Research Article

Analysis of Selected Properties of Fibreboard Panels Manufactured from Wood and Leather Using the Near Infrared Spectroscopy

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1. Introduction

The scarcity of resources can be a motor or motivation for innovation. The application of unused biogenic resources is one way where innovation can happen. With these new raw materials the product characteristics can change; thus material analysis and material test are necessary to determine the composite properties. Also guidelines for quality assurance of the new materials can aid in the transfer from laboratory to industrial conditions for consumer applications.

There is a lot of innovation regarding the use of by-product from biogenic resources in the forest product field [1]. Most of these investigations were dealing with an up- or recycling of by-products to find possible supplements to the fibres and particles for wood based panel production [2, 3]. Han et al. [4] and Halvarsson et al. [5, 6] produced fibreboards from wheat and reed remains in combination with urea formaldehyde (UF) and urea melamine formaldehyde (UMF) adhesives. Also, exotic materials such as coconut fibres [7, 8], steam exploded banana bunch fibres [9], and bamboo and rice straw [10] were investigated mechanically as well as physically and obtained meaningful insights into possible alternatives to conventionally produced fibreboards and particleboards. Kargarfard et al. [11] investigated encouraging agro-based materials as cotton and corn stalks in MDF. Lee [12] combined bagasse with other biobased materials, as bamboo, in particleboards and analysed their mechanical and physical properties.

Besides the biobased materials also the fossil-based materials, such as plastic waste [2], were examined in combination with wood particles and fibres, and its degradability was determined. A seminal way to upcycling by-products seems
to be the usage of waste materials, such as chicken feathers and leather shavings, which occur in the meat industry [13, 14]. Also, in the leather and tannery industry, a huge amount of waste incurs every year, alone in Europe more than 200,000 tons a year [15]. This waste is then disposed together with other urban wastes [16].

To produce a wood-based panel out of a mixture of wood fibres and leather particles is a quite new idea, which was patented by Lackinger [17] in the year 2009. Investigations by Rindler et al. [18], Solt et al. [19, 20], and Wieland et al. [21] describe the mechanical and physical properties of leather shavings in MDF. A multitude of fibreboards with different percentages of leather shavings and wood fibres were tested to analyse the influence on the leather of the mechanical properties (e.g., internal bond). The behaviour can be explained with the structural and the chemical properties of the various materials. Differences in functional groups of leather, wood, and adhesion were analysed by $^{13}$C-NMR [22], Raman spectroscopy [23], and FT-IR spectroscopy [24]. Moreover, the NIR spectroscopy in combination with multivariate data analysis was applied in the wood industry for a quality assurance control system [25]. Also detailed information about the application of NIR spectroscopy in wood and paper research is given by Tsuchikawa [26].

Therefore, based on their mechanical and technical properties the wood leather fibre composites are one of the most interesting wood engineered materials of the last years. These composite materials are highly sustainable because they can be produced by coupling wood fibres with industrial waste of tannery plants. The analysis of the wood leather fibreboard with the near infrared spectroscopy (NIRS) can provide a basis for further efforts in the upscaling from the laboratory to industrial conditions for consumer application concerning this tool for the development of quality assurance control system.

2. Experimental

2.1. Material. When leather gets produced, hides have to run through different production steps. After the withdrawal of the skin a preservation process has to be done to protect the freshly peeled skins against the influence of microorganisms. The next step, the tanning process, is used to protect the skin against enzymatic degradation and increase their resilience. Only after this production step the skins are called leather. For this investigation the leather types, wet blue and wet white, were used (Figure 1).

The leather particles accrue during the shaving process of hides preparation, where they got sliced to a specific thickness. These particles were dried to moisture content (m.c.) of $8\pm1\%$ in a small size dry kiln (Brunner-Hildebrand) of Salzburg University of Applied Sciences at Campus Kuchl at a temperature of 40°C.

Norway spruce wood fibres ($Picea abies$ (L.) [Karst.]) were used for this study. The fibres were produced in an industrial facility for MDF production and were unglued with an m.c. of $8\pm1\%$. 

![Figure 1: Raw materials to produce the composite materials: (a) wood fibres, (b) leather shavings wet blue, and (c) leather shavings wet white.](image-url)
2.2. Method

2.2.1. Fibreboard Manufacturing. The spruce fibres and wet white leather particles were glued based on the oven-dry density with 10% urea formaldehyde (UF) resin with 1% ammonium sulphate solution in a lab ENT WBH 75 ploughshare blender type with a Schlick two-substance nozzle up section. For the process a nozzle with a whole diameter of 2.3 mm and a pneumatic pressure of 2 bar was used. Further the glued fibres were distributed manually in a frame and were pressed to a final thickness of 8 to 20 mm in a Hoefer HLPO 280 automated hot lab press at 80°C with a pressing factor of 1 min/mm. Fibreboards with dimensions of 450 x 450 mm² with different thicknesses were produced under laboratory conditions. The ratio of different leathers to the wood fibres and the various thickness of fibreboard samples were selected from the results of various previous mechanical studies by Solt et al. [18, 19]. After the pressing process the samples were stored in a standard climate (20°C/65% r.H).

2.2.2. FT-NIR Spectroscopy. For FT-NIR measurements each raw material (e.g., leather particles and wood fibres) was milled with a cutting mill (Retsch) using solid CO₂ to pass a mesh of 500 μm and the fractions between 250 and 63 μm were separated with a sieving apparatus (Retsch). Then the powder was dried at 50°C for one week.

The FT-NIR spectra were obtained on the surface of wood leather fibreboards and on the milled fibres of each sample by an MPA spectrometer (Buker) equipped with a fibre probe (4 mm measurement diameter) at a resolution of 8 cm⁻¹ (32 scans). For every wood leather fibreboard and milled fibres five single spectra of the surface per sample were taken to minimize the influence of different concentrations of wood and leather shavings on various wood leather panels.

2.2.3. Mechanical Testing Methods. The sample preparation and the mechanical testing procedure for the modulus of rupture (MOR) and elasticity (MOE) were done according to the OENORM EN 326-1 [27] and the OENORM EN 310 [28]. To obtain meaningful results, each of the mechanical tests had an amount of 5 samples.

2.2.4. Data Analysis. The Unscrambler 10.3 software (CAMO, Norway) was used for the data analysis. The FT-NIR spectra were managed without data treatment and also were pretreated by using the second derivative (15 smoothing points). Principal Component Analysis (PCA) is a linear projection method to reduce the multidimensional data (e.g., NIR spectra) to only few orthogonal features (principal components (PCs)). The Partial Least Squares Regression (PLSR) method was applied to find the latent variables in X (e.g., NIR spectra) that would best describe the variables in Y (e.g., bending strength). Esbensen [29] and Kessler [30] give detailed information about the PLSR method. On one hand the NIR spectra were not pretreated. On the other hand the NIR spectra were pretreated by using the second derivative (15 smoothing points). The NIR data sets of various wood leather panels were regressed versus different physical and mechanical properties of the cross-validation (root mean square error of cross-validation (RMSECV)) of the models.

3. Results and Discussion

The chemical information relating to the two different milled leather powders and the milled wood powder was obtained by using the FT-NIR spectroscopy. Figure 2 shows the spectra in the region between the wave number range 9000–4000 cm⁻¹ of the wood fibres and the wet white (ww) and the wet blue (wb) leather fibres. The differences between the wood fibres spectrum and the leather fibres spectra can be observed. The bands around the wave numbers 6660, 4886, and 4587 cm⁻¹ corresponded with the structure of proteins, here especially the first overtone of N-H stretching, the single or combination of amide I or amide II, and the second overtone of N-H bending vibrations [31]. Also the second overtone of O-H bending vibrations at band around 5141 cm⁻¹ [31] is different compared to the wood fibres spectrum as this spectrum shows the first overtone O-H stretching vibrations at the wave number 5192 cm⁻¹ [32]. Furthermore, the spectrum of the wood fibres shows a significant difference at the band around 4751 cm⁻¹, which corresponded to the third overtone of the asymmetric C-O-O stretching and the O-H bending as well as the C-O stretching vibrations [31, 32]. The bands at around 5777 and 4373 cm⁻¹ corresponded to the first overtone of C-H stretching and the second overtone of C-H bending as well as CH₃ deformation vibrations [31].

These results show that the NIR spectroscopy is suitable for characterizing different materials of wood and leather, which was also depicted by the FT-IR and Raman Spectroscopy [23, 24]. However, the two types of leather fibres cannot be distinguished by the NIR spectra analysis.

Also, the NIR spectra of the various wood leather panels show differences in IR bands (Figure 3). The significant bands are around the wave numbers 4886 and 4587 cm⁻¹ for the leather particles and the wave number 4751 cm⁻¹ for the wood fibres. With the increase of the amount of ww leather shavings the changes in these wave numbers can be observed. Further analysis should be done to show the potential of NIRS for the classification of the leather and wood fibre amount in
Figure 3: FT-NIR spectra of wood leather fibreboards with various concentrations of wood and wet white (ww) leather particles in the wave number range between 9000 and 4000 cm\(^{-1}\).

Figure 4: Principal component (PC) analysis score plot of near infrared spectra of various wood fibre leather panels.

Figure 5: Loadings of the first two principal components (PCs) of near infrared spectra of various wood fibre leather panels.

The results of the physical and mechanical properties of various wood leather fibreboards are shown in Table 1. With increasing of the leather amount the values of mechanical properties of the wood leather boards were decreased. This phenomenon can be determined for various thicknesses of the wood leather fibreboards. All these results are consistent with the detailed analysis of mechanical properties by Solt et al. [19].

For the PLSR models the NIR data were pretreated by using the second derivative (15 smoothing points). A full cross-validation was performed for every sample. For wood leather fibreboard samples \((n = 18)\), the coefficient of determination \((R^2)\) was 77.35% and the RMSECV was 50.3% with two principle components (PCs). PLSR models for the bending strength (MOR) of the fibreboard were also calculated with 18 samples, resulting in \(R^2 = 93.55\%\) and RMSECV = 1.7% with two PCs. Figure 6 shows the measured versus the predicted density of various wood leather fibreboards. The sample amount was small and varied for the different material properties. However, with these results it seems that the FT-NIR spectroscopy is able to estimate the physical and mechanical properties (e.g., bending strength).

The relationship between the mechanical properties and the FT-NIR spectra demands further consideration. The physical and mechanical features of the wood fibre leather composite samples depend on the leather content of the fibreboards. The geometric form of leather particles is not to be compared with wood fibres. Therefore, the distribution of the leather particles is not homogeneous [23]. There are some areas with bigger leather accumulations. Simultaneously, the leather particles can fill in the void of the fibreboard [21]. These phenomena have been taken into account by the use of 4 mm measurement area of the NIR spectrometer. With changing leather contents the material properties of the fibreboards composite are also modified. Therefore, the leather particles are not only an additive of the wood based boards and can be used for new material resources for wood based panel industry.
Table 1: Estimated mean and standard deviation (SD) of the physical and mechanical properties of the wood leather fibreboards with different leather types (ww and wb) and various ratios of wood fibre and leather.

| Composition of the panel | Thickness (mm) | Density (kg/m³) | MOE* (N/mm²) | MOR# (N/mm²) |
|--------------------------|----------------|-----------------|--------------|--------------|
| 66.6 33.3 — — | 8 | 807 (49.9) | 1977.53 (148.30) | 23.32 (1.66) |
| 66.6 33.3 — — | 12 | 854 (31.4) | 1779.72 (120.77) | 17.11 (1.79) |
| 66.6 33.3 — — | 16 | 767 (39.1) | 1721.00 (156.89) | 17.14 (2.03) |
| 66.6 66.6 — — | 8 | 829 (27.7) | 1178.70 (95.74) | 6.35 (0.94) |
| 33.3 66.6 — — | 12 | 935 (59.1) | 1099.31 (237.85) | 9.72 (2.01) |
| — — 100 — | 20 | 725 (34.5) | 938.00 (208.01) | 10.30 (1.98) |
| 66.6 — 33.3 — | 12 | 772 (90.0) | 1513.00 (349.01) | 14.57 (3.42) |
| 66.6 — 33.3 — | 16 | 703 (60.5) | 1577.00 (98.86) | 11.80 (1.82) |
| 66.6 — 33.3 — | 20 | 694 (77.7) | 1488.00 (189.81) | 10.67 (1.93) |
| 33.3 — 66.6 — | 12 | 828 (30.4) | 1142.00 (277.49) | 12.59 (2.86) |
| 33.3 — 66.6 — | 16 | 777 (79.6) | 1122.00 (251.38) | 9.72 (2.01) |

1%10% UF adhesive content in every fibreboard.
* Modulus of elasticity (standard deviation).
# Modulus of rupture/bending strength (standard deviation).

4. Conclusions

Based on these results of this study, NIR spectroscopy can be used to distinguish wood fibres from different types of leather shavings (wet white and wet blue). These differences can be also observed of the wood leather fibreboard samples. The score plot from the PCA depicts a possible classification into various amounts of leather contents of the fibreboards. Moreover, the PLSR models for the prediction of the physical and mechanical properties were successfully developed. These results demonstrate that a classification of fibreboard composite and the estimation of density as well as bending strength features are possible with fast, nondestructive measurement methods. The methods may serve a basis to establish guidelines for quality assurance control systems of this new engineered wood leather composite fibreboard. These findings provide a basis for further efforts in the upscale from laboratory to industrial conditions for consumer applications.

Conflict of Interests

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

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