Research Article

Application of ATR-FTIR Spectroscopy to Compare the Cell Materials of Wood Decay Fungi with Wood Mould Fungi

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Wood fungi create vast damage among standing trees and all types of wood materials. The objectives of this study are to (a) characterize the cell materials of two major wood decay fungi (Basidiomycota), namely, \textit{Trametes versicolor} and \textit{Postia placenta}, and (b) compare the cell materials of decay fungi with four wood mould fungi (Ascomycota), namely, \textit{Aureobasidium pullulans}, \textit{Alternaria alternata}, \textit{Cladosporium cladosporioides}, and \textit{Ulocladium atrum}. Fourier transform infrared (FTIR) spectroscopy is used to characterize the microbial cellular materials. The results showed that the IR bands for the fatty acid at $\sim 2900 \text{ cm}^{-1}$ were different for the two-decay-fungi genre. \textit{Postia placenta} shows more absorbance peaks at the fatty acid region. Band ratio indices for amide I and amide II from protein amino acids were higher for the mould fungi (Ascomycota) than the decay fungi (Basidiomycota). Similarly, the band ratio index calculated for the protein end methyl group was found to be higher for the mould fungi than the decay fungi. Mould fungi along with the decay fungi demonstrated a positive correlation ($R^2 = 0.75$) between amide I and amide II indices. The three-component multivariate, principal component analysis showed a strong correlation of amide and protein band indices.

1. Introduction

Wood fungi are broadly classified into two major categories, namely, decay fungi and mould fungi \cite{1, 2}. Decay or deterioration is performed predominantly by brown rot and white rot fungi, while staining or discolouration of wood is performed by mould fungi \cite{3, 4}. Decay and staining reduce the performance of wood material either structurally or aesthetically. Cell structure of fungi has been widely studied for taxonomy and physiology by using various methods other than spectroscopy \cite{5–13}. The fatty acids that are produced in a fungal cell are species specific \cite{14}. Fungal identification libraries of \textit{Phanerochaete chrysosporium}, \textit{P. sordida}, \textit{P. sanguinea}, \textit{Trametes versicolor}, \textit{T. hirsuta}, and \textit{T. pubescens} have been made by extraction and esterification of fungal cellular fatty acids that were characterized by gas chromatography \cite{14}. However, analyzing the chromatographic result, the investigators observed wide variability in the maximum and minimum values. The variation in the amount of deoxyribonucleic acid (DNA) among fungi is often used to characterize species and strains. Attenuated total reflectance (ATR) Fourier transform infrared (FTIR) spectroscopy is a fast surface characterizing tool that can provide chemical information about a substrate up to several micrometres deep. Fungal characterization by FTIR spectroscopy, though not new, is, therefore, an exciting and challenging task as vast information is made available from the microbial structure without the extraction of cell materials \cite{6, 15–17}. The questions that originate from spectroscopic studies are as follows. (i) How to represent the spectroscopic findings? (ii) What information should be used for spectral analysis? (iii) How reliable are the spectroscopic data?

In this study, two naturally abundant wood rot fungi, \textit{Trametes versicolor} and \textit{Postia placenta}, and four common wood mould fungi, \textit{Aureobasidium pullulans}, \textit{Alternaria alternata}, \textit{Cladosporium cladosporioides}, and \textit{Ulocladium}}
atrum, were cultivated. The objectives of this study are to evaluate the applicability of ATR-FTIR spectroscopy to

1. identify cell materials of rot fungi (Basidiomycota);
2. identify cell materials of mould fungi (Ascomycota);
3. quantify spectral information of two rot fungi and four mould fungi.

2. Materials and Methods

2.1. Fungi Culture. The white rot fungus, *Trametes versicolor* (strain CTB 863A), was cultivated on agar plates containing 20 g Fluka malt, 10 g Fluka agar (4%), and 500 mL water. The brown rot fungus, *Postia placenta* (strain FPRL 280), was cultivated on agar plates containing 20 g Bacto malt, 10 g NMD agar (4%, NMD = Norsk Medisinaldepot AS), and 500 mL water. The wood mould fungi, *Aureobasidium pullulans* (strain DSM 2404), *Alternaria alternata* (strain DSM 62010), *Cladosporium cladosporioides* (strain DSM 62121), and *Ulocladium atrum* (strain DSM 63068), were cultivated separately in agar mediums containing 0.5 L of ionized water, 6.25 g Bacto malt, and 10 g NMD agar (NMD = Norsk Medisinaldepot AS). The mould fungi strains were procured from the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ, Germany). Fungi culture was performed at the Norwegian Forest and Landscape Institute, Ås, Norway.

2.2. Fungi Characterization. The white rot fungus, *Trametes versicolor* (strain CTB 863A), was cultivated on agar plates containing 20 g Fluka malt, 10 g Fluka agar (4%), and 500 mL water. The brown rot fungus, *Postia placenta* (strain FPRL 280), was cultivated on agar plates containing 20 g Bacto malt, 10 g NMD agar (NMD = Norsk Medisinaldepot AS). The mould fungi strains were procured from the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ, Germany). Fungi culture was performed at the Norwegian Forest and Landscape Institute, Ås, Norway.

OMNIC software was used for spectral analysis. Spectra with noise-free, sharp, and prominent peaks were selected to interpret the chemical structures, where absorbance ($A'$) of a pure specimen is governed by Beer-Lambert law as $A' = \log_{10}(1/T)$, with $T$ being the transmittance. Sigma Plot software (V. 11, Systat Software, Inc.) was used for plotting of graphs and for performing regression analysis. Scanning electron microscopic (SEM) back scatter electron (BSE) images, 15kV, were obtained on a carbon black tape background.

2.3. Statistical Analysis. Pearson’s correlation coefficient ($r$) was used to measure the strength of association between two variables. Coefficients of determination ($R^2$ ordinary and $R^2$ adjusted for coefficients) were used to indicate the amount of variation in the response variable explained by the independent variable for the linear regression model. Larger $R^2$ indicates more variability which is explained by the linear model. Chi-square distribution was used to ascertain the goodness of fit. Multivariate principal component analysis (PCA) was used to simplify the problem of variable groups by replacing a group of variables with a single new variable. Each principal component represents a linear combination of the original variables. The variance of this variable is the maximum among all possible choices of the axis. MINITAB software version 16.2.1 was used for statistical analysis. Three replicates from specimens of each fungal species were used for FTIR spectra representation and four replicates for each fungal species were used for regression analysis.

3. Results and Discussion

While fungi culture was performed for decay/rot fungi, the spectral findings of mould fungi were imported from a previous research work [18]. Several common IR bands were initially recognized in the FTIR vibrational spectra. Figures 2, 3, and 4 illustrate the FTIR absorbance spectra...
of the two wood rot fungi (Trametes versicolor and Postia placenta) and the four wood mould fungi (Aureobasidium pullulans, Alternaria alternata, Cladosporium cladosporioides, and Ulocladium atrum). The broad peak around ~3400 cm\(^{-1}\) represents the presence of stretching vibrations of hydrogen-bonded O-H (\(v_{\text{str}}\)) that comes from H\(_2\)O or bonded N-H. The amount of hydroxyl radical from the nutrient substrate contributes significantly to the degradation mechanism by rot fungi [19, 20]. Fatty acids show characteristic absorbance at the 2950–2845 cm\(^{-1}\) wavenumber region. The absence of high intensity peaks of fatty acids for Trametes versicolor at the fatty acid region is notable. Absorbances by amide and polysaccharide moieties were observed at wavenumbers ~ 1650–1050 cm\(^{-1}\). Amides show two absorption bands, amide I and amide II, at ~1700–1500 cm\(^{-1}\) due to vibrational coupling between stretching frequencies of C=O and N-H deformation. Amide I has high C=O character from \(\beta\)-sheet of protein, while amide II has lesser C=O character and demonstrates interactions between C=O and N-H deformation coupling. Phospholipids and PO\(_2\) show bands at 1237 cm\(^{-1}\) and 1082 cm\(^{-1}\) for asymmetric stretching and symmetric stretching vibrations, respectively [21]. Figure 3 for the two wood decay fungi and the four wood mould fungi shows the presence of aliphatic C-H \(v_{\text{str}}\) from esters or triglycerides at 2958 cm\(^{-1}\), from fatty acids, phospholipids, glycerol, and triglycerides at 2921 cm\(^{-1}\), 2915 cm\(^{-1}\), and 2849 cm\(^{-1}\), and from shoulders at 2872 cm\(^{-1}\) and 2852 cm\(^{-1}\), respectively.

3.1. Mould Fungi Figure 4 shows that, at lower wavenumbers, mould fungi cell lipids show band at 1456 cm\(^{-1}\) for CH\(_2\) bending mode and band at 1402 cm\(^{-1}\) for C=O symmetric stretching of COO groups as observed in previous research [18]. In general, aromatic and alkene C-H \(v_{\text{str}}\) bands appear slightly above ~3000 cm\(^{-1}\), while alkane C-H \(v_{\text{str}}\) appears below ~3000 cm\(^{-1}\). Glucans, chitins, amines, and acids are common fungal cell wall components [14, 22–27].

The cell carbohydrates of the four wood mould fungi (Aureobasidium pullulans, Alternaria alternata, Cladosporium cladosporioides, and Ulocladium atrum) show bands...
due to vibration modes at 1064 cm\(^{-1}\) for CH\(_2\)OH and C-O stretching vibration coupled with the C-O bending mode, 1773 cm\(^{-1}\) for C=O carboxyl stretching vibration, 1713–1711 cm\(^{-1}\) for carboxyl C=O, 1662–1659 cm\(^{-1}\) for amide C=O [28], 1592 cm\(^{-1}\) and 1615 cm\(^{-1}\) for asymmetric COO\(^{-}\), 1546 cm\(^{-1}\) and 1552 cm\(^{-1}\) for NH bending deformation, 1502 cm\(^{-1}\) and 1516 cm\(^{-1}\) for C=C aromatic, 1454 cm\(^{-1}\) and 1466 cm\(^{-1}\) for CH\(_2\) bending deformation of lipid and polysaccharide, 883–892 cm\(^{-1}\) for \(\beta\)-chitin and \(\gamma\)-CHx vibration at 962 cm\(^{-1}\) for \(\alpha\)-chitin, and 750–700 cm\(^{-1}\) for C-H out-of-plane bending vibration [18]. Alternaria alternata and Aureobasidium pullulans show two distinct peaks at 1733 cm\(^{-1}\) and 1717 cm\(^{-1}\), while Ulocladium atrum shows a broad peak at higher wavenumber of 1740–1733 cm\(^{-1}\) [18].

### 3.2. Decay Fungi.

Absorbance at \(\sim 2200–1900\) cm\(^{-1}\) is overshadowed by the interference from the diamond crystal absorption and spectral noise. Subsequently, there is no possibility to identify the presence of cumulative double bonds, antisymmetric C=C \(\nu_{as}\), at 2000 cm\(^{-1}\) (Figure 4). However, there is a sharp peak at 2323 cm\(^{-1}\) for both decay and mould fungi that could possibly have originated from ketenes, O=C=O. The absence of any strong band at \(\sim 2250\) cm\(^{-1}\) eliminates the probability of presence of nitrile group, characterized by C≡N \(\nu_{as}\) absorption. Trimetres versicolor shows higher intensities at the \(\sim 3400\) cm\(^{-1}\) region. Bonded amines, \(N\-H\ \nu_{as}\), display characteristic absorption at \(\sim 3300\) cm\(^{-1}\) region and \(\equiv C\-H\ \nu_{as}\) shows characteristic absorption band at \(\sim 3320\) cm\(^{-1}\). The band at \(\sim 2850\) cm\(^{-1}\) is the characteristic absorption band for symmetric stretching of CH\(_2\). C-H \(\nu_{as}\) absorbance peaks are more prominent for Postia placenta compared to Trimetres versicolor. The replicates of spectra from Trimetres versicolor yield broad bands at 2921 cm\(^{-1}\) and 2852 cm\(^{-1}\). The replicates of spectra from Postia placenta have strong absorption bands at 2958 cm\(^{-1}\), 2915 cm\(^{-1}\), 2872 cm\(^{-1}\), and 2849 cm\(^{-1}\). The absence of the band at 2872 cm\(^{-1}\) in the spectra of Trimetes versicolor indicates that possibly Postia placenta has higher numbers of alkyl and alkane chain numbers in fatty acid configuration.

The fingerprint region for the wood rot decay fungi, Trimetes versicolor and Postia placenta, is shown in Figure 4. The spectra for both fungi are largely similar in pattern except some more peaks for Postia placenta. Small shoulder at 1735 cm\(^{-1}\) of the spectrum originated from Trimetes versicolor shows the presence of C=O \(\nu_{as}\) band for uronic acids or from saturated esters. The overtone of C=O \(\nu_{as}\) band is not distinguishable at 3400 cm\(^{-1}\). A peak at 1717 cm\(^{-1}\) from both of the fungi and a small shoulder at 1700 cm\(^{-1}\) from Postia placenta suggest possible presence of carboxyl group in conjunction with a double bond or Glu (C=O) or Asp (C=O) amino acids [29]. The possibility of presence of the C=O group is further supported by a strong peak in the Postia placenta spectrum at 1683 cm\(^{-1}\) which could have resulted due to chelation between C=O group and a neighbouring O-H group. The presence of O-H along with C=O \(\nu_{as}\) is also evidenced by O-H deformation at 932 cm\(^{-1}\) found in the spectrum of both decay and mould fungi. Henceforth, these bands have originated from carboxyl groups present in the cell structures of the fungi. Two shoulders at 1654 cm\(^{-1}\) and 1652 cm\(^{-1}\) are probable peaks from characteristic absorptions of zwiterions -N\(^+\)-H \(\nu_{as}\) and -CO\(_2\) \(\nu_{as}\) from asymmetric stretching of amino acids, Arg (CN,HN), and acetamides from chitin [28]. A strong absorption at \(\sim 1556\) cm\(^{-1}\) is due to amide II from \(\alpha\)-helix of protein, Glu (COO\(^{-}\)), and/or chitin. The band at 1505 cm\(^{-1}\) shows the presence of \(\alpha\/-\mu\)-substituted aromatic units, indole groups of amino acids (CN, CH, and NH), that is, Trp amino acid. The band at 1496 cm\(^{-1}\) is assigned to aromatic (CC) ring, Tyr (O\(^\circ\), CC, CH), or Trp (CC, CH) amino acid [29]. The small band at 1456 cm\(^{-1}\) for asymmetric CH\(_2\) bending modes of end ethyl groups of protein and 1419 cm\(^{-1}\) could be due to absorption from carboxymethyl COO\(^{-}\) units [29]. A strong band at 1375 cm\(^{-1}\) is due to chitin. The strong band at 1317 cm\(^{-1}\) can be assigned to amide III from proteins and chitins, 1248 cm\(^{-1}\) for stretching of non-hydrogen-bonded PO\(_2\) group of phosphodiester groups of nucleic acid, 1202 cm\(^{-1}\) for P-O stretch, and 1148 cm\(^{-1}\) for \(\alpha\)-Glucans. Bands at 1146 cm\(^{-1}\) are due to CO/C/CC from \(\alpha\)-Glucan complexes, a shoulder at 1075 cm\(^{-1}\) is due to CO/CC from \(\beta\)-Glucan complexes, the band at 1023 cm\(^{-1}\) is for \(\alpha\)-Glucan complexes or cellulose-OH, the band at 930 cm\(^{-1}\) is for aromatic rings, and finally the bands at 888 cm\(^{-1}\) and 848 cm\(^{-1}\) represent C-1-H of \(\beta\)-Glucan and C-1-H of \(\alpha\)-Glucan units, respectively. Absorbance below 800 cm\(^{-1}\) could be due to the overtones of bands at higher frequencies.

There is an additional peak at 1683 cm\(^{-1}\) for Postia placenta. The observed differences between Trimetes versicolor and Postia placenta in Figures 3 and 4 have resulted due to the presence and/or absence of different proportions of hydrocarbons and fatty acids. Presence of the acid groups would have resulted in sharp peaks, while absence would have resulted in no absorbance. Extracellular OH groups are present in Trimetes versicolor [30]. Moreover, it is known that Postia placenta secretes oxalic acid into the sheath environment [19, 31]. Hence, the sharp peak at 1683 cm\(^{-1}\) in Figure 4 is probably due to the IR detection of carboxyl groups.

Overall, as observed by previous researchers [6, 32], it can be concluded that there is strong evidence of presence of chitin, Glucans, and glycoproteins that are covalently linked together in a dynamic process that could occur extracellularly or intercellularly. The probing depth of ATR-FTIR spectroscopy was enough to sense bond-vibration signal of these active groups.

### 3.3. Quantitative Spectral Analysis for Six Fungal Species

Based on the observations in Figures 2, 3, and 4, the ATR-FTIR band ratio indices at 1650 cm\(^{-1}\), 1546 cm\(^{-1}\), and 1456 cm\(^{-1}\) wavenumbers in the IR spectral region for the six fungal genre were calculated [18] in the following way:

\[
\text{Band ratio index} = \frac{I_a}{I_{2921}}, \quad (1)
\]

where \(I_a\) is the band intensity at the observed wavenumber “a” cm\(^{-1}\) and \(I_{2921}\) is the band intensity at wavenumber
The band ratio index calculated at 1456 cm\(^{-1}\) and at 1650 cm\(^{-1}\) for the decay fungi and the mould fungi is not linear as is observed in Figure 5. Observing the pattern for the fungi belonging to Ascomycota and Basidiomycota, it may be concluded that the amide I and amide II indices do not overlap. Figure 5 demonstrates that the band ratio index at 1456 cm\(^{-1}\) has higher values for decay fungi and the mould fungi is not similar to wood mould fungi resulting in \(R^2 < 1.00\). Previous research on bacterial cell component has reported similar positive correlation with the normalised intensities of amide II/C-H bands and the N/C elemental surface concentration ratio of the strains determined by X-ray photoelectron spectroscopy (XPS) [33]. The authors conferred that there is higher nitrogen content on the bacteria as compared to yeasts. Additionally, quantification of the peak areas at 1510 cm\(^{-1}\), 1138 cm\(^{-1}\), and 1758 cm\(^{-1}\) to follow degradation changes in wood substrate by ATR-FTIR spectroscopy showed increasing amount in lignin/carbohydrate ratio with degradation period [34–36].

Table 1 shows the descriptive statistics for each variable considered in Figure 5. Assuming normal distribution, Chi-square test was performed with hypothesized standard deviation \(\sigma = 0.5\). From the \(p\) value \(\{\text{probability}(X > \text{test statistic})\}\) in Table 1, it can be concluded with 95% confidence limit that the band index calculated at \(I_{1456}/I_{2921}\) does not follow a multinomial distribution with specific proportion.

An IR spectrum contains more information than is needed. The entire spectrum is used to obtain absorbance values of a few peaks of interest. However, by using peak indices as variables \(I_{1650}/I_{2921}, I_{1546}/I_{2921},\) and \(I_{1456}/I_{2921}\) in principal component analysis (PCA) technique, redundant data that are not useful can be reduced. A principal component analysis (PCA), assuming normal distribution, shows (Table 1) that the band index calculated at \(I_{1546}/I_{2921}\) has the largest percentage of the total data variation, followed by index at \(I_{1456}/I_{2921}\) and \(I_{1650}/I_{2921}\). The larger correlations are in bold fonts in Table 1. The first principal component of a variable has the largest sample variance amongst all normalized linear combinations of the columns. Subsequent principal components have maximum variance subject to being orthogonal to the earlier ones. The first principal component (PC1) correlates most strongly with \(I_{1456}/I_{2921}\). Moreover, all the 3 variables show correlation with a value above 0.5, suggesting that these three band indices vary together to certain extent for the first component. The second principal component (PC2) correlates strongly with \(I_{1456}/I_{2921}\) and has negative correlation with \(I_{1650}/I_{2921}\). PC2 can be viewed as the measure of \(I_{1456}/I_{2921}\) and lack of \(I_{1650}/I_{2921}\). The third principal component (PC3) has the strongest negative correlation with \(I_{1546}/I_{2921}\) and positive correlation with \(I_{1650}/I_{2921}\). Henceforth, PC3 can be viewed as the measure of \(I_{1546}/I_{2921}\) with lack of \(I_{1546}/I_{2921}\). Therefore, a regression model in future with IR band ratio index would exclude the component that has the largest variation.
Table 1: Summary of the Chi-square test and principal component analysis (strong interactions are in bold letters).

| Variable (n = 6) | Standard deviation | Chi-square | Principal component analysis |
|-----------------|--------------------|------------|-----------------------------|
|                 |                    | 95% confidence interval | Test statistic | p value | PC1      | PC2      | PC3      |
| \(I_{1650}/I_{2921}\) | 0.670              | (0.418, 1.643)         | 8.98           | 0.220   | 0.563    | −0.546   | 0.621    |
| \(I_{1546}/I_{2921}\) | 0.576              | (0.359, 1.412)         | 6.63           | 0.500   | 0.528    | −0.341   | −0.778   |
| \(I_{1456}/I_{2921}\) | 0.811              | (0.506, 1.989)         | 13.15          | 0.044   | 0.636    | 0.765    | 0.096    |

Figure 6: SEM image of fungal spores and hypha.

Figure 6 shows the scanning electron microscopic image of Basidiomycota fungal spores on a carbon black tape background. Researches have shown that there could be crystal depositions on the hypha depending on the wood substrate [19]. The small dimensions of the spores and hyphae from Figure 6 suggest that the ATR-FTIR signal from a single beam spot on fungal surface could not be singly due to the hypha; however, it could be from both the hyphae and amino acid rich spores.

4. Conclusions

Fourier transform infrared (FTIR) spectroscopy is used to compare cellular materials of wood decay and wood mould fungi. Infrared (IR) bands originating from six fungal species showed marked differences. Specifically, it was observed that

(1) chitin and fatty acid were present in all six fungi;

(2) spectra from *Trametes versicolor* (Basidiomycota) yield broad bands at 2921 cm\(^{-1}\) and 2852 cm\(^{-1}\). Contrarily, the spectra from *Postia placenta* (Basidiomycota) have strong absorption bands at 2958 cm\(^{-1}\), 2915 cm\(^{-1}\), 2872 cm\(^{-1}\), and 2849 cm\(^{-1}\);

(3) Ascomycota and Basidiomycota amide I and amide II indices (for indices at 1650 cm\(^{-1}\) and 1546 cm\(^{-1}\)) do not overlap. The calculated amide II index at 1546 cm\(^{-1}\) is less for the two decay (Basidiomycota) fungi than for the four mould (Ascomycota) fungi;

(4) multivariate analysis provides Pearson’s correlation coefficient, \(r = 0.8\), for amide I and amide II (for indices at 1650 cm\(^{-1}\) and 1546 cm\(^{-1}\)) indicating a positive linear correlation;

(5) additionally, 75% of the variation could be explained in the linear relationship between band indices of amide I at 1650 cm\(^{-1}\) and amide II at 1546 cm\(^{-1}\) (\(R^2 = 0.75\));

(6) the protein ethyl index at 1456 cm\(^{-1}\) shows a decreasing trend for the four mould fungi *Cladosporium cladosporioides > Alternaria alternata > Aureobasidium pullulans > Ulocladium atrum*;

(7) principal component analysis was found to be useful to define the nature of correlation between the bands indices calculated from fingerprint regions. It was observed that the band index calculated at \(I_{1546}/I_{2921}\) has the largest percentage of the total data variation.

In conclusion, this study shows that ATR-FTIR spectroscopy provides information about fungal cellular structure that is specific to genus and species. Inter- and intraspecies comparison by FTIR spectroscopy shows the potential to generate future databanks to identify various fungi and pathogens.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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