

Impacts of Impregnation with Fire Retardant Chemicals on the MOE in Bending of Some Woods

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ABSTRACT

This study has been performed to determine the effects of impregnation with fire retardant chemical materials on the modulus of elasticity (MOE) in bending of Oriental beech, European oak and Scotch pine wood materials. To achieve this goal, test samples prepared from woods of Oriental beech, European oak and Scotch pine according to TS EN 345 regulations were impregnated with ammonium-sulfate [(NH₄)₂SO₄], sodium acetate (NaC₂H₃O₂·3H₂O), aluminum chloride (Al₂C₆I₂H₂O), borax [Na₂B₄O₇·5H₂O], boric acid [H₃BO₃] and, borax + boric acid (w:w=%50:50). The modulus of elasticity in bending of impregnated wood samples were determined according to TS EN 408. Consequently, according to wood species; modulus of elasticity in bending was found the highest value at beech (10350 N/mm²) and the lowest value at pine wood (9501 N/mm²). According to variety of impregnation; modulus of elasticity in bending values were found no statistical difference between control samples and impregnated test samples. Considering the interaction of wood type and process; modulus of elasticity in bending was found the highest value at beech + borax (11450 N/mm²) and the lowest value at pine + control samples (8223 N/mm²). As a result, in the massive construction and furniture elements that the modulus of elasticity in bending after the impregnation with borax is of great concern, Oriental beech wood materials could be recommended.

Keywords: Fire retardant chemicals, modulus of elasticity in bending, impregnation, wood.

ÖZ

Bu çalışma, Yangın geciktirici kimyasallar ile empenye etmenin ağaç malzemelerin eğilmede elastiklik modülüne etkilerini belirlemek amacıyla yapılmıştır. Bu maksatla, ülkemiz orman ürünleri endüstrisinde yaygın olarak kullanılan Doğu kayını, sapsız meşe ve sarıçam odunlarından TS EN 345 esaslarına göre hazırlanan deney örnekleri; amonyum sülfat [(NH₄)₂.SO₄], sodyum asetat (NaC₂H₃O₂·3H₂O), alüminyum klorit (Al₂C₆I₂H₂O), boraks [Na₂B₄O₇·5H₂O], borikasit [H₃BO₃] ve boraks + borikasit (w:w=%50:50) ile ASTM D 1413 esaslarına uyularak vakum yöntemi ile empenye edilmiştir. Empenye edilen deney örneklerinin eğilmede elastiklik modülü değerleri TS EN 408 standardı esaslarına belirlenmiştir. Sonuç olarak; ağaç türüne göre, eğilmede elastiklik modülü değerleri en yüksek kayında (10350 /mm²), en düşük sarıçam odununda (9501 N/mm²) elde edilmiştir. İşlem çeşidine göre; kontrol örnekleri ile işlem çeşidi sonuçları arasındaki fark istatistiksel olarak önemsiz çıkmıştır. Ağaç türü + işlem çeşidi etkileşimine göre eğilmede elastiklik modülü değeri en yüksek kayın + boraksda (11450 N/mm²), en az sarıçam + kontrolde (8223 N/mm²) bulunmuştur. Buna göre eğilmede elastiklik modülü değerinin önemli olduğu yapı ve mobilya elemanları üretiminde boraks ile empenye edilmiş kayın odunu tercih edilebilir.

Anahtar Kelimeler : Yangın geciktirici kimyasallar, eğilmede elastiklik modülü, empenye, ağaç malzeme.

1. INTRODUCTION

Wood is an environmentally desirable material for fiber and structural use. It is efficient in both economic and environmental costs to users. To extend its utility into new markets, wood is sometimes treated with chemicals [1]. One of the major objections to the use of wood for many purposes is of course the question of its long-term resistance to the natural processes of degradation, particularly at sites and in situations where there is high biological hazard and where no form of chemical or physical protection is afforded to the material. With an increased demand for timber worldwide and moves towards fast-grown plantation species, the need to impart additional protection, usually in the form of chemical treatment, has become necessary to confer long-term

performance in these wood products [2]. In case wood is not impregnated but only painted and varnished instead, the prevention on the surfaces is limited to a maximum of two years [3].

It is reported that, in mines, as a result of the impregnation of the beech and spruce wood with water-soluble salts, the bending, tensile and impact strength decreased a little whereas compression strength increased [4]. In another research concerning the impregnation of pine, spruce, fir, beech and poplar woods with antrasen, it was found that, the compression strength increased by 6-40 % and bending strength increased by 10-22 % [5]. In the impregnation of pine and beech wood with UA salts and tar oil, the tar oil increased compression strength by 10 % and UA salts increased with a small rate. On the other hand, the tar oil increased the bending strength whereas the UA salts diminished the bending strength. [6].

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Vologdin declared that, among the materials used for the impregnation of pine; Sodium pentachlorofenol, Cuppersulphate and Sodium fluoride increased the compression strength respectively by 95 %, 25 % and 3 % whereas Zinc chloride decreased the compression strength by 9%. Sodium pentachlorofenol also increased the bending strength [7]. In another study, pressure treatment caused to a decrease of 8-10 % in the bending strength of different wood types [8].

It was assessed that, salty impregnation materials increased the compression strength by 4.6-9.6 %, whereas decreased the bending strength by 2.9-16 % [9]. After the impregnation of pine wood samples by hot-cold open tank method with eleven preventives, no significant difference was observed in the bending strength except the decreasing effects of fluotox containing acid florid [10]. In another study, chromate copper arsenate (CCA) and arsenate copper arsenate (ACA) salts did not caused any significant impact on modulus of elasticity in bending [11].

Fire retardant chemicals can also reduce the strength of lumber or plywood, an effect related to the nature of the chemicals and to the re-drying temperatures used in the treating process [12].

In this study, Oriental beech, European oak, Scotch pinewoods commonly being used in furniture manufacturing and massive constructions were examined with respect to the effects of impregnation with fire retardant chemical materials on the modulus of elasticity in bending of Oriental beech, European oak and Scotch pine wood materials.

2. MATERIAL AND METHOD

2.1. Materials

2.1.1. Solid Woods

The solid woods to be used as test samples were randomly selected from the timber merchants of Ankara. Specific pains were taken for the selection of wood materials. Accordingly, non-deficient, proper, knotless, normally grown (without zone line, without reaction wood and without decay, insect mushroom damages) wood materials were selected.

2.1.2. Protective Chemicals

In this study, preservative chemicals; borax, boric acid, boric acid + borax (w:w=%50:50) ammonium sulfate, sodium acetate, aluminum chloride, are used to impregnate the test samples.

Borax and boric acid were obtained from Etibank - Bandırma (Turkey) borax and acid Factory. Boric acid [H_3BO_3] contains %56.30 $\frac{1}{2} B_2O_3$ %43.7 H_2O with a molecular weight 61.84, density $1.4 g.cm^{-3}$, melting point $171 ^\circ C$. Borax [$Na_2B_4O_7 \cdot 5H_2O$] contains %21.28 Na_2O %47 B_2O_3 , %30.9 H_2O with a molecular weight 291.3, density $1.8 g/cm^3$, melting point $741 ^\circ C$ [13].

Ammonium sulfate ($(NH_4)_2SO_4$) is the molecular weight of 132.14, decompose above $^\circ C$ 280. Aqueous solution

shows weak acid functionality. Aluminum chloride ($AlCl_3 + 3KCl + Al$) is density of $2.698 g/cm^3$ at $25 ^\circ C$, a melting point of $659.7 ^\circ C$ boiling point is $2057 ^\circ C$. Sodium acetate ($C_2H_3NaO_2$) is obtained from the reaction of acetic acid, sodium carbonate or sodium hydroxide. There are $58 ^\circ C$ water lost trihydrate and anhydrous forms. Water-soluble and ethoxyethyl Zealand; slightly soluble in ethanol [14].

2.2. Method

2.2.1. Determination of density

The densities of wood materials, used for the preparation of test samples were determined according to TS 2472 [15]. For determining the air-dry density, the test samples with a dimension of $20 \times 30 \times 30$ mm were kept under the conditions of $20 \pm 2 ^\circ C$ temperature and 65 ± 3 % relative humidity until they reached to a stable weight. The weights were measured with an analytic scale of ± 0.01 g sensitivity. Afterwards, the dimensions were measured with a digital compass of ± 0.01 mm sensitivity. The air-dried densities (δ_{12}) of the samples were calculated by the formula;

$$\delta_{12} = \frac{W_{12}}{V_{12}} g.cm^{-3} \quad [1]$$

Where W_{12} is the air-dry weight (g) and V_{12} is the air-dry volume (cm^3).

The samples were kept at a temperature of $103 \pm 2 ^\circ C$ in the drying oven until they reached to a stable weight for the assessment of oven-dry density. Afterwards, oven-dry samples were cooled in the desiccator containing phosphorus pentoxide (P_2O_5). Then, they were weighted on a scale of ± 0.01 g sensitivity and their dimensions were measured with a digital compass of ± 0.01 mm sensitivity. The volumes of the samples were determined by stereo metric method and the densities (δ_o) were calculated by the formula;

$$\delta_o = \frac{W_o}{V_o} g.cm^{-3} \quad [2]$$

Where W_o is the oven-dry weight (g) and V_o is the oven-dry volume (cm^3).

2.2.2. Determination of humidity

The humidity of test samples before and after the impregnation process was determined according to TS 2471 [16]. Thus, the samples with a dimension of $20 \times 20 \times 20$ mm were weighed and then oven-dried at $103 \pm 2 ^\circ C$ till they reach to a constant weight. Then, the samples were cooled in desiccator containing phosphorus pentoxide (P_2O_5) and weighed with an analytic scale of 0.01 g sensitivity. The humidity of the samples (r) was calculated by the formula;

$$r = \frac{Mr - Mo}{Mo} \times 100 \quad [3]$$

Where Mr is the initial weight (g) and Mo is the oven-dry weight (g).

2.2.3. Preparation of the test samples

The rough drafts for the preparation test and control samples were cut from the sapwood parts of massive woods and conditioned at a temperature of 20±2 °C and 65±3 % relative humidity for three months until reaching an equilibrium in humidity distribution. The samples for compression strength test, with a dimension of 20x20x400 mm were cut from the drafts having an average humidity of 12 % according to TS EN 408 [17]. The densities and humidity values of all test samples were measured before the impregnation process.

The test samples were impregnated according to ASTM D 1413 [18], TS 344 [19] and TS 345 [20]. The samples were dipped in the impregnation pool immersing 1 cm below the upper surface for 2 hours for medium-term dipping. The specifications of the impregnation solution were determined before and after the process. The processes were carried out at 20±2 °C temperature. Retention of impregnation material (R) was calculated by the formula;

$$R = \frac{G.C}{V} 10^3 kg.m^{-3} \quad G = T_2 - T_1 \quad [4]$$

Where G is the amount of impregnation solution absorbed by the sample (g), T₂ is the sample weight after the impregnation (g), T₁ is the sample weight before the impregnation (g), C is the concentration (%) of the impregnation solution and V is the volume of the samples (cm³).

Impregnated test samples were kept under a temperature of 20±2 °C and 65±3% relative humidity until they reach to a stable weight.

2.2.4. The modulus of elasticity (MOE) in bending

The tests for modulus of elasticity in bending were carried out with the Universal Testing Equipment, according to TS EN 408. The capacity of the Universal Testing Equipment was 400 N. Deformations on the test samples were measured in the middle of the specimen within a zone of five times the width of the sample by comparator. The deformations by incrementally increasing the forces were assessed with a sensitivity of 0.01 mm. In the elastic deformation zone, modulus of elasticity (MOE) was calculated with the following formula [21, 22];

$$MOE = \frac{\Delta F.L^3}{4.b.h^3.\Delta f} \quad N \text{ mm}^{-2} \quad [5]$$

Table 2. The results of analysis of variance for retention amounts

SOURCE	Degrees of freedom	Sum of square	Means of square	F value	Sig. α ≤ 0.05
Wood types (A)	2	4224.239	2112.120	71.0438	0.0000
Impregnation chemicals (B)	5	4167.535	833.507	28.0360	0.0000
Interaction AB	10	2238.093	223.809	7.5281	0.0000
Error	162	4816.234	29.730		
Total	179	15446.101			

where ΔF is the difference between the arithmetic average of the upper and lower limits of applied force in the elastic deformation zone (N), Δf is the net elastic deflection - difference between the measured elastic deflection in the upper and lower loading limits- (mm), L is the span (mm), b is the cross-sectional width of test sample (mm), h is the cross-sectional thickness of the test sample (mm)

2.3. Data Analysis

A total of 99 samples (3 x 3 x 11) were prepared. The effects of wood material and impregnation method on the modulus of Elasticity in Bending were analyzed by Analysis of Variance. Duncan’s Multiple Range Test was also applied where appropriate.

3. RESULT AND DISCUSSION

3.1. Density

Statistical values of average air-dried densities that are used in the experiments samples have been shown in Table 1.

Table 1. Statistical values of air-dried density averages

Statistical values	Wood Materials		
	Beech	Oak	Pine
X (g.cm ⁻³)	0.692	0.665	0.573
Ss (g.cm ⁻³)	0.01961	0.02368	0.01305
v (s ²)	0.00042	0.00062	0.00019
min (g.cm ⁻³)	0.648	0.624	0.552
max (g.cm ⁻³)	0.715	0.695	0.595
N	10	10	10

X: Arithmetic mean, v: Variance, Ss: Standard deviation, N: Number of samples

According to the Tab.1 the highest air-dried density value was found in beech and the lowest in pine wood. The air-dried density values of massive wood materials in literature; Ash (0.690 g.cm⁻³), beech (0.660 g.cm⁻³), oak (0.650 g.cm⁻³), walnut (0.680 g/cm³), pine (0.520 g.cm⁻³), poplar (0.502 g.cm⁻³) [23]. These values have shown parallelism with air-dried density values of experimented wood materials.

3.2. Retention Amount

The results of multiple variance analyses with regard to the effects of wood type and impregnation chemicals are given in Table 2.

The results of the analysis of variance indicated that the effects of the wood types, impregnation chemicals and their interaction were found to be statistically significant ($\alpha \leq 0.05$). Average values the retention amounts of different wood and process types are given in Tab. 3.

Table 3. Average the retention amounts of different types of wood and types of process

WOOD TYPES	X	HG
Pine (I)	19.39	A
Beech (II)	18.60	A
Oak (III)	8.74	B
IMPREGNATION CHEMICALS	X	HG
Borax (Bx)	22.70	A
Borax + Boric acid (Bx+Ba)	19.77	B
Sodium acetate (Sa)	17.90	B
Boric acid (Ba)	12.46	C
Aluminum chloride (Ac)	10.37	C
Ammonium sulfate (As)	10.26	C

LSD : $\pm 1,964$, X: Average value, HG: Homogeneous group
As shown in Table 3, according to types of wood highest retention amount was obtained in pine (19.39 kg/m³) and

chemicals are shown Tab.4 and the graphic is shown in Figure 1.

According to interaction of wood type and impregnation chemicals highest retention amount was obtained in pine impregnated with Bx (32.67 kg/m³) and the lowest in oak impregnated with As (6.355 kg/m³) samples.

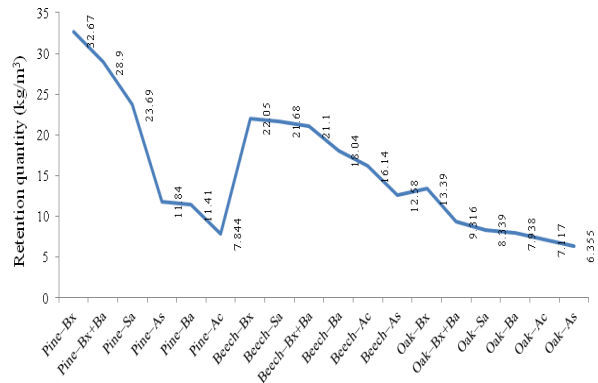


Figure 1. The retention amount of the interaction of wood type and impregnation chemicals

Table 4. The Duncan test results of the interaction of wood type and impregnation chemicals

Type of wood and impregnation chemical	X	HG	Type of wood and impregnation chemical	X	HG
Pine (I)-Bx	32.67	A*	Beech (II)-Ba	18.04	CDE
Pine (I)-Bx+Ba	28.90	A	Beech (II)-Ac	16.14	DEF
Pine (I)-Sa	23.69	B	Beech (II)-As	12.58	FGH
Pine (I)-As	11.84	FGHI	Oak (III)-Bx	13.39	EFG
Pine (I)-Ba	11.41	FGHI	Oak (III)-Bx+Ba	9.316	GHI
Pine (I)-Ac	7.844	GHI	Oak (III)-Sa	8.339	GHI
Beech (II)-Bx	22.05	BC	Oak (III)-Ba	7.938	GHI
Beech (II)-Sa	21.68	BC	Oak (III)-Ac	7.117	HI
Beech (II)-Bx+Ba	21.10	BCD	Oak (III)-As	6.355	I**

LSD : ± 4.812 , * : the highest, ** : the lowest

Table 5. The results of analysis of variance for modulus of elasticity in bending

SOURCE	Degrees of Freedom	Sum of Square	Means of Square	F Value	Sig. $\alpha \leq 0,05$
Types of wood (A)	2	26226815.29	13113407.64	2.9744	0.0535
Imp. chemicals (B)	6	16691460.67	2781910.11	0.6310	0.0512
AB	12	86576799.79	7214733.31	1.6364	0.0845
Error	189	833265099.32	4408810.04		
Total	209	962760175.08			

the lowest in oak (8.74 kg/m³) wood samples. This situation could be attributable to the impact of permeability of pine wood and due to tyloses in oak wood samples. According to impregnation chemicals, highest retention quantity was obtained in Bx (22.70 kg/m³) and the lowest in As (10.26 kg/m³). This may be due to chemical properties of Bx and As. The Duncan test results of the interaction of wood type and impregnation

3.3. MOE in Bending

The results of multiple variance analyses with regard to the effects of wood type and impregnation chemicals for modulus of elasticity in bending are given in Table 5.

The results of the analysis of variance indicated that the effects of the wood types, impregnation chemicals and their interaction were not found to be statistically

significant ($\alpha \leq 0.05$). The average modulus of elasticity in bending values of different types of wood and types of process are given in Table 6.

Table 6. Average the MOE in bending values of different types of wood and types of process (N/mm²)

WOOD TYPES	X	HG*
Beech (II)	10350	A
Oak (III)	10090	B
Pine (I)	9501	B
IMPREGNATION CHEMICALS	X	HG**
Bx	10520	A
Bx+Ba	10140	A
Control	10030	A
Sa	9971	A
Ba	9894	A
Ac	9747	A
As	9554	A

*LSD : ± 699.7 , **LSD: ± 1069 , X: Average value, HG: Homogeneous group

As clearly shown in Table 6, according to types of wood the highest modulus of elasticity in bending value was obtained in beech (10350 N/mm²) and the lowest in pine (9501 N/mm²) wood. According to impregnation chemicals, the highest modulus of elasticity in bending value was obtained in Bx (10520 N/mm²) and the lowest in As (9554 N/mm²). Also results of control samples and impregnation chemicals were not found to be statistically significant.

The Duncan test results of modulus of elasticity in bending of the interaction of wood type and impregnation chemicals are shown Tab.7 and the graphic is shown in Figure 2.

Table 7. The Duncan test results of the interaction of wood type and impregnation chemicals

Type of wood and impregnation chemical	X	HG
Pine (I)-Sa	10500	ABC
Pine (I)-Bx+Ba	10410	ABCD
Pine (I)-Bx	9968	ABCD
Pine (I)-Ac	9839	ABCD
Pine (I)-As	9054	BCD
Pine (I)-Ba	8511	CD
Pine (I)-Control	8223	D**
Beech (II)-Bx	11450	A*
Beech (II)-Control	10800	AB
Beech (II)-Sa	10200	ABCD
Beech (II)-As	10180	ABCD
Beech (II)-Ba	10090	ABCD

Beech (II)-Bx+Ba	10030	ABCD
Beech (II)-Ac	9665	ABCD
Oak (III)-Ba	11080	AB
Oak (III)-Control	11050	AB
Oak (III)-Bx	10130	ABCD
Oak (III)-Bx+Ba	9965	ABCD
Oak (III)-Ac	9738	ABCD
Oak (III)-As	9432	ABCD
Oak (III)-Sa	9214	ABCD

LSD: ± 1851 , *: the highest, **: the lowest

According to interaction of wood type and impregnation chemicals the highest modulus of elasticity in bending value was obtained in impregnated beech with Bx (11450 N/mm²) and the lowest in control pine (8223 N/mm²) samples.

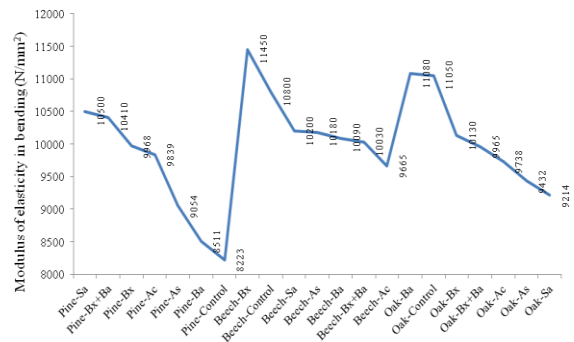


Figure 2. MOE of the wood type and impregnation chemicals interaction

4. CONCLUSION

The retention amounts were found different depending on wood type and impregnation materials. According to types of wood highest retention amount was obtained in pine wood (19.39 kg/m³ -21.81%) and the lowest in oak wood samples (8.742 kg/m³-9.15%); according to impregnation chemicals highest retention amount was obtained in Bx (22.70 kg/m³) and the lowest in As (10.26 kg/m³). According to interaction of wood type and impregnation chemicals highest retention amount was obtained in pine impregnated with Bx (32.67 kg/m³) and the lowest in oak impregnated with As (6.355 kg/m³) samples. Accordingly, the highest values were obtained in pine samples. This case may be due to structural properties of pine wood. In the literature, Özçiftci and Batan [24] reported that retention amount of Scotch pine higher than Uludag fir and Oriental beech.

Modulus of Elasticity in Bending; according to types of wood the highest modulus of elasticity in bending value was obtained in beech (10350 N/mm²) and the lowest in pine (9501 N/mm²) wood. Depending on the species of impregnation chemicals, the highest modulus of elasticity in bending (10520 N/mm²) was that of Bx, followed by Bx+Ba, control samples, Sa, Ba, Ac and As (10140 N/mm², 10030 N/mm², 9971 N/mm², 9894

N/mm², 9747 N/mm², and 9554 N/mm², respectively). According to this, it can be said that, Bx and Bx+Ba showed an increasing effect on the modulus of elasticity in bending compared to the control samples, however other impregnation chemicals showed reducing effect on the modulus of elasticity in bending.

According to interaction of wood type and impregnation chemicals the highest modulus of elasticity in bending value was obtained in impregnated beech with Bx (11450 N/mm²) and the lowest in control pine (8223 N/mm²) samples. In general, impregnation chemicals showed an increasing effect on pine wood, however decreasing effect showed on beech and oak wood, excluding Bx for beech and Ba for oak. Colakoglu et al. [25] reported that modulus of elasticity in bending values of laminated veneer lumber treated with 1 % boric acid were decreased for 5.1 % compared to untreated control sample. Gerhards [26] found that fire retardant chemical treatment and kiln-drying reduce the MOE of wood for 5 % in average. Consequence, in the massive construction and furniture elements that the bending strength after the impregnation with borax is of great concern, Oriental beech wood materials could be recommended.

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