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Performance evaluation of wood-based panels under a mild accelerated aging treatment

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Abstract

Accelerated aging treatments are often much more severe than necessary for final panel products that are being installed. It is thought that tests under these conditions unable to exhibit properly the degradation mechanism in approach for evaluating durability performance of wood panel products. This study aimed to determine the effects of a mild accelerated aging treatment and to characterize the degradation behavior of wood-based panels under this condition. We used four groups of panels: plywood, oriented strand board, particleboard, and medium density fiberboard. The panels were subjected to a wet-dry cycle treatment for 80 cycles. We used a vibrational non-destructive test to evaluate the strength properties and trace degradation behaviors of the panels during the cycle treatments. The mild aging condition that we chose generated fairly small weight and thickness changes, which indicated that there was mild intensity in the aging process. Elastic constant values decreased, whereas tan δ values increased when the number of cycles increased. The wet-dry cycle treatments were clearly responsible for the reduced strength properties and the degradation behavior of each panel product. However, our results suggest that the proposed mild aging conditions used in this study, though not certified for use, may be sufficient to degrade wood-based panels in ways that lead to predictable durability performance.


Introduction

The use of wood-based panels, as commonly for housing and building construction purposes, in Japan has been steadily increasing [1]. As a building material, wood-based panels are exposed to the environment during their use. Therefore, the durability performance of panel products must be determined as it affects the longevity of the product over its service life.

Durability performance can be assessed in two ways. The products can be directly exposed to the real usage environment and accelerated aging test. Accelerated aging treatments are practical for evaluating the durability performance of panel products. They can be used for quality control of panel products and can estimate how long a product will last. The accelerated aging test can be carried out in long term test and short term test. Outdoor exposure test is kind of long term accelerated aging test while simulated condition is a short term aging test. The outdoor exposure test can provide basic data because it combines various influencing factors. Many researchers have conducted these tests, including Deppe and Schmidt [2], Krahmer et al. [3], Hayashi et al. [4, 5], and Sekino and Suzuki [6], to study various panel types and locations over many years. However, the test has some shortcomings, such as requiring long exposure periods for testing. Deppe [7] reported that getting reliable data from outdoor exposure tests required at least five years of exposure, and since the product tests apply to a specific location, the test results cannot be generalized for use elsewhere.

The simulated condition test is employed over a particular time frame and is commonly used for short period tests. The test is intended to rapidly approximate the effects of many years of outdoor exposure and avoid the time consuming of long-term test. A number of studies have used this kind of aging test to determine the durability performance of panel products by using either standardized or laboratory-scale methods. Kajita et al. [8], Alexopoulos [9], Kojima et al. [10], and Kojima and Suzuki [11, 12] have worked with standardized methods such as the ASTM 6-cycle [13], V313 [14], APA D-1 [15], and JIS [16]. Others such as McNatt and Link [17] and McNatt and McDonald [18] have used
laboratory-scale methods, with some modified procedures. Some studies have also discussed this method’s relationship with outdoor exposure. Most of the results from these standardized methods tests have found much more severe damage than those from outdoor exposure tests. As a result, simulated tests cannot be used to reliably estimate outdoor exposure, as reported by Hann et al. [19], Vick [20], River [21], and Korai et al. [22].

The severity of simulated aging tests is caused by the extreme conditions of the treatment, particularly those involving liquid water, such as boiling, soaking, spraying, and vacuum pressure soaking. Therefore, several studies have attempted to use vapor water conditions in their aging tests. Chiu and Biblis [23] and Pu et al. [24] have reported on humid conditions in aging tests over short periods, which involved several hours of cycles that were evaluated using destructive bending tests. However, researchers have recognized a key shortcoming of the destructive testing: once the specimen is broken, it cannot be used again and many specimens are needed to conduct the test. Non-destructive evaluation has been well investigated for wood products, and some (e.g. Zhenbo et al. [25], and Bos and Casagrande [26]) have focused on wood-based panels. Nevertheless, there are few studies that focus on non-destructive evaluations under accelerated aging tests. However, Sun and Arima [27] have performed these tests using particleboard and oriented strandboard as specimens. The specimens were treated with boiling water for 60 minutes and then evaluated for strength properties and dimensional stability over several time periods. We observed that elongated cycle times with milder aging treatments led to only small losses in the product’s properties. Thus, this method may effectively degrade a product in ways that emulate environmental conditions. Furthermore, using non-destructive test helps to gather information about the degradation behavior of the wood panels over the course of the treatment.

Our research proposes milder conditions for the simulated aging test over longer time exposure. We consider a humid condition as a mild treatment, and this involves using vapor-based water to degrade panel products. Our objective in this research is to investigate the effects of the proposed aging condition, to assess the degradation rate of the panel
products, and to distinguish degradation behaviors induced by our mild aging test. Degradation behaviors can provide practical information on the durability of a panel product, which is especially relevant for commercial panel products in Japan.

**Materials and methods**

*Specimen preparation*

We used four groups of commercial panel products as specimens: plywood (PW), oriented strand board (OSB), particleboard (PB), and fiberboard (MDF). Each group of panels consisted of two panel types that differed in adhesive type, panel thickness, or wood species. The characteristic for each panel type are shown in Table 1. All specimens were prepared to be 50 mm wide and 250 mm long with four replications for each measurement. The specimens were initially conditioned at 40°C for 48 hours, after which we measured the dimensions and weight of each specimen and specified as the initial dimensions and weight.

*Mechanical properties*

We conducted a dynamic bending test using transverse vibration. A microphone was placed at one end of the specimen and the specimen was hit using a small hammer at the opposite end. The vibration signal, which was obtained by a microphone, was converted into a power spectrum by a fast Fourier transform (FFT) analyzer. We calculated the elastic constant ($E_d$) and the loss tangent ($\tan \delta$) from the peak resonance frequency, which corresponded to the first vibration mode and the amplitude of the resonance curve, respectively, as shown in the following equations:

\[
E_d = \frac{8\pi^2m_0^2a^4\rho^2}{800.5c^5} \quad \text{(GPa)}
\]

(1)

\[
\tan \delta = \frac{\lambda}{\pi}
\]

(2)
where $\rho$ is density (g/cm$^3$), $L$ is the length (mm), $f$ is the frequency of the resonance peak (Hz), $t$ is the thickness (mm), and $\lambda$ is the logarithmic decrement of the resonance amplitude which calculated by Hilbert transformation.

An accelerated aging treatment

We used an accelerated aging treatment designed as a mild test. Figure 1 shows the schematic diagram of the treatment. The treatment included a wet-dry cycle where each cycle had wet and dry states; 40°C and 90% relative humidity (RH) for 120 hours was characterized as the wet state, while 40°C for 48 hours without humidity control was characterized as the dry state. We repeated this treatment for 80 complete cycles in a climate-controlled chamber.

At the end of every state, we measured the dimensions and weight of each specimen and determined $E_d$ and $\tan \delta$ via the dynamic bending test. We calculated weight change (WC) and thickness swelling (TS) in every cycle on the basis of the initial weight and dimensions of the specimen, respectively. For conformity intensity of the condition test, we determined $\Delta WC$ and $\Delta TS$ with the following formulas:

$$\Delta WC_i = WC_{wi} - WC_{di} \quad (3)$$

Where $WC_{wi}$ is WC of wet condition at $i$ cycle and $WC_{di}$ is WC of dry condition at $i$ cycle.

$$\Delta TS_i = TS_{wi} - TS_{di} \quad (4)$$

Where $TS_{wi}$ is TS of wet condition at $i$ cycle and $TS_{di}$ is TS of dry condition at $i$ cycle.
Results and discussion

Intensity of the wet-dry cycle treatment condition

The effects of the aging treatment are shown in Figs. 2 and 3. Figure 2 shows the mean value and standard deviation of $\Delta WC$ for 80 cycles. We found that the wet-dry cycle treatment caused 8–11.5% $\Delta WC$ in the panels. There were a few differences in $\Delta WC$ for the mat-formed panel types (PB, MDF, and OSB) and in the veneer-based type. The $\Delta WC$ of plywood was approximately 11.5%, whereas the mat-formed panels ranged from 8–10%. This suggests that plywood absorbs much more moisture than do mat-formed panels. The mean value and standard deviation of $\Delta TS$ for all panel types are given in Fig. 3. We found that the wet-dry cycle treatment caused a 2.5–6% $\Delta TS$. The $\Delta TS$ for the plywood was 2.5% and the mat-formed panels ranged from 5–6%.

Figures 2 and 3 show that the intensity of the wet-dry cycle treatment accelerated the mat-formed panels. The conditions of this treatment yielded a $\Delta WC$ of ~10%. However, author private note by Kojima et al. [28], $\Delta WC$ of ASTM 6-cycle test has been found that could reach almost 53%. For $\Delta TS$, the values reached approximately 6% for the wet-dry cycle treatment, while a study by Kojima et al. [28] found a value of approximately 26%. Our wet-dry cycle treatment resulted in smaller $\Delta WC$ values, indicating that this treatment is milder than the standardized method. However, we consider that 10% of the moisture change in the wet-dry cycle treatments could be quite high. As a result, our method could still misrepresent degradation in the real environmental conditions.

Interestingly, the particleboards TS for each 1% change in moisture content was estimated using approach of weight change value and found to be about 0.4–0.6%. This was quite close to the thickness change in standard particleboard, which is approximately 0.3–0.5% for each 1% change in moisture content [29]. This confirmed that the conditions designed for this wet-dry cycle treatment yielded fairly small thickness changes for the panels and was nearly ideal for evaluating moisture changes that would occur during alternating wet
and dry conditions in the environment. This is relevant though our method is not yet certified.

TS of all panels increased with increasing the number of cycle with the exception of veneer base panels. Although the trend was similar but the increasing TS among the panels has differently proceeded during the wet-dry cycle treatment. To clarify the effects of the treatment on TS behavior, we observed OSB and PW for comparison as shown in Fig. 4. TS increased during the wet state because panels swell from moisture absorption and decreased during the dry state because panels shrink from moisture desorption. We dropped points and drew lines for the TS values in each state during the cycles. There was an obvious and repeated rise and fall over the 80 cycles. The thickness swelling of the OSB rapidly increased in the early cycles of the treatment, and then tended to level off after the first 20 cycles.

When panel products absorb moisture, either in vapor or liquid form, overall TS occurs both from the swelling of the particles themselves and the springback [30–31]. Particle swelling occurs because wood, from which the particles are made, is naturally hygroscopic. Furthermore, absorbed moisture could cause bond breakage and bring about particle separation. Springback is a result of releasing the stress of the board, which is caused by the compression of pressing. Nevertheless, when a board desorbs moisture and returns to a dry stage, there is part of the overall swelling never recovers to its original form. That is springback which generates permanent change of thickness and henceforth called irrecoverable TS.

The overall TS of the OSB after completed cycle was approximately 15% and about 9% of that did not recover which we defined as irrecoverable TS. When OSB was exposed to a wet state in the first cycle, TS was about 9% and when OSB continued to a dry state, only 6% of its swelling recovered, which we defined as recoverable TS. Notably, 3% was irrecoverable TS. The TS from the 20th cycle to the end of the cycles was quite similar. This indicates that the amount of irrecoverable TS increased from the first cycle up to 20 cycles but that the amount of recoverable TS in each subsequent cycle did not increase. Unlike
OSB, the overall TS for the PW panel was about 3% and the PW panel almost retained its initial thickness when it was exposed to a dry state in subsequent cycle. Even at the end of cycle, the treatment resulted in almost 3% TS value for PW. The amplitude of recoverable TS is almost same with advancing cycles in PW. As a result, we found that a wet-dry cycle treatment is not adequate for assessing the degradation of PW.

Overall, our wet-dry cycle treatment can cause TS to reach 9% after completing cycle. Conversely, ASTM 6-cycles can cause TS to reach about 24% [28]. Thus, the wet-dry cycle treatment resulted in smaller thickness changes than did the ASTM 6-cycles. Therefore this condition was well designed to act as a mild accelerated aging test.

**Strength retention under the wet-dry cycle treatment**

We examined strength property changes during the treatment using $E_d$. $E_d$ values during 80 cycles for all panel types are shown in Fig. 5. $E_d$ values also show an alternating rise and fall during the treatment, which corresponded to changes in TS and WC. These rise-fall curves illustrate $E_d$ behaviors during the wet-dry cycle treatment. Generally, they decreased gradually during the early cycles and then levelled off with later cycles.

There were some differences in $E_d$ amplitude, which we calculated from dry to wet state between the panel types. In other words, the effect of this treatment was different among panels. To explore this, we selected two of the eight panel types for comparison i.e. PBa and MDFa which had same 1.1 GPa initial amplitude. At the end of total cycle, it became 0.4 GPa for PBa and 0.8 GPa for MDFa. Thus, PBa experienced a larger loss of strength than did MDFa.

We tested the extent of $E_d$ retention in the wet state of cycle treatment and defined as follows:

$$E_d \text{ retention (\%)} = \left( \frac{E_d \text{ wet state}}{E_d \text{ initial}} \right) \times 100$$  \hspace{1cm} (5)
Ed retention of panel products is shown in Fig. 6. The wet-dry cycle treatment resulted in decreasing Ed values over 80 cycles, with different changes for each panel. Retention values were largest in PB, then followed by MDF, OSB, and PW. Ed retention in the wet state were 70–90% in the first cycle, then declined to 50–80% at the end of the cycle. Furthermore, the Ed retention value at the end of the complete cycle for the PW panels was higher than that for the mat-formed panels. That indicates that PW panels retained greater strength than did mat-formed panels. The rate of retention decreased fairly rapidly during the early cycles and then continued to decrease gradually until the final cycle. These retention values represent a loss of strength due to moisture deteriorating the structure of the panels during the adsorption process.

The Ed retention for all panels in Fig. 6 shows a clear decreasing trend, but in fact retention behavior during the wet-dry cycle treatment was very different for the different panels. To further explore this phenomenon, we used a degradation model developed by Suzuki and Saito [32] to elucidate the strength loss behavior for each of the panels over a prolonged treatment. Calculating Ed retention behavior for all the panels is based on the data in Fig. 6. The degradation model was as follows:

\[
F(t) = A + (100 - A) \exp \left( -t/B \right)
\]  

(6)

Coefficient \(A\) is the saturation value of the Ed retention and refers to the durability of the panel. Coefficient \(B\) is the decreasing rate and refers to how fast retention decreases. When panels are continuously subjected to the same treatment condition, we assumed that their strength would never reach zero. This means the panels would retain their strength after an infinite number of cycles. Using this model, we can predict the specific saturation value for strength retention after successive treatment cycles. Table 2 shows the coefficient values for all the panels, which are determined by Equation (6).

The coefficients in Table 2 were determined by non-linear least-squares regression analysis and show great variation. These coefficients also provided a useful overview of the different Ed retention behaviors among the panels over certain cycles. Except PWb, all
panels show that the values of $A$ are higher in the dry state than in the wet state. Another finding, except MDFb, the PW types had higher values of $A$ than did the mat-formed panel types. Comparing between mat-formed panels during the wet state, the values of $A$ for PBa and OSBa were 52% and 63%, respectively. These results indicate that, by this calculation, PBa was less durable than OSBa during the wet-dry cycle treatment, even though thickness swelling of PBa was slightly less than that of OSBa.

Based on the values of $B$, some panels experienced a decrease in strength at the very beginning of the cycles, and others toward the end of the cycles. However, to better explain this difference, we compared PBa and OSBa during their dry states. PBa, where $B$ is 13, has a higher coefficient than OSBa, where $B$ is 8. Since the rate of decrease corresponds to the number of cycles, the strength retention of PBa needed more time to reach its saturation value than did that of OSBa.

To clearly identify the different behaviors of strength loss for both PBa and OSBa panels, we plotted all $Ed$ retention data during the wet-dry cycle treatments and drew regression lines calculated by non-linear least-squares regression. Plots show the values of $Ed$ retention rates, while the solid and dotted lines represent the predicted value of the strength loss. Figure 7 shows the relationship between $Ed$ retention, when panels were tested at a dry state, and the number of cycles. As shown in those curves, the wet-dry cycle treatment degradation rates of the two panels differed significantly. The strength loss of PBa decreased continuously and moved more slowly toward its saturation value, whereas OSBa rapidly decreased within the first few cycles and then became relatively stable, as it had almost reached its saturation value.

The regression lines agreed well with the $Ed$ retention values from our experimental data as shown in Fig. 5. This means that Equation (3) can be used effectively to evaluate the strength loss of panels under wet-dry cycle treatments for an infinite number of cycles.
Estimating internal bond strength

The loss tangent (tan $\delta$) values of all the panels, which were computed by Equation (2), are plotted in Fig. 8 for each number of cycles. The tan $\delta$ behavior of the panels during the wet-dry cycle treatment is shown in Fig. 8. Loss tangent is a rheological term that corresponds to internal loss when a material vibrates. This is likely related to element attribution inside a board. Logically, this could be used as an indicator of internal bond strength.

The values of tan $\delta$ increased with each increase in the number of cycles. As shown by Obataya et al. [33], a TS of wood that results in $E_d$ reduction also results in increasing tan $\delta$. Notably, the internal bond strength, which is represented by the tan $\delta$ value, decreased for all panels during the wet-dry cycle treatment. The tan $\delta$ values showed an overall increasing trend, growing for the first 20 cycles and subsequently levelling off. This trend resembled the TS trend. This is understandable because moisture changes occurred during the treatment by adsorption and desorption, which caused TS in the panels. TS reduces the number of bonding points inside the board and leads to the degradation of internal bond strength.

Figure 8 shows the changes of tan $\delta$ during the wet-dry cycle treatment for 80 cycles. The tan $\delta$ values differed among the panels. The largest tan $\delta$ occurred in MDFa, which ranged from 0.03 at the first cycle to 0.045 at the end of cycles. The smallest change in tan $\delta$ from the first cycle to the last occurred in PBa, which ranged from 0.025 to 0.03. There was no big difference for both PW types while OSB types and PBb were found to be quite same trend. However, under this aging condition, the changes of tan $\delta$ were not consistent when associated with sequences of the changing TS values. We assumed that this was due to differences in wave propagation through the board, as element size, element direction, and adhesive type varied among panels. We thought that tan $\delta$ could be used as an index of mechanism degradation inside the panel. Therefore, relation between tan $\delta$ and durability is important issue to be discussed furthermore.
Conclusion

We subjected four groups of commercial panel products to a wet-dry cycle treatment. The proposed aging conditions yielded an 8–11.5% ΔWC and a 2.5–6% ΔTS. Our test used considerably milder conditions than those used in the ASTM 6-cycle standardized method. Our wet-dry cycle treatment caused a rise and fall of thickness swelling during the treatment and resulted in a significant irrecoverable thickness swelling for mat-formed panel types, though not for plywood types.

Our wet-dry cycle treatment considerably reduced the $E_d$ value over increasing numbers of cycles. The rate of $E_d$ retention decreased fairly rapid during early cycles in the treatment and then tended to level off. The residual $E_d$ retention rate of the panel products in the wet state was between 50–80% after completing 80 cycles.

The degradation behavior of each panel product was caused by extended wet-dry cycle treatments. This was well traced using vibrational non-destructive test and was also obvious from the predictive model. The value of tan $\delta$ increased with increasing numbers of cycles. However, our results suggest that this proposed aging condition was fairly well designed and may be effective for providing degradation information on wood-based panels. This could help predict the durability performance of mat-formed panel products.

Acknowledgments

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References


Table 1. Characteristics of the specimens

<table>
<thead>
<tr>
<th>Panel type</th>
<th>Abbreviation</th>
<th>Adhesive</th>
<th>Thickness (mm)</th>
<th>Density*2 (g cm⁻³)</th>
<th>MOR*3 (MPa)</th>
<th>MOE*4 (MPa)</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plywood</td>
<td>PWa</td>
<td>PF*⁵</td>
<td>12.2</td>
<td>0.60</td>
<td>49.3 ± 13.4</td>
<td>6.55 ± 0.84</td>
<td>5 plies</td>
</tr>
<tr>
<td></td>
<td>PWb</td>
<td>PF</td>
<td>8.8</td>
<td>0.58</td>
<td>71.8 ± 13.1</td>
<td>8.78 ± 1.16</td>
<td>3 plies</td>
</tr>
<tr>
<td>Oriented strand board</td>
<td>OSBa</td>
<td>PF</td>
<td>12.4</td>
<td>0.63</td>
<td>37.7 ± 8.9</td>
<td>4.90 ± 0.69</td>
<td>Aspen</td>
</tr>
<tr>
<td></td>
<td>OSBb</td>
<td>PF</td>
<td>11.8</td>
<td>0.65</td>
<td>36.0 ± 6.9</td>
<td>4.68 ± 0.62</td>
<td>Pine</td>
</tr>
<tr>
<td>Particleboard</td>
<td>PBa</td>
<td>PF</td>
<td>12.2</td>
<td>0.74</td>
<td>21.6 ± 3.5</td>
<td>3.44 ± 0.46</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>PBb</td>
<td>MDI*⁶</td>
<td>12.1</td>
<td>0.77</td>
<td>29.7 ± 2.4</td>
<td>3.97 ± 0.19</td>
<td>-</td>
</tr>
<tr>
<td>Medium density fiberboard</td>
<td>MDFa</td>
<td>MUF*⁷</td>
<td>12.2</td>
<td>0.75</td>
<td>44.9 ± 3.0</td>
<td>4.07 ± 0.22</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>MDFb</td>
<td>MDI</td>
<td>9.1</td>
<td>0.71</td>
<td>33.8 ± 1.4</td>
<td>3.10 ± 0.15</td>
<td>-</td>
</tr>
</tbody>
</table>

*¹ Lower-case letters are standing for distinguishing among two panels in the same panel type  
*² Density are given as average values  
*³ Modulus of Rupture, given as average ± standard deviation  
*⁴ Modulus of Elasticity, given as average ± standard deviation  
*⁵ Phenol formaldehyde  
*⁶ Methylene diphenyl diisocyanate  
*⁷ Melamine urea formaldehyde
Table 2. Coefficient values of the panels for the wet-dry cycle treatment

<table>
<thead>
<tr>
<th>Panel type*1</th>
<th>Dry state</th>
<th>Wet state</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>$A^2$</td>
<td>$B^3$</td>
</tr>
<tr>
<td>PWa</td>
<td>87</td>
<td>15</td>
</tr>
<tr>
<td>PWb</td>
<td>82</td>
<td>9</td>
</tr>
<tr>
<td>OSBa</td>
<td>73</td>
<td>8</td>
</tr>
<tr>
<td>OSBb</td>
<td>78</td>
<td>12</td>
</tr>
<tr>
<td>PBa</td>
<td>60</td>
<td>13</td>
</tr>
<tr>
<td>PBb</td>
<td>74</td>
<td>15</td>
</tr>
<tr>
<td>MDFa</td>
<td>73</td>
<td>13</td>
</tr>
<tr>
<td>MDFb</td>
<td>89</td>
<td>19</td>
</tr>
</tbody>
</table>

*1 The abbreviations refer to Table 1.
*2 The saturation value of the $Ed$ retention.
*3 The decreasing rate.
Fig. 1 Schematic diagram of a wet-dry cycle treatment
Fig. 2 Δ Weight change (%) of panel products after a complete 80 cycles
Fig. 3 Δ Thickness Swelling (%) of panel products after a complete 80 cycles
Fig. 4 The fluctuation of Thickness Swelling (%) for PWa and OSBa over 80 cycles
Fig. 5 $E_d$ (GPa) changes during each state over 80 cycles
Fig. 6. Ed retention (%) of the panel products in the wet state
Fig. 7. Ed retention and retention model (F(t)) of PBa and OSBa during the wet-dry cycle treatment in the dry state.
Fig. 8. Tan $\delta$ of the panel products in relation to the wet state during the wet-dry cycle treatment