EFFECT OF THERMAL TREATMENT ON PHYSICAL AND MECHANICAL PROPERTIES OF BIRCH AND PINE WOOD

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Abstract
No simple method has yet been found for satisfactory wood bio-resistance improvement regarding material performance in its end-use. An attempt to obtain material with proper strength and bio-durability by combined wood thermal modification and impregnation with a biocide is being researched. To select the most appropriate treatment conditions for the combined process, changes in wood physical and mechanical properties depending on the treatment temperature were investigated in the present study. For the investigation, in Latvia the most widespread wood – softwood pine (Pinus sylvestris L.) and hardwood birch (Betula spp.) was used. Changes of wood mechanical and physical properties due to thermal modification were investigated and effect of treatment temperature and relative humidity on wood characteristics evaluated. It was found that, due to different degree of changes, no identical treatment conditions suit for birch and pine wood. Birch wood is considerably more sensitive to temperature and acceptable strength was maintained only for birch wood treated at 150 °C and for pine wood treated at 160 °C. Nevertheless at higher environmental humidity equilibrium moisture content and consequently radial and tangential swelling increased for all studied wood types, substantially smaller changes due to elevated humidity were detected for modified wood.

Key words: thermal treatment, birch, pine, hardness, bending strength, swelling, capillary water uptake.

Introduction
Over recent decades, different treatment methods have been proposed for restriction of such wood drawbacks as dimensional instability and low resistance to biodegradation (Homan & Jorissen, 2004; Gerardin, 2016). Commercially the most successful has been wood thermal modification and different treatment processes have been introduced in production on industrial scale (Hill, 2006). One advantage of wood thermal modification in comparison with other wood modification methods is that no chemicals are used to alter wood properties as the changes are caused by autocatalytic reactions of wood chemical components during its exposure to elevated temperature. This makes wood thermal modification relatively environmentally friendly (Sandberd, Haller, & Navi, 2013). Another advantage is that the equipment is simple and comparatively low capital expenditure is needed to launch manufacturing. Therefore, it is foreseen that thermal modification of wood will continue growing (Militz & Altgen, 2014).

A lot of different wood thermal modification methods are known which vary with the treatment atmosphere, temperature, and duration (Hofmann et al., 2013; Militz & Altgen, 2014; Gerardin, 2016). They all include exposure of wood to elevated temperature (150 – 260 °C) in absence of oxygen which result in complex reactions including certain destruction of low-molecular substances and hemicelluloses, reorganisation of lignin and cellulose and evaporation of volatile compounds (Sandberd, Haller, & Navi, 2013). These wood chemical transformations result in changed wood colour, improved dimensional stability and enhanced bio-durability. However, alteration of wood chemical composition and structure due to thermal treatment causes reduction in wood mechanical strength (Boonstra et al., 2007; Arnold, 2010; Welzbacher et al., 2011; Widmann, Fernandez-Cabo, & Steiger, 2012). Therefore, when selecting the modification method and processing parameters, it is important to find a trade-off between benefits and losses taking into consideration the requirements for the material end-use.

Despite numerous investigations, no simple method has been found yet for fully satisfactory result regarding material potential performance in its end-use. Thus, different combined treatment processes, including pre- or post-treatment of thermally modified wood, are now intensively investigated and the results are being reported (Ahmed, Hansson, & Moren, 2013; Ferrari et al., 2013; Wang, Zhu, & Cao, 2013). The present research is a part of the investigation aimed at improving wood bio-durability by combined wood thermal treatment and impregnation with a commercial biocide. It is found that substantial enhancement of bio-durability can only be reached by wood thermal treatment at the temperature range at which wood mechanical strength significantly decreases consequently restricting the application area of the material (Kamdem, Pizzi, & Jermannaud, 2002; Metsä-Korteleinen & Viitanen, 2010; Candelier et al., 2017). It is expected that by applying the proposed combined treatment process a material with decent mechanical strength and meeting the requirements of the use class 3 according to the EN 335-1 standard will be obtained.
The objective of the present study was to evaluate changes in wood physical and mechanical properties due to thermal modification depending on the treatment temperature. It will let select the treatment conditions for the combined process. For the investigation, in Latvia the most widespread wood - softwood pine (Pinus sylvestris L.) and hardwood birch (Betula spp.) was used.

Materials and Methods

For the thermal modification, kiln-dried boards of birch (Betula spp.) and pine (Pinus sylvestris L.) wood measuring 700 x 100 x 25 mm were used. The modification was carried out in a multifunctional wood modification device of WTT (Denmark) production. The boards were thermally modified in a water vapor medium under elevated pressure (0.6 – 0.8 MPa depending on the temperature) for 1 h at the peak temperature. Each of the wood species was treated at three peak temperatures: birch wood at 150, 160 and 170 °C and pine wood at 160, 170 and 180 °C. The boards were weighed before and after modification and mass losses calculated. The modified boards were conditioned (RH 65 ± 5%; 20 ± 2 °C) for at least two weeks before preparing specimens for testing mechanical and physical properties.

Determination of wood equilibrium moisture content (EMC) and wood radial and tangential swelling was performed with specimens measuring 20 x 20 x 10 mm (r x t x l) and having the annual ring orientation strictly parallel to the edge. Ten replicates were used per each modification temperature and the untreated wood. Before starting the test, specimens were oven-dried (102 ± 2 °C) and their mass (with accuracy of 0.0002 g) and dimensions (with accuracy of 0.02 mm) were determined. Further the specimens were conditioned until reaching constant weight at 20 ± 2 °C temperature and fixed relative humidity (RH) conditions in increasing sequence: 45%, 65%, 85%. The specimen equilibrium mass and dimensions were recorded for each tested RH and calculations according to DIN 520184 were performed to establish wood swelling characteristics. The moisture exclusion efficiency (MEE) was calculated as given below (Eq. 1).

\[ \text{MEE} = \frac{\text{EMC}_{\text{mod}} - \text{EMC}_{\text{mod}}}{\text{EMC}_{\text{mod}}} \times 100\% \]  

Capillary water uptake (CWU) through radial and tangential surfaces was tested using cubic specimens (20 x 20 x 20 mm) with annual ring and grain orientations strictly parallel to the edges. All sides, except two opposite faces (radial or tangential), one of which was intended for CWU evaluation, were sealed with waterproof coating. After conditioning (RH 65 ± 5%; 20 ± 2 °C) until constant weight, the specimens were installed into a frame that restricted water evaporation from the container and fixed the specimens in a position in which the contact surface was 2 ± 0.2 mm under the water. The container was filled with distilled water the level of which was monitored and adjusted every day. After 10 days, the specimens were removed from the water, excess water wiped off with a paper towel and the specimen weight recorded with an accuracy of 0.0002 g. The experiment was carried out in a room with controlled RH (65 ± 2%) at 20 ± 2 °C. Ten replicates were used per each wood type.

Wood bending strength was determined according to DIN 52186 in a three-point bending test using a material strength testing device ZWICK Z100. Before the test was carried out, the specimens measuring 360 x 20 x 20 mm with the fibres parallel to the sample longitudinal axis were conditioned at two different RH (65% and 85%) till constant mass. At least 30 specimens obtained from different boards were tested for each modification and RH. The loading speed was adjusted for each of wood type to reach the destruction maximum within 90 ± 10 sec.

Wood surface hardness was determined according to the Brinell test procedure and meeting the requirements of the EN 1534 standard. The hardness for radial and tangential directions of specimens conditioned till constant mass at two RH (65% and 85%) were evaluated. A force of 1 kN was applied with reaching it within 15 sec and maintaining it for 25 sec by using a universal test device ZWICK Z100 and a metal ball of diameter 8 mm as an intender. Ten specimens were examined for each wood type and eight measurements were performed on radial and tangential surfaces of each specimen.

Results and Discussion

Density of wood before and after THT as well as mass loss due to thermal treatment, are presented in Table 1.

The wood mass loss increases with increasing THT temperature for both species at the temperature range used in the study. However, birch wood is considerably more sensitive to temperature compared to pine as equal mass losses are detected for birch at 160 °C and pine at 170 °C. Other authors have also reported similar findings (Rowell et al., 2009; Chaouch et al., 2010). It is explained by higher content of hemicelluloses, lower content of lignin and high content of syringyl groups in hardwood. Unlike the mass loss, the density dependence on temperature is significantly less pronounced and, compared to the initial material, the decrease for both woods varies in the range of 7 – 8%.

Examination of wood mechanical properties showed that both bending strength and hardness
decrease with increasing treatment temperature (Table 2 and 3). For samples conditioned at RH of 65% (20 °C), it was found that untreated pine wood is relatively soft with its Brinell hardness being approximately 16 – 18 N mm⁻². Treating of pine at the THT temperature range 160 – 180 °C, results in wood hardness decrease by 18 – 23% in the radial direction and by 19 – 23% in the tangential direction. However, increase of the THT temperature by 20 °C, causes hardness decrease by only 4 – 5%, whereas hardness reduction by 16 – 33% in the radial direction and by 27 – 48% in the tangential direction was detected for birch after THT treatment. Moreover, birch wood is affected to a greater extent by THT with hardness decrease almost twice due to rising the treatment temperature by 20 °C (from 150 °C to 170 °C). The birch wood surface hardness loss is significant, and it is an important indicator, especially for the articles subjected to horizontal loads (e.g., terraces).

Bending strength is more affected by thermal modification than hardness. Already at lower treatment temperatures, the bending strength for birch and pine decreases by 33% and 34%, respectively (Tables 3 and 4). The decrease in bending strength is close to linear within a modification temperature range of 160 – 180 °C for pine. For birch, the largest decrease in bending strength is caused by the increase in temperature from 150 °C to 160 °C. At the maximum treatment temperature of 170 °C, the bending strength for birch wood decreases by 56%. Bending strength for pine wood at this temperature decreases by 51% and by 60%, if the treatment temperature is increased up to 180 °C.

It is well known that the RH of the environment substantially affects wood properties (Simpson & TenWolde, 1999). For both unmodified and thermally modified samples conditioned at higher RH, the mechanical strength is significantly reduced comparing with samples conditioned at lower RH (Tables 2 and 3). At RH 85% for untreated pine, bending strength decreases by 25%, but for birch – by 32%, compared to similar parameters at RH 65%. However, for thermally modified wood, the decrease in bending strength at elevated humidity is

### Table 1

<table>
<thead>
<tr>
<th>Wood</th>
<th>Birch</th>
<th>Pine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment</td>
<td>unmodified</td>
<td>THT 150</td>
</tr>
<tr>
<td>ρ, g cm⁻³</td>
<td>0.618</td>
<td>0.593</td>
</tr>
<tr>
<td>STDEV</td>
<td>0.046</td>
<td>0.045</td>
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<td>Mass loss, %</td>
<td>-</td>
<td>0.1</td>
</tr>
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</table>

### Table 2

<table>
<thead>
<tr>
<th>Birch</th>
<th>RH 65%</th>
<th>RH 85%</th>
<th>Pine</th>
<th>RH 65%</th>
</tr>
</thead>
<tbody>
<tr>
<td>unmodified</td>
<td>Rad direct.</td>
<td>Tg direct.</td>
<td>Rad direct.</td>
<td>Tg direct.</td>
</tr>
<tr>
<td>THT 150</td>
<td>22.3 (3.4)</td>
<td>17.9 (2.7)</td>
<td>15.4 (1.5)</td>
<td>THT 160</td>
</tr>
<tr>
<td>THT 160</td>
<td>18.7 (3.2)</td>
<td>15.7 (3.2)</td>
<td>13.6 (1.5)</td>
<td>THT 170</td>
</tr>
<tr>
<td>THT 170</td>
<td>18.3 (2.3)</td>
<td>14.2 (1.9)</td>
<td>11.5 (1.3)</td>
<td>THT 180</td>
</tr>
</tbody>
</table>

(standard deviation in parentheses).

### Table 3

<table>
<thead>
<tr>
<th>Birch</th>
<th>RH 65%</th>
<th>RH 85%</th>
<th>Pine</th>
<th>RH 65%</th>
<th>RH 85%</th>
</tr>
</thead>
<tbody>
<tr>
<td>unmodified</td>
<td>116.8 (16.4)</td>
<td>79.2 (10.4)</td>
<td>unmodified</td>
<td>93.4 (13.9)</td>
<td>69.6 (8.9)</td>
</tr>
<tr>
<td>THT 150</td>
<td>78.7 (18.9)</td>
<td>60.2 (14.1)</td>
<td>THT 160</td>
<td>61.5 (15.9)</td>
<td>47.8 (11.7)</td>
</tr>
<tr>
<td>THT 160</td>
<td>57.9 (15.5)</td>
<td>41.2 (8.6)</td>
<td>THT 170</td>
<td>46.2 (18.3)</td>
<td>40.3 (18.1)</td>
</tr>
<tr>
<td>THT 170</td>
<td>59.2 (17.2)</td>
<td>40.7 (13.2)</td>
<td>THT 180</td>
<td>37.8 (19.6)</td>
<td>33.2 (12.7)</td>
</tr>
</tbody>
</table>

(standard deviation in parentheses).
significantly smaller. Similar trend has been reported by Arnold (2010). However, different results were obtained for thermally modified pine and birch wood regarding the effect of treatment temperature on strength reduction due to elevated RH. The reduction in bending strength of pine caused by rising RH decreased with increase in THT temperature. For the THT treated birch wood the decrease does not depend on the modification temperature. Similar effect of elevated RH was observed also with respect to the surface hardness of both tree species. The surface hardness of unmodified pine decreases by 29% in the radial direction and by 34% in the tangential direction. The surface hardness of THT pine wood also decreased but to a level that measuring failed and therefore no data on pine at RH 85% are presented in the table. The unmodified birch surface hardness decreases by 34% in the radial direction and by 23% in the tangential direction, compared to the samples conditioned at RH 65%. For the birch modified at 150 °C, the conditioning at RH 85% reduces the hardness in the radial and tangential direction by 24%. However, elevation of RH almost does not change the hardness for THT treated birch wood at higher (160 – 170 °C) temperatures. The differences between both species regarding the influence of humidity on hardness in different directions may be explained by the significantly different anatomical structure.

The changes in the wood component composition as a result of the thermal action have significant influence on the wood/moisture/water interaction. As a result of the degradation and/or mutual interactions in wood, with decreasing hydrophilic components, the equilibrium moisture, linear swelling and capillary water uptake decrease. The changes in wood EMC at different RH and depending on the modification temperature are presented in Fig. 1 and 2.

For both unmodified and THT wood, the EMC increases with rising RH, but for the modified one, EMC is significantly lower and decreases with increasing treatment temperature.

MEE is an important characteristic of modified wood that shows how much the equilibrium moisture decreases by modifying wood (Van Acker et al., 2015). This characteristic implies on improvement. In accordance with the normative documents, MEE at RH 85% must be > 40%. Figures 3. and 4. give MEE values for THT birch and pine.

In our case, this requirement is ensured for birch, modifying at 160 °C, whereas for pine – at only 180 °C.
The effect of thermal modification on the linear swelling of wood is shown in Table 4. As it can be seen, both the pine and birch wood swell less after thermal treatment and the swelling at similar RH decreases with increase in THT temperature. However, the results, obtained by soaking the wood in water up to maximum linear swelling was reached, show that there is no effect of the modification temperature on the wood dimensional stability above wood fibre saturation point. It agrees with the findings that changes in some other wood properties are dependent on the modification temperature only up to a certain

Table 4

<table>
<thead>
<tr>
<th>Modification temperature</th>
<th>Rad direction</th>
<th></th>
<th></th>
<th></th>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RH 45 %</td>
<td>RH 65 %</td>
<td>RH 85 %</td>
<td>(max)</td>
<td>RH 45 %</td>
<td>RH 65 %</td>
<td>RH 85 %</td>
</tr>
<tr>
<td>Birch</td>
<td>untreated</td>
<td>1.5 (0.1)</td>
<td>2.4 (0.3)</td>
<td>4.0 (0.5)</td>
<td>5.0 (0.7)</td>
<td>1.7 (0.2)</td>
<td>2.8 (0.3)</td>
</tr>
<tr>
<td>THT 150</td>
<td>0.9 (0.2)</td>
<td>1.3 (0.2)</td>
<td>2.0 (0.4)</td>
<td>2.5 (0.3)</td>
<td>1.2 (0.2)</td>
<td>1.8 (0.3)</td>
<td>3.0 (0.5)</td>
</tr>
<tr>
<td>THT 160</td>
<td>0.8 (0.1)</td>
<td>1.1 (0.2)</td>
<td>2.0 (0.4)</td>
<td>3.1 (0.9)</td>
<td>1.0 (0.1)</td>
<td>1.5 (0.2)</td>
<td>2.9 (0.4)</td>
</tr>
<tr>
<td>THT 170</td>
<td>0.7 (0.2)</td>
<td>1.1 (0.2)</td>
<td>1.8 (0.4)</td>
<td>3.6 (0.8)</td>
<td>0.9 (0.1)</td>
<td>1.4 (0.2)</td>
<td>2.4 (0.3)</td>
</tr>
<tr>
<td>Pine</td>
<td>untreated</td>
<td>1.4 (0.3)</td>
<td>2.0 (0.3)</td>
<td>3.2 (0.5)</td>
<td>5.2 (0.9)</td>
<td>2.3 (0.2)</td>
<td>3.4 (0.3)</td>
</tr>
<tr>
<td>THT 160</td>
<td>0.7 (0.1)</td>
<td>1.1 (0.1)</td>
<td>2.0 (0.2)</td>
<td>3.4 (0.4)</td>
<td>1.4 (0.2)</td>
<td>2.2 (0.2)</td>
<td>3.9 (0.3)</td>
</tr>
<tr>
<td>THT 170</td>
<td>0.7 (0.1)</td>
<td>1.1 (0.2)</td>
<td>1.9 (0.5)</td>
<td>3.0 (0.8)</td>
<td>1.1 (0.2)</td>
<td>1.8 (0.3)</td>
<td>3.4 (0.7)</td>
</tr>
<tr>
<td>THT 180</td>
<td>0.6 (0.1)</td>
<td>1.0 (0.3)</td>
<td>1.8 (0.4)</td>
<td>3.4 (0.8)</td>
<td>0.9 (0.1)</td>
<td>1.8 (0.3)</td>
<td>3.2 (0.6)</td>
</tr>
</tbody>
</table>

(standard deviation in parentheses).
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Conclusions

1. The results showed that, due to the significant strength losses during wood thermal treatment, only modification at 150 °C for birch wood and at 160 °C for pine wood is admissible to obtain intended material by the combined wood treatment.
2. The higher treatment temperatures resulted in greater improvement of wood hydrofobicidy; however, substantial decrease in wood swelling and equilibrium moisture content is obtained also at lower temperatures at which hardly any losses of wood mass were detected.

References

The authors gratefully acknowledge the financial support by the European Regional Development Fund project.

Johansson, Sehlstedt-Persson & Moren (2006). The explanation of the differences between the THT birch and THT pine CWU could be the differences in structural changes during thermal treatment, but it needs further research. However, these results suggest that impregnation regimes should be adjusted for each of THT treated species.
References


