additional PW medium was prepared without MA salts as a control.

A clear growth difference was observed, particularly regarding the color of the conidia, which turned green on PW but stayed yellow on media without PW (Fig. 5). Moreover, *T. reesei* yielded a significantly higher spore number on the plates with PW. The addition of glucose to the MA agar only slightly increased sporulation, and PW medium without the addition of MA salts was not sufficient for optimal growth (Fig. 5).

**Liquid cultivation in PW supernatant with surplus of nutrients**

In the following, we cultivated *T. reesei* RUT-C30 in liquid cultures with different concentrations of PW supernatant (after centrifugation) to observe the effects on endoglucanase and endoxylanase production. Here, the liquid cultures were formulated with the complete MA medium and hence, the ion concentrations varied slightly, depending on the used PW volumes.

The enzymatic activities of the culture supernatants showed substantial enhancement in PW media compared to the control condition (Fig. 4a). Most noticeably, only the PW-supplemented cultures
displayed enzymatic activities already at day 3, whereas the control media with 0 % PW needed several days more to reach the same level of activity. The activities seen in the culture supernatants with PW continued to be significantly higher than the controls throughout the cultivations, except for the endoxylanase activity in 100 % PW, which was noticeably lower than the other PW conditions. However, the gap between the activities measured in 0 % PW and 25 – 75 % PW decreased over the course of the cultivation period. For instance, the endogluanase activities measured in the PW supernatants were on average 8.8-fold higher at day 5 and only 2.5-fold higher at day 7 compared to the control without PW. Similarly, the endoxylanase activity measured between 25 % and 75 % PW displayed a 2.3-fold and 1.9-fold increase relative to the control at day 5 and 7, respectively. No significant difference in enzymatic activities was observed among the cultures with different PW dilutions. These results demonstrate that PW addition can robustly enhance the production of cellulase and xylanase enzymes in *T. reesei* RUT-C30 over a broad concentration window.

**Liquid cultivation in PW with replacement of salts and C-source**

Since PW contains several nutrients in relevant quantities (mainly trace elements but also theoretically hydrolysable solids; Table 1, Fig. 2d), we wanted to test whether some of the components of the conventional fungal cultivation medium (MA) could be replaced by raw PW (with solids, not centrifuged). The minimal PW concentration at which the trace elements could be eliminated from the MA medium was 55 % raw PW, as calculated based on the zinc concentration (Table 1). Since cobalt was not measured in the PW it had to be supplemented separately. However, at this dilution level, iron and manganese concentrations are higher than in the conventional MA medium. Particularly high concentrations of Fe$^{3+}$ ions were reported to be inhibitory for the saccharification of cellulose in other fungi [47]. Therefore, to elucidate whether an overdose of trace elements would have negative effects, we prepared a control condition with 50x trace elements. Furthermore, the carbon source Avicel was replaced based on the theoretically hydrolysable solids in the PW (Table 2). The soluble monosaccharides were not considered in this case.

The distribution of the volume-weighed particle size of the PW supernatant was measured by laser diffraction. The particles in the PW supernatant (after centrifugation at 4,000 rcf for 15 min) showed a monomodal distribution, with most of the particles displaying a diameter of 1 µm and a maximum particle size of ca. 100 µm (Fig. 2c). Further separation of these suspended particles would require centrifugation at higher speeds, longer settling times or extensive filtration. The PW was fractionated by means of centrifugation and membrane filtration into sedimented macro solids, filtered micro solids, and dissolved substances, which were quantified gravimetrically. The total concentration of solids and dissolved substances in PW amounted to 10.38 ± 0.07 g L$^{-1}$ (Fig. 2d). Dissolved substances represented the largest fraction of the PW with 6.31 ± 0.04 g L$^{-1}$. The concentration of macro solids (at
3.40 ± 0.30 g L^{-1}) and micro solids (at 0.67 ± 0.05 g L^{-1}) together accounted for about 40 % of the total dry mass.

An acid hydrolysis was performed to further investigate the composition of the macro solids in PW (Fig. 2e). Three fractions were obtained and quantified, namely hydrolysable polysaccharides (Hydrolysate), acid-insoluble residues (AIR), and acid-soluble aromatics (ASA). The hydrolysable polysaccharides in the solids of the PW accounted for 51.2 % (w/w). Assuming that these are predominantly cellulose and hemicelluloses, a maximum of 1.73 g L^{-1} of the PW solids could potentially serve as a carbon source and inducer to produce cellulolytic enzymes in fungal fermentations.

To better elucidate the composition of the acid-insoluble residues (making up 47.3 % (w/w)), we performed a pyrolysis GC/MS to identify the major pyrolysis products. The largest peaks were distributed in three main groups, corresponding to phenols (2-methoxy-4-methylphenol, 1,2-dihydroxybenzene, 2-methoxy-4-vinylphenol), fatty acids (oleic acid), and phytosterols (campesterol, stigmasta-3,5-diene) (Fig. 2f). The absence of levoglucosan was a good indication for the successful hydrolysis, since it is a pyrolysis product from carbohydrates. These results suggest that the polysaccharides in the PW solids were effectively hydrolyzed leaving only the phenol-rich polymers and some resin derivatives behind.

**Dissolved phenolic compounds**

To analyze the dissolved phenolic compounds we concentrated the PW using a solid phase extraction (SPE) on C18 silica and the methanol soluble fraction was analyzed using GC/MS. The chromatogram (Additional file: Fig. S6) showed the presence of some phenolic substances, like 3-(4-hydroxyphenyl)-1-propanol, and sugar derivatives. However, the flavonoid taxifolin was found to be the most abundant compound in the PW at a concentration of 7.4 % (w/w) in the extract.

**Dissolved ions**

The most abundant ions measured according to standard methods for the examination of water in the PW were potassium (K^{+}; 130 mg L^{-1}), calcium (Ca^{2+}; 49.2 mg L^{-1}), and SO_{4}^{2-} (38 mg L^{-1}) (Table 1), followed by iron (Fe^{2+}; 20.10 mg L^{-1}), magnesium (Mg^{2+}; 14.60 mg L^{-1}), manganese (Mn^{2+}; 4.83 mg L^{-1}), sodium (Na^{+}; 4.28 mg L^{-1}), and zinc (Zn^{2+}; 0.61 mg L^{-1}). As expectable from wood, PW was found to be a poor source of nitrogen with nitrate concentrations below detection range and only 1.43 mg L^{-1} of ammonium.

**Sugar analysis**

The most abundant monosaccharides in the PW measured using HPAEC-PAD were fructose and glucose with 0.60 g L^{-1} and 0.24 g L^{-1}, respectively, followed by galactose (0.09 g L^{-1}) and arabinose (0.03 g L^{-1}).
Xylose \((13.81 \text{ mg L}^{-1})\) and mannose \((2.77 \text{ mg L}^{-1})\) were the least abundant detectable sugars. The disaccharide concentrations of cellobiose and sucrose were \(2.77 \text{ mg L}^{-1}\) and \(1.9 \text{ mg L}^{-1}\), respectively. The combined concentration of mono- and disaccharides in the PW amounted to \(0.99 \text{ g L}^{-1}\) (see Additional file: Fig. S1).

Comparable to the results seen on PW supernatant, the effect of the raw PW was most noticeable at day 3, where endoglucanase and endoxylanase activities of the cultures were significantly higher than the MA control (Fig. 4b). The highest endoglucanase activities were observed after 7 days of cultivation in 55% \((11.5 \pm 1.2 \text{ U mL}^{-1})\) and 25% PW \((11.3 \pm 0.5 \text{ U mL}^{-1})\). The endoxylanase activities showed a clear advantage at 25% PW, which yielded significantly higher activities than the other PW conditions, reaching a maximum activity of \(20.3 \text{ U mL}^{-1}\) after 7 days. PW concentrations above 55% showed no significant enhancement of the endoxylanase activity.

The 50-fold increased concentration of trace elements in the medium led to overall slightly higher enzymatic activities, but the difference was only significant for the endoglucanase activities at day 5 compared to the 1x control.

These results indicate that it is possible to replace some of the salts, Avicel, and trace elements present in MA medium by PW while maintaining the enhanced endoglucanase and endoxylanase activities up to 55% raw PW. However, as demonstrated by the 50x trace elements control, the growth enhancement seen in the PW cultivations can only partially be explained by the increased concentration in salts or trace elements. Nevertheless, no significant inhibition was observed due to the surplus of iron.

**Liquid cultivation with addition of free sugars simulating 25% PW**

To determine whether the free sugars present in the PW contribute to the observed enhanced enzymatic activities, we simulated the conditions present in the 25% PW by supplementing MA media with glucose, fructose, arabinose, galactose, and cellobiose in the respective concentrations (Fig. 3b). The addition of free sugars to the cultivation medium resulted in no significant difference compared to the control condition with only Avicel (Fig. 4c). On the other hand, the addition of 25% PW supernatant recurrently resulted in significantly higher enzymatic activities compared to the two conditions without PW. This suggests that the free sugars in PW are not responsible for the observed beneficial effects of PW during the cultivations.

**Liquid cultivation with PW solids as C-source**

To estimate to what extent *T. reesei* RUT-C30 can use the PW solids as a carbon source and inducer, liquid cultures were prepared with 1% ball-milled PW solids, non-pressed Douglas-fir wood and Avicel as standard carbon source. Although the fungus seemed to grow, there was a clear difference in the development of the fungal biomass in the liquid cultures with PW solids and Douglas-fir wood vs. the Avicel-grown cultures (Fig. 5b – d) and almost no enzymatic activities could be measured in the culture.
supernatants, even after 10 days (Fig. 5a). These results suggest that *T. reesei* is unable to utilize the PW solids or the Douglas-fir wood powder as a carbon source.

**Growth rate of several basidiomycetes in the presence of PW**

The growth rate of a series of wood-degrading basidiomycetes was measured on agar plates containing yeast malt extract agar (YEMA) supplemented with PW from Douglas-fir (25 % and 75 % v/v) to assess the potential of wood PW as a substrate for lignocellulolytic fungi (Fig. 6a). Considering the ability of the tested strains to degrade lignin, we decided to use a batch of PW from Douglas-fir wood chips but with bark (PWB), which had a darker color and higher organic load (Additional file: Fig. S2).

We found that PWB enhanced the growth rates of all tested white rot fungi but not for the brown rot *Rhodonia placenta*. The strongest improvement was observed for *Pleurotus ostreatus*, for which the growth rate was accelerated from 0.18 mm h⁻¹ (0 % PW) to 0.54 mm h⁻¹ (25 % PWB), representing a 3-fold improvement over YEMA. Despite showing a less drastic effect, a clear tendency of faster growth with increasing PWB concentrations was also observed for the used strains of *Dentipellis fragilis*, *Schizophyllum commune*, *Hericium coralloides* and *Trametes versicolor* (Fig. 6a).

**Bio-clarification of the PW**

To assess the potential to reduce the turbidity of PW (with and without bark), as beneficial trait for biological wastewater treatment, we cultivated *T. versicolor*, *G. applanatum*, and *P. chrysosporium* in liquid cultures of PW supplemented with potato dextrose liquid medium (PDY) and measured the optical density of the culture supernatants. A turbidity reduction compared to controls incubated without fungal biomass was observed for all the tested strains in PW and PWB. The highest turbidity reduction was observed in *T. versicolor* cultures (Fig. 6b). Furthermore, all strains displayed a change in coloration of the PW, indicating the uptake, degradation or metabolization of some PW components (see Additional file: Fig. S4, Fig. S5).

**Discussion**

The optimization of the wood chip drying process by implementing mechanical drying in conventional sawmills could lead to higher energy efficiency and reduced fuel consumption during the drying process. Thus, this seemingly small improvement could have considerable effects on the entire sawmilling process [48]. The COD values of the PW are comparable with effluents seen in the pulp and paper industry (thermo-mechanical pulp processing effluents and black liquor), which cause considerable operating costs due to the requirement of wastewater treatment [49–53]. Therefore, the discharge and treatment of large PW volumes could require more energy than the energy saved by the mechanical drying compared to the conventional thermal drying, hence becoming a financial burden [18]. Although no correlation between the COD and the amount of dissolved and undissolved substances in the PW was
observed, it could be used as an orientation value to estimate the PW quality. Thus, higher COD and PW volumes, which translate to higher concentrations of substrates, would be beneficial for fungal fermentations. In parallel, the utilization of the PW would reduce the need for freshwater in the fermentation process, which is an important aspect for a more sustainable industry [54].

Wood type, felling season, and the location in which the trees have been growing ultimately define the composition and concentration of dissolved substances in the PW [55, 56]. A direct comparison of sugar concentrations found in the PW and literature values is limited, since the substances in the PW are not actively extracted using solvents, temperature, nor hydrolyzing the wood. Glucose and fructose can be found as free sugars in the xylem of Douglas-fir, although in low concentrations compared to the branches or foliage and it might vary depending on biotic and abiotic growth influencing factors [56, 57]. We were unable to detect rhamnose in the PW, which coincides with the low rhamnose concentrations observed in Douglas-fir wood [58]. The occurrence of cellobiose, the characteristic building block of cellulose, shows that there is a certain mechanical degradation of the cell wall, because it is not a naturally occurring disaccharide in wood [59]. Although Douglas-fir as a typical softwood contains more mannans than xylans (see Additional file: Table S2), we observed considerably lower mannose concentrations compared to xylose within the PW (Fig. 3b). Also the arabinose and galactose concentrations were higher than the expected concentrations found in wood compositional analyses [58, 60, 61]. The branched structure and the nature of the neighbouring molecules influences the stability of the hemicelluloses. It is therefore possible that mannose is more stably embedded in the cell wall than xylose and thus more resistant against extraction by mechanical pressure [61, 62].

One of the concerns of supplementing the PW to fungal media was the possible presence of bioactive substances, like phenolic compounds from lignin or tannins, that could inhibit fungal growth [63, 64]. In contrast to the wood-degrading basidiomycetes, T. reesei RUT-C30 lacks significant capacity to degrade lignin [65]. Considering that bark is the most extractive-rich tissue of Douglas-fir, the growth experiments of T. reesei were made using PW from debarked wood chips, thus minimizing any unforeseen interaction with potential inhibitors [66–68].

One of the major limitations of the PW as a substrate for liquid cultures is the lack of a nitrogen source, which is indispensable for fungal growth and enzyme production. However, this is a common issue among lignocellulosic substrates from agro-industrial or industrial wastes, such as sugarcane bagasse, soybean hulls, pulp and paper sludges, and even pretreated wood chips [29, 69–71], the only exception being manure [72]. Compared to the aforementioned substrates, the PW is not a rich source of cellulose and hemicelluloses, hence the supplementation of a carbon source, like Avicel, is necessary. Despite this, the PW was able to significantly enhance the enzymatic activities in the culture supernatants when supplemented to the medium (Fig. 4a). In contrast to PW, the utilization of several agro-industrial and industrial wastes has some limitations regarding the production of cellulolytic enzymes. The utilization of sugarcane bagasse and molasses often requires genetic strain optimization to overcome the effect of repressing sugars or the utilization of sucrose [29]. In the case of waste paper sludge, nonproductive binding of the substrate or inhibition from mineral paper additives are factors that render this substrate
unsuitable for the enzyme production [69, 73]. Similarly, steam-pretreated wood as a carbon source and inducer for enzyme production is limited by the presence of inhibiting sugar degradation products [71]. Sophorose and lactose are more suitable supplements for the production of cellulases in *T. reesei* as soluble inducers and carbon sources [74]. However, sophorose is prohibitively expensive and lactose, although cheap and largely available, induces only a limited amount of cellulases compared to cellulosic substrates [75–78].

The presence of lignin in the PW solids was confirmed by Py-GC/MS of the acid insoluble residues (Fig. 2f), corresponding to typical lignin fragments found in pyrolyzed Douglas-fir wood [79, 80]. Additionally, the presence of some small phenolic substances and one characteristic flavonoid found in Douglas-fir, taxifolin, was confirmed in the methanolic fraction separated with SPE (C18 silica) of the dissolved substances in the PW. However, contrary to the literature reports, no substantial inhibition of endoglucanase activity and only a slight reduction of endoxylanase activity was observed, despite the presence of phenolic substances in the PW (see Additional file: Fig. S3) [66, 81–83]. Moreover, the lack of nutrients, especially a nitrogen source, had a stronger influence on the growth of *T. reesei* than the presence of PW (Fig. 3a). These results indicate that the concentration of potentially inhibiting compounds was insufficient to cause significant inhibition of *T. reesei* enzymes.

At low PW concentrations (25 – 55 % PW) the enhancing effect for enzyme production was similar for the cultivation with PW supernatant and with raw PW. The enzymatic activity of the cultures with PW was consistently higher after 3 days of cultivation compared to the control, even after the replacement of salts and trace elements (Fig. 4b). The consistent results suggest that the addition of 25 – 55 % PW to the cultivation medium has a stronger effect on the enzymatic activities than varying the concentration of salts or trace elements in the medium (Fig. 4a – b). On the other hand, at higher concentrations (75 – 100 % PW) the measured enzymatic activities were less consistent. The endoxylanase activity was the most affected by the increased concentration of raw PW (Fig. 4b) compared to the same conditions with PW supernatant (Fig. 4a). Since the ion concentrations had no measurable effect on the enzymatic activities, the only difference between the PW supernatant and raw PW was the presence of PW solids.

The carbon source and inducer molecules constitute a major cost factor in the production of lignocellulolytic enzymes, and thus finding cheaper alternatives would be beneficial [29, 84]. The (partial) replacement of Avicel by the solids in PW has the potential to reduce some of the costs. However, *T. reesei* RUT-C30 was unable to grow on PW solids as sole carbon source as well as on non-pressed Douglas-fir wood powder (Fig. 5a), suggesting that the cellulose in the solids is too difficult for *T. reesei* to access efficiently in comparison to the microcrystalline cellulose of Avicel [65]. This observation is also in line with reports about *Trichoderma spp.* preferably growing on wood that has been previously degraded or pre-treated either chemically or by other fungi [85, 86]. Therefore, the reduced concentration of Avicel in the cultivations with raw PW-replaced media could have contributed to the reduced endoxylananase activity at higher PW concentrations in the late stages of the fermentations. Moreover, the measurable reduction of the endoxylananase activity in presence of PW (Additional file: Fig. S3) suggests that inhibitory effects might also be due to direct interactions of PW components with the secreted enzymes.
Lignin has been suggested to bind hydrophobic faces such as for example on the cellulose-binding module of cellulases, thus inactivating the enzymes by non-productive binding [87–90]. Consequently, it is possible that enzymes bound to the surface of the PW solids, thus reducing the enzymatic activity of the culture supernatants at higher concentrations of raw PW (Fig. 4b) [89]. In addition to that, several phenolic compounds generated from lignin degradation have been reported to inhibit a GH11 endoxylanase from *Thermobacillus xylanilyticus* in a non-competitive multisite mechanism [91]. This suggests that some components in the PW might interact specifically with the secreted endoxylanases. On the other hand, phenolic substances in the black liquor from the Kraft pulping process have been reported to modify the protein structure of commercial xylanases, enhancing the hydrolysis of xylan [92]. This might have contributed to the enhanced activities seen at low PW concentrations (25 – 55 %), while at higher PW concentrations (75 – 100 %) the positive effects are offset by the non-productive binding on the PW solids and inhibition by other substances.

Cellobiose is known to induce the production of cellulases in *T. reesei* [93]. Furthermore, oligosaccharides derived from cellulose have a strong induction effect on the cellulase expression [94, 95]. However, the addition of surplus cellobiose and other monosaccharides to the cultivation medium corresponding to the concentrations found in 25 % PW showed no significant effect by itself (Fig. 4c). Consequently, neither the dissolved sugars, nor the suspended solids of the PW were found to be causative for the positive effects of the PW cultures, and that the minerals had only a small positive influence (Fig. 4b). Therefore, we suggest that the enhancing effect might come from other, so far unidentified, organic molecules dissolved in the PW. There is a possibility that some components in the PW also directly interact with some signaling pathways or induce some modifications on the hyphal cell walls, similar to N, N-dimethylformamide, which was reported to enhance cellulase production via calcium signaling and permeabilization of the hyphal cell wall in *T. reesei* RUT-C30 [96]. It has been suggested that the micromorphology of *T. reesei* influences the enzyme productivity [97]. Although the micromorphology of *T. reesei* RUT-C30 was not measured in this study, an analysis could reveal if the enhanced enzymatic productivity seen in the PW cultures is related to changes in hyphal morphology. A deeper understanding of the composition of the organic fraction of the PW combined with an RNAseq analysis of *T. reesei* RUT-C30 in presence of PW to identify internal signaling processes would allow to elucidate how the PW interacts with the fungus and induces higher endoglucanase activities.

The growth experiments on agar plates with the white rots *P. ostreatus*, *T. versicolor*, *D. fragilis*, *S. commune*, *H. coralloides* and the brown rot *R. placenta* were made using PWB as a supplement, which contained more dissolved substances than PW (Additional file: Fig. S2, Table S1). The tested white rot strains showed a significant growth enhancement regardless of the PWB concentration, except for *D. fragilis*, where an increment from 25 % to 75 % caused a growth rate reduction (Fig. 6a). Interestingly, *R. placenta*, the only brown rot strain that was tested, showed no significant growth rate change in the presence of PWB. While more species will have to be tested, it is intriguing to speculate that the differences seen among the basidiomycetes is related to the presence or absence of ligninolytic enzymes [35, 36]. Altogether, no growth inhibition was observed on all the tested fungal strains, hence the PW of Douglas-fir was considered non-toxic. A deeper examination of the effect of PWB components on the
transcriptome of white rots would be required to determine the effect of PW on the expression of lignin-degrading peroxidases of basidiomycetes.

One interesting alternative to the conventional wastewater treatment that might offer a quick solution for the high turbidity and high COD would be fungal-assisted bio-clarification of wood PW. Our results demonstrated that the turbidity was substantially reduced after a few days of cultivation (Fig. 3b). Fungal mycelia entrapping the suspended solids in the PW is a practical solution to overcome longer settling times or extensive filtration [98]. The advantage of using wood-degrading basidiomycetes goes beyond the capture of suspended solids. We observed a decrease in absorption over the entire visible spectrum (230 – 800 nm) (Additional file 4; Figure S3), indicating some degradation of dissolved substances in the PW probably due to the secretion of ligninolytic enzymes [99–101], which nevertheless needs to be verified.

**Conclusion**

The PW of Douglas-fir wood chips is a complex sawmill side stream with a high COD, demanding costly wastewater treatment. However, several of the dissolved components as well as the solids are also part of conventional fungal growth media, and despite the presence of potential phenolic inhibitors, the PW was found to be a suitable non-toxic media supplement for several basidiomycetes as well as the industrially employed ascomycete *T. reesei*. The supplementation of cleared PW to liquid cultures of *T. reesei* RUT-C30 reduced the lag-phase and significantly enhanced the endoglucanase and endoxylanase activities in the supernatant. The supplementation of 55 % PW to the cultivation media allowed the replacement of 100 % trace elements (Fe^{2+}, Mn^{2+}, Zn^{2+}) of the conventional MA medium as well as 12 %, 13 % and 6 % of Ca^{2+}, Mg^{2+}, and K^{+}, respectively, without losing the positive influence on enzyme production. Furthermore, PW allowed to replace an equal volume of fresh water. The utilization of PW in fungal cultivations could therefore combine a bio-clarification of this sawmill effluent with the creation of added-value by lowering costs of media formulations and an increased product yield.

A further analysis of the molecular pathways being activated by PW as well as an advanced qualitative and quantitative chemical analysis will therefore be crucial to understand the mechanisms of the positive effects seen in this study and pinpoint the responsible compounds for the growth-enhancing effects of the PW on the tested fungal strains.

**Materials And Methods**

**Press water samples**

The press water used in this study was obtained by pressing debarked wood chips of *Pseudotsuga menziesii* (Douglas-fir) with a roller press (wood chips squeezer) located at Bohnert Technik GmbH in Seebach, Germany. Douglas-fir wood chips with and without bark were obtained from local sawmills. The wood chips were fed evenly on a patented plate conveyor chain, which passes between two large rollers
Physico-chemical analysis of PW

The PW was centrifuged at 4000 rcf for 15 min in a centrifuge (Heraeus Megafuge 40R, Thermo Scientific). The PW supernatant was carefully decanted, aliquots of 100 mL were transferred to lyophilisation flaks, and the same was done for the sediment. The total mass content was determined gravimetrically after drying the samples in a freeze dryer (Christ Alpha 1-2LDplus). Fine particles that remained suspended after the centrifugation were quantified gravimetrically using cellulose acetate filter membranes with pore size of 0.45 µm (Sartorius).

Particle size distribution

A laser diffraction system HELOS (Sympatec, Germany) with the RHODOS dispersing unit was used to measure the particle size distribution over a wide range of sizes of the PW supernatant.

Chemical oxygen demand

The chemical oxygen demand (COD) and dissolved ions concentrations were measured by the Chair of Urban Water Systems Engineering at the Technical University of Munich, according to the German standard methods for the examination of water [102–105].

Acid hydrolysis

The determination of acid insoluble residues in the solids of the press water was conducted according to the standard procedures TAPPI T 249 and T 222. Briefly, 1 g of the lyophilized and homogenized PW solids was incubated for 2 h with 15 mL of 72 % sulphuric acid in a water bath at 20 °C. Then, the acid was diluted to 3 % and incubated for 4 h at 100 °C. After hydrolysis the sample was filtered, the acid-insoluble residue (AIR) was determined gravimetrically and the acid-soluble aromatics (ASA was determined spectroscopically at 205 nm (ε 110 g L⁻¹ cm⁻¹) [106–108].

Py-GC/MS and GC/MS analysis and sample preparation

Solid samples were measured using the pyrolysis gas chromatography coupled to a mass spectrometer (Py-GC/MS), VLMSD 5975C (Agilent Technologies) equipped with a VF17 MS 30 m x 250 µm x 0.25 µm column (Agilent Technologies). The pyrolysis was performed in a single shot analysis at 450 °C for 0.2 min. For liquid samples, PW was filtered followed by an extraction step using a SPE C18ec cartridge.
and methanol to elute the hydrophobic substances (Chromabond, Macherey-Nagel). The GC/MS conditions: Inj. 300 °C with a split of 40:1 and temperature program: $T_1 = 100$ °C for 1 min, $R = 10$ °C/min, $T_2 = 300$ °C for 4 min. Liquid samples were silylated prior to injection. The mass spectra were evaluated using the NIST MS library (NIST20). Taxifolin, was verified and quantified using a calibration curve with a reference (Sigma-Aldrich).

**HPAEC-PAD analysis of the PW**

Free neutral sugars (list all sugars) were determined on a Dionex ICSW 3000 HPAEC-PAD instrument setup with a Dionex AS Autosampler, a Dionex gradient mixer GM-3 (Dionex Corp., California USA) and a CarboPacPA1 preparative IC column (4 x 250 mm) equipped with a CarboPacPA1 standard bore guard column (4 x 50 mm) (Thermo Fisher Scientific Inc., Massachusetts USA). The analysis was carried out with a 27 min isocratic method with a 10 mM sodium hydroxide solution for monosaccharides in deionized water with low total organic carbon at 1 mL min$^{-1}$ flow rate and 30 °C. For the analysis of disaccharides, the sodium hydroxide concentration was elevated to 100 mM.

The PW samples were filtered with a PES membrane with a pore size of 0.2 µm and cleaned using an anion exchange SPE cartridge (Strata XA, Phenomenex) following the protocol of the manufacturer. The samples were further diluted with ddH2O before measurement in duplicate.

**Strain cultivation**

*T. reesei* strain RUT-C30 was propagated on potato dextrose agar (Carl Roth) plates in the dark at 30 °C for two days, then switched to constant light at 25 °C for conidiation. All liquid media cultivations were carried out in 250 mL flasks without baffles containing 50 mL medium and shaken at 250 min$^{-1}$ (25 mm throw) and at 30 °C in darkness, if not otherwise mentioned. For the inoculation, a respective volume of conidial suspension was added after optical density measurement to a final concentration of $10^6$ conidia ml$^{-1}$. Cultures were always grown in triplicates if not otherwise mentioned.

*Schizophyllum commune* and *Ganoderma applanatum* were isolated by Philipp Benz in Freising, Germany. *Dentipellis fragilis* and *Hericium coralloides* were isolated by Markus Blashke in the natural forest reserve Gitschger, Germany. *Pleurotus ostreatus* FPRL 40C, *Trametes versicolor* BAM116 (CTB863) and *Rhodonia placenta* BAM 113 (FPRL 280) were obtained from the German Federal Institute for Materials Research (BAM). *Phanerochaete chrysosporium* (DMSZ 1556) was obtained from the German Collection of Microorganisms and cell Cultures GmbH. Basidiomycete strains were cultivated on yeast malt extract agar (YMEA) containing 10 g L$^{-1}$ glucose, 5 g L$^{-1}$ peptone, 3 g L$^{-1}$ malt extract, 3 g L$^{-1}$ yeast extract and 20 g L$^{-1}$ agar until the mycelium covered the entire plate.

**Phenotype analysis and growth rate on PW agar**
The agar plates used for the phenotype experiments consisted of 2 % agar adjusted to pH 5.0. The PW agar was prepared either without or with the addition of Mandels-Andreotti (MA) medium components and no addition of glucose [40]. MA minimal media agar was prepared with 0.2 g L⁻¹ glucose to simulate the glucose concentrations present in the PW. The plates were inoculated with 2 µl from a spore solution containing 0.5 x 10⁶ conidia ml⁻¹.

After ten days of cultivation at 25 °C with constant light. The conidia were harvested by washing the agar plates with 5 ml H₂O ten times covering one half of the plate and then repeating the procedure with another 5 ml of H₂O for the other half of the plate. Conidia were filtered with glass wool, centrifuged, and finally resuspended in 5 ml H₂O. The quantification was done by measuring the OD at 600 nm.

For the growth analysis of the Basidiomycete strains, 5 mm plugs were cut out of pre-cultures using a coring tool and then transferred to the middle of new plates with YMEA with no PW, with 25 % or 75 % (v/v) PW from Douglas-fir wood chips with bark (Additional file 1; Figure S1). All plates were prepared with 20 mL agar and the plugs were cut from the peripheral growth zone of the fungal cultures. The plates were incubated at 25 °C. The growth was observed daily and recorded with digital photographs using a camera equipped with a 60 mm macro lens (Nikon) next to a reference scale. The growth rate was calculated from the fitted curve of biological triplicates and technical replicates were measured in different directions.

**Press water liquid media for cellulase production**

For the cultivation with PW supernatant, the raw PW was centrifuged at 4000 rcf for 15 min (Heraeus Megafuge 40R, Thermo Scientific) and the supernatant was decanted to be used in the liquid medium. The PW supernatant was diluted with ddH₂O to different concentrations, namely 25 %, 50 %, 75 %, and 100 % (v/v). The complete MA medium was added to each PW condition, so that only ddH₂O was replaced in each condition. All liquid media were adjusted to pH 5.0 with 0.1 M phosphate-citrate buffer and 1 % (w/v) Avicel PH-101 (Sigma-Aldrich) was used as a carbon source and inducer, if not otherwise mentioned.

The sedimented PW solids, after the centrifugation step mentioned above, were used as a carbon source, and compared to Avicel and wood powder obtained from unpressed Douglas-fir wood chips. The PW solids were washed with ddH₂O and dried overnight in a drying oven. To achieve a particle size like that of Avicel (500 µm), we treated the PW solids and the *Douglas-fir* powder in a ball-mill (MM200, Retsch). Each flask with MA medium contained 1 % (w/v) of the respective solid as a carbon source.

The cultivation with PW with replacement of salts and carbon source was carried out with uncentrifuged PW in different concentrations. The percentage of salts, trace elements and Avicel to be replaced were calculated based on the molar concentrations of dissolved ions (Table 1) and the total hydrolysable solids in the PW (Fig. 2e). A PW concentration of 55 % (v/v) was chosen as the minimal concentration
needed to achieve a complete replacement of trace elements. No trace elements were added to the conditions containing more than 55% PW. Only (NH₄)₂SO₄, peptone, urea, and CoCl₂ were not replaced in any condition (Table 2).

**Table 1** Ion concentration in PW and relative to Mandels-Andreotti medium (MA)

|          | PW    | MA     | covered by PW, % |
|----------|-------|--------|------------------|
|          | mg L⁻¹ | mM     | mM               |                   |
| PO₄³⁻    | 16.90 | 0.18   | 14.70            | 1.21              |
| SO₄²⁻    | 38.00 | 0.40   | 11.85            | 3.34              |
| NH₄⁺     | 1.43  | 0.08   | 21.19            | 0.37              |
| NO₃⁻     | < 5   | -      | -                | -                 |
| Na⁺      | 4.28  | 0.19   | -                | -                 |
| Ca²⁺     | 49.20 | 1.23   | 2.72             | 45.13             |
| Mg²⁺     | 14.60 | 0.60   | 1.22             | 49.38             |
| K⁺       | 130.00| 3.33   | 14.70            | 22.62             |
| Trace elements |       |        |                  |                   |
| Fe²⁺     | 20.10 | 0.36   | 0.02             | 2001.60           |
| Mn²⁺     | 4.83  | 0.09   | 0.01             | 874.88            |
| Zn²⁺     | 0.61  | 0.01   | 0.005            | 184.86            |

**Table 2** Mandels-Andreotti medium constituents modified for the cultivation of *T. reesei* with raw PW (% v/v) with replacement of Avicel
| Components       | Unit     | MA media           | PW media          |
|------------------|----------|--------------------|-------------------|
|                  |          | 1x | 50x TE | 25% | 55% | 75% | 100% |
| **Salt solution**|          |    |        |     |     |     |      |
| KH$_2$PO$_4$     | g L$^{-1}$ | 2.00 | 2.00 | 1.89 | 1.75 | 1.66 | 1.55 |
| CaCl$_2$ · 2H$_2$O | g L$^{-1}$ | 0.40 | 0.40 | 0.35 | 0.30 | 0.26 | 0.22 |
| MgSO$_4$ · 7H$_2$O | g L$^{-1}$ | 0.30 | 0.30 | 0.26 | 0.22 | 0.19 | 0.15 |
| **Trace elements**|          |    |        |     |     |     |      |
| FeSO$_4$ · 7H$_2$O | mg L$^{-1}$ | 5.00 | 250 (*350) | 5.00 | -  | -  | -   |
| MnSO$_4$ · H$_2$O | mg L$^{-1}$ | 1.60 | 80 | 1.60 | -  | -  | -   |
| ZnSO$_4$ · 7H$_2$O | mg L$^{-1}$ | 1.40 | 70 | 1.40 | -  | -  | -   |
| **Carbon source**|          |    |        |     |     |     |      |
| PW (v/v) %       |          | -  | 25 | 55 | 75 | 100 |
| Avicel           | g L$^{-1}$ | 10 | 9.57 | 9.05 | 8.7 | 8.27 |

*Iron concentration used in the iron-enriched control medium MA 50x TE + 20x Fe

Cobalt had to be supplemented to the medium, since considerable differences in the cellulase. 50x TE: 50 times the concentration of trace elements (TE) was used as a control.

The addition of free sugars to the cultivation medium was based on the sugar concentration measured in the PW (Additional file: Fig. S1). Glucose (0.061 g L$^{-1}$), fructose (0.150 g L$^{-1}$), galactose (0.022 g L$^{-1}$), arabinose (0.007 g L$^{-1}$), cellobiose (0.003 g L$^{-1}$), xylose (0.003 g L$^{-1}$), concentrations that correspond to 25 % PW, were added to MA medium with 1 % Avicel and autoclaved. MA medium with no sugars and 25 % PW supernatant, both with 1 % Avicel, were used as control.

**Enzymatic assays**

*Endo*-1,4-β-D-glucanase and *endo*-1,4-β-D-xylanase activity assays were carried out according to the protocols (S-ACMC and S-AXBL) of the manufacturer (Megazyme, Ireland), slightly modified and adapted for a 96-well microplate. Each sample was tested in technical duplicate. The mean and the standard deviation are calculated from the biological triplicates and technical duplicates.

Inhibition of the enzymatic reactions in the presence of PW was verified using the same method as described above. As enzyme we used supernatant of a 5-day old *T. reesei* RUT-C30 culture grown with 1 %
Avicel. Each reaction consisted of 10 µl enzyme and increasing concentrations of PW, namely 20 % (v/v) and 60 % (v/v). The reaction volume was adjusted to 25 µl with ddH₂O. These tests were performed in 6 technical replicates for each condition.

**Bio-clarification of PW**

Liquid pre-cultures of *T. versicolor*, *G. applanatum* and *P. chrysosporium*, were made in shaking flasks with 50 mL potato dextrose yeast (PDY) in 250 mL Erlenmeyer flasks until enough biomass was formed. These strains were chosen due to their fast growth in liquid media. The biomass was then homogenized using an Ultra Turrax (IKA Werke GmbH, Germany) and transferred to the PDY media containing 50 % PW or PWB. The cultures were inoculated with 10 % biomass suspension and were cultivated in 100 mL Erlenmeyer flasks with 30 mL medium for 4 days at 28 °C, 100 rpm (50 mm throw). As a control condition PW and PWB flasks without fungal biomass were incubated together with the cultures. The entire volume of the cultures was collected in centrifugation tubes and centrifuged 2 min at 1000 rcf. The supernatants were carefully transferred to cuvettes and the OD was measured at 600 nm to calculate the turbidity change against the control condition.

**Statistical analyses**

Statistical analyses were performed by applying one-way repeated measures analysis of variance (ANOVA) followed by Dunnett’s test using the growth condition without PW as control in OriginPro 2021 (OriginLab Corporation). Significance level of p< 0.05.

**Declarations**

**Authors’ contributions**

JPB, EWH and KR conceived of the study and supervised it. MR, JPB and EWH designed the experiments. MR and RG performed the experiments and acquired the data. MR, RG, EWH and JPB analyzed and interpreted the data. MR drafted the manuscript, which was critically revised by JPB, RG, EWH and KR. All authors read and approved the final manuscript.

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**Competing interests**
The authors declare that they have no competing interests.

**Consent for publication**

Not applicable.

**Ethics approval and consent to participate**

Not applicable.

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**Figures**

**Figure 1**

Schematic diagram of the processes and respective products associated with the mechanical drying of wood chips intended for energy production and the resulting side-product, press water (PW), tested here for its suitability in enzyme production with *T. reesei* RUT-C30 and Bio-clarification with wood-degrading basidiomycete fungi.

**Figure 2**

Microscopic photographs of raw PW (**a**) and PW supernatant (**b**). **c** Volume-weighed particle size distribution of Douglas-fir PW supernatant by laser-diffraction. **d** Concentration of macro solids, micro solids, and dissolved substances in the PW. **e** Mass percentage of the main fractions of PW solids after acid hydrolysis, namely hydrolysate, acid soluble aromatics, and acid insoluble residues. **f** Chromatogram of the Py-GC/MS of PW AIR with the predominant compound classes with exemplary formulas for 2-methoxy-phenol, oleic acid, and campesterol.

**Figure 3**

Spore concentration and phenotype of *T. reesei* RUT-C30 grown on various agars with combination of MA salts (MA), 0.2 g L⁻¹ glucose (Glc), and press water (PW). Conidia were washed from the plates with equal amounts of water and quantified after 10 days, incubated at 25 °C in constant light.

**Figure 4**

Endoglucanase and endoxylanase activities of *T. reesei* RUT-C30 cultivated in shaking flasks at 30 °C and 250 rpm. Avicel (1 % w/v) was used as a carbon source. **a** MA medium supplemented with different PW supernatant concentrations (25, 50, 75, 100 %). **b** MA medium replacing salts (KH₂PO₄, MgSO₄, CaCl₂), trace elements, and Avicel according to the raw PW concentration. MA 50x TE and 50x TE + 20x Fe are controls with increased trace element (TE) concentrations and iron, respectively. No trace elements were supplemented to PW concentrations starting at 55 %. **c** MA medium with 1 % (w/v) Avicel, 1 % (w/v)
Avicel + sugars based on the concentration in 25 % PW, and 25 % PW supplemented with MA medium. Error bars represent the standard deviation (n=3). Significant differences (p>0.05) relative to the control (0 % PW) are indicated by asterisk.

Figure 5

a Endoglucanase and endoxylanase activity of *T. reesei* RUT-C30 cultivated in MA medium with 1 % (w/v) Avicel, Douglas-fir wood powder, and PW solids. Shaking flasks cultivated at 30 °C at 250 rpm. Micrographs at 10x of 5 day old cultures with b Avicel, c ball-milled Douglas-fir wood, d ball-milled PW solids. Error bars represent the standard deviation (n=3).

Figure 6

a Growth rate (by colony diameter) of several basidiomycetes on yeast malt extract agar supplemented with 25 % and 75 % (v/v) press water Douglas-fir with bark (PWB). Cultivation at 25 °C. Growth measured daily over a period of 2 weeks. Growth rate was calculated from the slope of the growth curve. Significance (p<0.05) in relation to the control without PWB. b Turbidity (600 nm) of culture supernatants incubated in shaking flasks 4 days at 28 °C and 100 rpm. Error bars represent the standard deviation of biological replicates (n=3), thus the PW and PWB controls are displayed without error bars.

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