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2 Softwood-lignin/natural rubber composites containing novel plasticizing agent: Preparation and characterization

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ABSTRACT: Composite materials based on natural rubber were obtained by using 8 9 glycerolysate (decomposition product of polyurethane) as a novel plasticizer. In order to determine the effect of various lignin content, four different filler amounts were 10 used, namely 5 phr (parts per 100 parts of natural rubber) of lignin (WLI5G), 10 phr of 11 12 lignin (WLI10G), 20 phr (WLI20G), and 40 phr (WLI40G). The reference specimen without lignin (WLI0G) was also prepared. The resulting vulcanizates were analyzed 13 by Fourier Transform Infrared Spectroscopy (FTIR) to determine the chemical 14 interaction between the lignin powder and the natural rubber chain. The SEM 15 analysis of the cross-sections of the obtained materials was carried out to determine 16 17 the adhesion between lignin and rubber. The results of dynamic mechanical analysis (DMA) and thermogravimetric analysis (TGA) showed that the samples containing 5 18 and 10 phr of lignin had the best thermal properties. Also, the measured mechanical 19 properties, such as tensile strength, hardness, resilience and abrasiveness, 20 confirmed these findings. 21

22 KEYWORDS:

23 Softwood Lignin;

24 Natural rubber composit	es;
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- 25 Scanning Electron Microscopy;
- 26 Thermal analysis;
- 27 Mechanical properties;
- 28 Equilibrium swelling measurements
- 29

30 1. INTRODUCTION

At present, many scientific teams work on the possibility to employ natural 31 resources in the field of materials science and engineering. It is a consequence of the 32 33 ending stocks of petrochemical resources such as coal, natural gas, and crude oil. One of the most important issues related to green chemistry is the utilization of 34 renewable resources as a new application in the existing products. Moreover, the 35 polymer recycling process and the utilization of polymer recycling products constitute 36 the second most urgent task due to the increase in the waste quantity. It has been 37 proven that the renewables and the products of polymer recycling can partially 38 replace the primary resources used in the polymer synthesis and preparation. The 39 resulting materials display the same, approximately the same, or even better 40 properties. 41

One of the most recent issues in green chemistry is finding the possibility to maximize the application of lignin by-products, which are produced in vast amounts by the paper and pulp industries, in the field of polymer technology. Presently, ca. 50 million tons of lignin by-products produced annually by the industries are used as a

fuel for the energy production (Faruket al., 2016a). Only 2 % of this valuable raw material is used for other applications because of the complex structure and heterogeneity of lignin, which causes difficulties during delignification on a commercial scale that is conducted by the pulp and paper industries (DeWild et al., 2014; Neutelings, 2011).

Lignin is a highly branched bio-macromolecule. It is composed of units such as 51 paracoumaryl alcohol, coniferyl alcohol and sinapyl alcohol (Jianget al., 2014). These 52 53 phenylpropane units have none, one, or two methoxyl groups at the positions 3 or 5 in the phenolic ring (Faruket al., 2016a). The aforementioned molecules are linked 54 together by different bond types, i.e. 5-O-4, β -O-4, β -1, β -5, β - β , etc. (Rogers, 2015). 55 Chakar et al. (Chakarand Ragauskas, 2004) presented the percentages of different 56 bonds in softwood lignin, β -O-4 being the most common one. This particular bond 57 type is present in up to 50 % of total bonds in softwood-lignin. 58

The varying occurrence of bond types depends on the origin of lignin, i.e. the 59 sources of softwood or hardwood, which results in different percentages of bond 60 content.Bjornsson indicated that generally hardwoods contain less lignin than 61 62 softwoods (Bjornsson, 2014). The exact lignin structure not only depends of the type of biomass, but also of the type of delignification process used, which modifies lignin 63 64 to a certain degree (Hatakeyama and Hatakeyama, 2010). The application of various 65 delignification methods results in a variety of lignin products. The main types of lignin can be divided into lignosulfonates, kraft lignin, and organosolv lignin (Holladay et al., 66 2007). 67

68 The delignification process affects the content of impurities in the obtained 69 products and, consequently, their further applicability. Lignosulfonates can be used

as dispersants, emulsion stabilizer, carbon black, industrial binders, agricultural 70 71 chemicals or concrete additives due to their medium purity (residual sulfur). Kraft lignin also contains some ash and sulfur, and can be used as emulsifiers, 72 dispersants, carbon fibers or binders. Organosolv lignin, which is sulfur free, has the 73 highest degree of purity. Because of this property, it is possible to use organosolv 74 lignin for the synthesis of aromatic polyols, new diacids, carbon fibers, activated 75 carbon, phenolic resins, phenol derivatives and antioxidants (Faruket al., 2016a; 76 Holladay et al., 2007). 77

Plasticizers, represented by freely available, non-volatile compounds, are 78 widely used in the polymer production due to the important role they play in the 79 resulting products. They improve processability during the polymer preparation as 80 well as add flexibility to the final materials (Vieiraet al., 2011). Over the past years, 81 due to the increasing interest of polymer industries in biopolymers, biorenewables 82 and the products of the chemical recycling of polymer waste, many publications 83 appeared in which the addition of a biopolymer plasticizer and/or biodegradable 84 plasticizing materials of natural origin had been described. Altenhofen Da Silva and 85 co-workers utilized the product of the polyesterification of rice fatty acid as a 86 plasticizer in the poly(vinyl chloride) and natural rubber films. The results of this 87 research indicate that the addition of natural plasticizer increased the elongation at 88 break compared to pure polymer film. In terms of thermogravimetric analysis no 89 significant differences were detected between the plasticized material and pure 90 natural rubber-based product (Altenhofen Da Silva et al., 2011). Alexander and 91 Thachil investigated differences between cardanol and aromatic oil emloyed as 92 plasticizers. It was demonstrated that cardanol, used as a plasticizing agent in the 93 94 natural-rubber matrix, gave mechanical properties similar to those obtained with the aromatic oil-based materials (Alexander and Thachil, 2006). The same observation
was later reported by Mohapatra and Nando(Mohapatra and Nando, 2014).

The main aim of this research was to prepare and characterize natural rubber-97 based composites filled with various amounts of lignin, and obtained with the use of 98 99 glycerolysate. This experimental set up was an exemplification of the application of the chemical recycling of polymer product as a novel plasticizer in the natural rubber 100 matrix. The synthesis of the plasticizer used in our study had been based on the 101 102 decomposition of polyurethane waste by means of chemicals, heat and catalysts. Polyurethanes are in the sixth place in the ranking of the most used polymers in the 103 global market. The polyurethane waste constitutes ca. 6 % of all plastic waste 104 (Kopczyńska and Datta, 2016). As a result, the chemical recycling processes of 105 polyurethanes are one of the most prolific development tasks in the materials 106 engineering (Simonet al., 2014). Moreover, the most investigated are the wastes of 107 108 flexible polyurethane foams due to their extensive application and the resulting high volume of waste that entails environmental and economic problems (Nikjeand Nikrah, 109 2007). Glycerolysate is the product of thermo-mechanical recycling of polyurethane, 110 111 with the use of glycerine as a decomposition prime mover. The process is called glycolysis or glycerolysis, and it is the transesterification reaction between hydroxyl 112 groups in glycol or glycerol (glycerine), respectively, which interchanged the ester 113 groups in the polyurethane chains (Simónet al., 2016). The decomposition product of 114 polyurethane is a mixture of compounds and monomers. Until now, glycerolysates 115 were most frequently used to synthesize polyurethanes, mainly in the form of foams 116 (Simónet al., 2016) or elastomers (Datta, 2010; Datta and Pasternak, 2005), in which 117 polyols were partially or completely replaced by the product of the chemical recycling 118 119 of polyurethane.

This paper describes novel elastic composites containing four different levels of lignin, and the same amount of plasticizer in the form of glycerolysate. The influence of lignin content on the structure, morphology, and the selected mechanical, thermal and chemical properties of the obtained composites was investigated.

125 **2. EXPERIMENTAL**

126 **2.1. MATERIALS**

Natural rubber used to prepare the composites was purchased from Torimex 127 Chemicals Ltd Sp. z o. o., KonstantynówŁódzki, Poland (density 0.92 g/cm³, weight 128 average molecular weight, $M_w = 800\ 000\ mol/g$). Lignin (INDULIN AT – kraft pine 129 lignin - softwood lignin) used in this study was obtained from MeadWestvaco 130 Corporation, Specialty Chemical Division, South Carolina, USA. INDULIN AT with a 131 density of ca. 1.25 g/cm³ was dried prior to use at 100°C for 12 hours in air. After 132 drying, it was applied as a filler in the rubber mix. Sulfur with a density of about 1.8-133 2.1 g/cm³ and a molecular weight of 32.1 g/mol was also purchased from Torimex 134 Chemicals Ltd Sp. z o. o. Other ingredients, listed below, were purchased from 135 BrenntagPolska Sp. z o. o., Kędzierzyn - Koźle, Poland: 136

- stearic acid with a density of about 0.85 0.99 g/cm³ and a molecular weight of 284.5 g/mol,
- zinc oxide with a density of about 5.6 g/cm³ and a molecular weight of 81.4 g/mol,
- stabilizer AR (fenyl-β-naphthylamine): density 1.16 g/cm³, molecular weight 219.3 g/mol,

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143 accelerator T (tetramethylthiuram disulfide): density 1.5 g/cm³, molecular weight 240 g/mol. 144

The glycerolysate with a number average molecular weight of about 902.4 g/mol and 145 a hydroxyl number of about 186.5 mg KOH/g was used as a plasticizer. This 146 component was produced at the Department of Polymer Technology, Gdańsk 147 University of Technology. 148

2.2. THE PREPARATION OF COMPOSITES 149

Five different composite specimens were prepared. Four composite samples 150 contained different lignin contents, namely, 5, 10, 20 and 40 phr, and were coded 151 WLI5G, WLI10G, WLI20G and WLI40G, respectively. The reference sample without 152 lignin was also prepared (WLI0G). Before mixing, the natural rubber was annealed in 153 154 air to improve the mastication process. The natural rubber was subjected to the mixing process in a BUZULUK open-roll machine (Dattaand Głowińska, 2011). Then 155 the ingredients were added in the quantities shown in Table 1. The friction ratio 156 157 between the two rolls was about 1.1:1. All composites were vulcanized at a temperature of 146 °C with the use of hydraulic press produced by ZUP Nysa. The 158 applied pressure was ca. 5 MPa. The vulcanization temperature was chosen in 159 accordance with the reports by Jacob et. al. (Jacobet al., 2004) and Chonkaew et. al. 160 (Chonkaew et al., 2010). 161

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 Table 1 Composition of lignin-filled natural rubber-based composites.

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2.3. CHARACTERIZATION OF THE COMPOSITES

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Fourier Transform Infrared Spectroscopy was used to obtain the spectra of the samples of five composites, pure lignin, natural rubber, and glycerolysate. The measurements were carried out using a Nicolet 8700 FTIR spectrometer (Thermo Electron Corporation) with the use of ATR technique. The resolution was 4 cm⁻¹. Sixty-four scans in the wavenumber range from 4500 to 500 cm⁻¹ were taken.

Scanning Electron Microscopy was used to characterize the cross-section morphology of the composites. The study was performed with the use of a Phenom G2 PRO scanning electron microscope (Phenom-World corporation) at the accelerating voltage of ca. 5 kV.

Dynamic mechanical tests were carried out with the use of a DMA Q 800 analyzer (TA Instruments). The measurements were performed in accordance with ISO 6721-1, which allowed to obtain the values of storage modulus and tangent delta (damping factor) curves. The specimens with dimensions $30 \times 5 \times 2$ mm were analyzed at a heating rate of ca. 4 °C/min for the temperature range from -100 to 150°C. The tests were performed in air, with a frequency of 1 Hz.

Thermogravimetric analysis allowed to characterize the thermal stability of the prepared composites. The measurements were carried out with the use of a NETZSCH TG 209F3 analyzer. The specimens weighing ca. 5 mg each were analyzed under nitrogen atmosphere. The temperature used ranged from 35 to 600°C at a heating rate of 20 °C/min.

The mechanical properties, such as tensile strength, elongation at break, and permanent elongation after break, were determined with the use of a Zwick/Roell Z020 universal testing machine. The tests were performed in accordance with ISO 37, with the crosshead speed set to 300 mm/min and the 20 kN load cell. The

dumbbell-shaped specimens of all composites were tested. The obtained results arereported as average values calculated from three samples.

The hardness measurements were performed with a Shore type A Durometer (Zwick/Roell). The circular specimens with a thickness of 6 mm were tested in accordance with the standard ISO 868. The presented results are the mean values of hardness based on ten independent measurements.

A Schob machine was used to analyze the rebound resilience of the obtained materials. Tests were performed on the circular, 6-mm thick samples in accordance with ISO 4662. The mean values calculated from ten measurements are reported for all the composites.

The density of the produced materials was determined by using an electronic analytical balance equipped with a kit for measuring the density of solids. During the single test the sample was weighed in air and in the liquid of known density, namely, methanol with a density of ca. 0.790 g/cm³. All measurements were performed at a temperature of 23°C in accordance with ISO 2781. The presented results are the mean values calculated from three independent measurements.

The abrasion resistance was investigated with the use of a Schopper-Schlobbach instrument. All specimens were circles of diameter 17 mm. Tests were performed on three samples of each material in accordance with ISO 4649. The average values were calculated from three independent measurements. The abrasiveness V (cm³) was defined as a volume loss, and determined from equation (1):

213
$$V = (m_0 - m_1) * m_t / (\rho * \Delta m_w)$$

where: $m_0(g)$ and $m_1(g)$ are the sample mass before and after the test, respectively; m_t (g) is a theoretical loss of weight of the reference mixture (assumed value of 0.2 g); ρ (g/cm³) is the density of tested sample; and Δm_w (g) is the average weight loss of reference mixture with the known abrasiveness (assumed value of 0.1095 g).

219 The swelling parameters, such as swelling index, volume fraction of rubber network, molecular weight between crosslinks, and the crosslink density, were 220 221 determined by employing the equilibrium swelling method. The tests were carried out 222 according to the descriptions presented in the literature (Abdelmouleh et al., 2007; Bahl et al., 2014; Gregorová et al., 2006; Kosikova et al., 2007). The rectangular 223 specimens with the average dimensions 15 x 15 x 2 mm were immersed in pure 224 toluene (purchased from POCH, Gliwice, Poland). The measurements were carried 225 out at room temperature for 7 days (168 hours). In this time period, samples were 226 227 taken out from the immersion liquid to determine the weight change. The samples were dried on paper for one minute in air, then weighted and placed again in toluene. 228 After obtaining the equilibrium swelling state, the samples were dried in air for 7 days 229 and then subjected to annealing at a temperature of 80°C for 3 hours (Riyajan, 2015; 230 Stelescu et al., 2014). The swelling ratio was calculated according to the following 231 equation (2): 232

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(2)

where: $m_b(g)$ and m_a (g) are the mean weights of analyzed samples before and after swelling in the immersion liquid, respectively. In the next step of the procedure, the volume fraction of rubber network, $V_{fr}(-)$, was calculated from equation (3):

239

240
$$V_{fr} = V_r / (V_r + V_s) = (m_r / \rho_{rr}) / (m_r / \rho_{rr} + m_s / \rho_s)$$
 (3)

241

where: V_r (cm³) and V_s (cm³) are the mean volumes of rubber and solvent in the swollen sample, respectively; $m_r(g)$ and m_s (g) are the mean weights of rubber and solvent in the analyzed samples after swelling in the immersion liquid, respectively; $\rho_{rr}(g/cm^3)$ and ρ_s (g/cm³) are the respective densities of rubber (0.9125 g/cm³ for natural rubber) and solvent (0.867 g/cm³ for toluene) in the composites.

The average molecular weight between crosslinks, M_c (g/mol) was determined by using the Flory–Rehner theory according to the following equation (4):

249

$$M_{c} = \frac{-\rho_{r} \times V_{s} \times \left(V_{fr}^{\frac{1}{2}} - \frac{1}{2}V_{fr}\right)}{\ln(1 - V_{fr}) + V_{fr} + \left(\chi \times V_{fr}^{2}\right)}$$
(4)

251

250

where: $\rho_r(g/cm^3)$ means the density of lignin-rubber composites; V_s is the molar volume of solvent (106.52 cm³/mol for toluene); and χ is the Huggins parameter, defining the interaction between the polymer and solvent (0.38 for natural rubbertoluene). 256 The crosslink density, v (mol/cm³),was determined from equation (5): 257 258 $v = \rho_r / M_c$ (5)

259

where: $\rho_r(g/cm^3)$ means the polymer density (lignin-rubber composite), and M_C (g/mol) is the average molecular weight of polymer.

262 3. RESULTS AND DISCUSSION

263 3.1. FOURIER TRANSFORM INFRARED (FTIR) SPECTROSCOPY

Fourier Transform Infrared analysis was used to investigate the chemical structure of the prepared lignin/rubber composites, pure lignin, natural rubber and glycerolysate used. The FTIR spectra of all composite samples (Figure 1a) demonstrated similar profiles.

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Figure 1 a) FTIR spectra of the composites, pure lignin, natural rubber and glycerolysate; b) FTIR spectra of the obtained specimens for the wavenumbers ranging from 1800 to 800 cm⁻¹.

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273 More differences were associated with the varying intensity of characteristic 274 peaks. For the samples without the natural rubber, the characteristic wide vibration 275 was present in all spectra in the wavelength range between 3570 and 3170 cm⁻¹, 276 which was attributable to the stretching vibrations of hydroxyl groups derived from lignin and glycerolysate (Kubačková et al., 2013). The strong peaks with the highest
intensity at 2960, 2918 and 2850 cm⁻¹ correspond to the stretching vibrations of the
CH₃, CH₂, CH groups from cis-1,4-polyisoprene macromolecules and glycerolysate
(Riyajan, 2015) , however, the lignin particles also revealed a small shift in this
wavenumber range.

For a better visibility, the FTIR spectra of samples in the wavelength range 282 from 1800 to 800 cm⁻¹ were presented in Figure 1 b. The intensive peak at 1596 cm⁻¹ 283 is attributable to the symmetric aromatic skeletal vibration indicated by lignin 284 macromolecules (Faruket al., 2016c). Other strong bands at 1515 cm⁻¹ are related to 285 the asymetric aryl ring stretching corresponding to lignin (Faruket al., 2016c). The 286 absorption at ca. 1445 cm⁻¹ was assigned to the deformation vibrations of methyl 287 groups -CH is related to the lignin aromatic rings, and also to the natural rubber 288 chains (Josephet al., 2010). The band near 1377 cm⁻¹ is attributable to the 289 asymmetric vibrations of the methyl -CH groups derived from the natural rubber 290 chain as well as glycerolysate (Dattaand Włoch, 2015). The absorption at ca. 1266 291 cm⁻¹ was assigned to the stretching vibrations of the -C-O groups present in the 292 phenolic rings of lignin. The peaks at ca. 1127 and 1030 cm⁻¹ were attributed to the 293 aromatic –CH groups in-plane deformation stretching originating from lignin (Faruket 294 al., 2016c). In the case of glycerolysate, the most intensive peak at a wavenumber of 295 1094 cm⁻¹ was attributable to the stretching vibration of the C-O group. The peak 296 near the wavelength of ca. 839 cm⁻¹ was related to the variations of -CH groups in 297 298 the rubber chains (Linetal., 2015).

299 It is noteworthy that the intensity of characteristic peaks did not increase with 300 the increasing amount of lignin in the composites. That can be explained by the formation of lignin agglomerates, which occurs as the amount of lignin increases. The
 accumulation of lignin particles results in the disturbance of the spectrum.

The interaction between the lignin units and the natural rubber chain can be 303 explained based on the FTIR results. Jiang et al. (Jianget al., 2013) described the 304 305 interaction between lignin and the poly(diallyldimethylammonium chloride) (PDADMAC) chain by comparing the shifts in the FTIR peaks corresponding to the 306 stretching vibration of the -CH groups due to the presence of double bonds. The 307 308 authors also indicated that lignin interacts with the rubber chains in the same way because of the similarity between the chain skeletons of NR and PDADMAC. In 309 Figure 1 b, the peaks at 1515 and 1596 cm⁻¹ in the spectrum of pure lignin shift to the 310 respective locations at 1545 and 1600 cm⁻¹ in the spectra of composites. A small shift 311 from a wavenumber of 1445 cm⁻¹ to 1450 cm⁻¹ is also visible, which is associated with 312 the vibration of the natural rubber chain. This little shift indicates a noncovalent 313 interaction between the NR chain and lignin due to the adsorption of NR onto lignin. 314 The similar results were presented by Pillai and Renneckar(Pillaiand Renneckar, 315 2009), and Yang and co-workers (Yanget al., 2005). 316

317 3.2. SCANNING ELECTRON MICROSCOPY (SEM)

The cross-sections of specimens were assessed by the scanning electron microscopy in order to characterize the lignin dispersion in the composites. In addition, the SEM micrographs revealed the interaction between the lignin powder and the natural rubber matrix. Figures 2a and 2b show the specimens without lignin and those containing 5 phr of lignin, respectively. It has been demonstrated that the dispersion of this lowest quantity of lignin was satisfactory, without any visible agglomerates. Other specimens, i.e. WLI10G, WLI20G and WLI40G showed the

accumulation of lignin particles. The lignin agglomerates were of irregular shape with 325 approximate dimensions from several to several tens of micrometers (Jianget al., 326 2015). This finding explains the formation of hydrogen bonds between the lignin 327 particles. It also indicates that the aggregation of lignin is unavoidable in the natural 328 rubber matrix without additional modifications (Jianget al., 2013). Moreover, it is well 329 known that the composites with highly dispersed filler in the rubber can bear higher 330 stress in comparison to the composites containing the filler agglomerates. The 331 concentration of lignin particles in the obtained composites results in a decrease in 332 resistance (Jianget al., 2015). It is noteworthy that the observed agglomerates did not 333 334 expand with increasing lignin content in the composites, however, their number increased. This can be explained by the glycerolysate content, which can suppress 335 the accumulation of lignin particles into larger agglomerates despite the lignin 336 polydispersity, and its propensity to form larger agglomerates with increasing lignin 337 concentration. 338

339

Figure 2SEM images of a) reference samples, WLI0G; b) samples containing 5 phr
of lignin, WLI5G; c) samples with 10 phr of lignin, WLI10G; d) samples with 20 phr of
lignin, WLI20G; and e) samples with 40 phr of lignin, WLI40G.

343

344 3.3. DYNAMIC MECHANICAL ANALYSIS (DMA)

The dynamic mechanical behavior of the lignin-rubber composites was investigated with the use of dynamic mechanical analysis. The variation of the storage modulus logarithm, log E' versus temperature, and variation of tangent delta, Tan δ versus temperature were determined (Figures 3 and 4).

Figure 3 The logarithm of storage modulus (Log E') of the lignin-rubber composites
plotted versus temperature.

352

The characteristics of the Log E' vs. temperature curve are typical for thermoplastic 353 polymers with high molecular weights. The relationship between the storage modulus 354 355 and temperature for the temperature range from -100 to ca. -50°C was linear. The underlying cause of this behavior is the presence of natural rubber which, at the 356 aforementioned temperature range, occurs is the glassy state, and does not generate 357 358 the reinforcing effect with the filler (Braset al., 2010). After achieving the temperature of ca. -50°C, a sharp decreasing trend was observed in the curve. This phenomenon 359 is related to the alfa transition temperature of the natural rubber matrix. The midpoint 360 of the decreasing curve at the temperature range from ca. -50 to ca. 0°C indicates 361 the position of the glass transition temperature, T_g(Chonkaew et al., 2010; 362 363 Geethamma et al., 2005). This point is closely associated with the relaxation of the long chain sequences of natural rubber. Cooperative motion of macromolecules 364 induce the dissipation of energy at the temperature of maximum tan δ (Figure 4) 365 (Abdelmouleh et al., 2007). The curve characteristics at around 0°C are related to the 366 melting of the remaining crystalline regions in the macromolecule matrix (Gopalan 367 Nair and Dufresne, 2003). 368

369

371

Figure 4 Tan δ as a function of temperature of the lignin-rubber composites.

The addition of the various amounts of lignin to the composites resulted in a 372 373 decrease in the alfa transition temperature. The lowest value, i.e. ca. -44 °C was determined for the specimens containing 10 and 40 phr of lignin (Table 2). These 374 results can be explained by impaired chain mobility due to the lignin addition, which is 375 an obstacle for generating significant reinforcement in the composite (Chonkaew et 376 al., 2010). The highest tan δ peak was observed in the reference sample. With 377 increasing lignin content, the peak intensity decreased. Because the filler particles 378 were surrounded by the rubber matrix and the hydrogen bonds formed, the 379 movement of macromolecule chains was reduced, which resulted in the decreased 380 381 intensity of the peaks (Kargarzadeh et al., 2015).

382

Table 2 Alfa transition temperature [°C] of the lignin-rubber composites.

384

385 3.4. THERMOGRAVIMETRIC ANALYSIS (TGA)

The thermal stability of the composites, reference sample and pure lignin was investigated with the use of thermogravimetric analysis. The measurements allowed for plotting the thermogravimetric (TGA; Figure 5) and differential thermogravimetric (DTG; Figure 6) graphs.

390

Figure 5 Thermogravimetric (TGA) curves for pure lignin, reference samples, and
 lignin-rubber composites with different lignin content.

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393

Figure 6Differential thermogravimetric (DTG) curves for pure lignin, reference
 samples, and lignin-rubber composites with different lignin content.

396

Both aforementioned figures present one step of thermal decomposition for all 397 samples. The highest rate of weight loss was observed in the sample of pure lignin. 398 The samples of lignin-rubber composites showed an increase in the weight loss rate 399 with increasing lignin content. In comparison to the reference sample, the composites 400 containing higher lignin levels lost weight at the lower temperature (a 5% weight loss 401 402 temperature). This was caused by the effect of lignin on the thermal properties of the prepared composites. Despite the aforementioned finding, the maximum rate of mass 403 loss for all the composites occurred at the same temperature (390°C), except for the 404 405 reference sample and pure lignin. For the latter two, the maximum rate of weight loss was observed at the lower temperature. In the case of temperatures of 50% and 80% 406 weight loss, the samples displayed a similar range of maximum rate. Table 3 407 presents the thermal degradation characteristics for all the analyzed samples. 408

409

Table 3 Thermal degradation characteristics of the lignin-rubber composites and pure lignin, where $T_{5\%}$ is a temperature of 5% weight loss, $T_{50\%}$ is a temperature of 50% weight loss, $T_{80\%}$ is a temperature of 80% weight loss, and T_{max} is a temperature of the maximum rate of weight loss. Residue at 600°C [%].

414

The residual weight after TGA analysis (at 600°C) increased with increasing lignin content. This resulted from the presence of the highest lignin content. The

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residue at 600°C comprised of ash, which consisted of organic and inorganic impurities. The impurities play an important role because they influence the thermal properties of lignin. The content of contaminants reduced the possibility of lignin softening in the composites, and decreased the quality of the filler. Moreover, due to the presence of impurities, thermal motion is hindered thereby preventing the thermal processing (Hu,2002; Kadla et al., 2002).

423 **3.5. MECHANICAL PROPERTIES**

The mechanical properties of the composites, such as tensile strength, elongation at break, and permanent elongation after break, were investigated. In order to characterize the prepared materials, their hardness, rebound resilience, density and abrasiveness were measured. Figure 7 shows the effect of varying amounts of lignin on the tensile strength of the composites.

429

430 **Figure 7**Stress-strain curve of the rubber composites.

431

It was found that the tensile properties of the samples decreased with 432 increasing lignin content. The composites with a low amount of filler (WLI5G and 433 WLI10G) showed a decrease in the elongation at break and tensile strength. The 434 specimens containing 10 phr of lignin displayed the characteristics which were similar 435 to those of the reference sample, however, they had lower tensile strength. In the 436 437 case of the samples containing 20 and 40 phr lignin, they were characterized by the higher (WLI20G) or similar (WLI40G) elongation at break, and lower tensile strength 438 than the reference sample. The lignin admixture in the composites caused a 439

decrease in the deformation resistance of rubber chains. Yu et al. (Yuetal., 2015)
obtained similar results. Table 4 presents the results of tensile measurements.

442

Table 4The effect of lignin content on the tensile properties of lignin/natural rubbercomposites.

445

The interaction between the rubber chain and the filler particles hindered the return of macromolecules to the pre-test state. For this reason, the lignin rubber composites showed an increase in the permanent elongation after break. The specimen containing 5 phr of lignin was an exception because its permanent elongation decreased.

In Table 5 the results of hardness, resilience, density and abrasiveness
 measurements are presented.

453

Table 5 The results of hardness, resilience, density and abrasiveness tests for theanalyzed composites.

456

For the samples with 5 and 10 phr of lignin added, hardness of the analyzed composites slightly increased. In the case of the remaining filled samples, the hardness values decreased. It is known that the hardness of composites is closely related to the crosslink density (Stelescu et al., 2014). The maximum hardness at ca. 45 °Sh A was measured in the sample containing 5 phr of lignin. A decrease in

462 hardness with increasing lignin content was correlated with the reduced crosslink 463 density. The differences were observed in the case of the composite samples 464 containing 5 phr of lignin. Those samples were characterized by higher hardness 465 than the reference sample and other specimens. Thereby, the crosslink density of 466 this particular composite displayed the greatest value.

Rebound resilience is also correlated with the crosslink density of composites. It was clearly demonstrated in this study that with increasing lignin content the crosslink density decreased (except for WLI5G), and thus the elasticity was also reduced which led to more rigid composites. Stelescu et al. obtained similar results (Stelescu et al., 2010).

472 Density is strictly related to the lignin amount, therefore, the highest density 473 was measured in sample WLI40G, while the lowest, in WLI0G.

The composite with a 5 phr of lignin admixture showed the lowest abrasiveness compared to the remaining composites. This finding is clearly connected to the crosslink density of this specimen, which was the highest. For the remaining composites, the abrasion resistance decreased with increasing lignin content.

479 3.6. EQUILIBRIUM SWELLING PROPERTIES

The effect of lignin content on the composite network structures was investigated via the equilibrium swelling test. The following parameters were determined: the volume fraction of rubber, molecular weight between the crosslinks, crosslink density, and swelling ratio all of composites. Table 6 presents the results of those measurements.

Table 6 The results of equilibrium swelling measurements performed on the ligninrubber composites.

488

With increasing lignin content the swelling ratio of the composites decreased 489 (Table 6) in comparison to the reference specimen. The swelling ratio increased only 490 491 in the case of the composites with the highest lignin content. M. Jacob et al. (Jacobet al., 2004) pointed out that the solvent diffusion mechanism in the rubber matrix is 492 correlated with the polymer's ability to form voids suitable for the solvent uptake. With 493 494 increasing lignin content in the composites, the number of voids in the rubber matrix decreased, locking the pathways for the solvent, and thus the solvent uptake also 495 decreased. The highest value of the swelling ratio of the specimen with a 40 phr 496 lignin content is related to the most numerous formation of agglomerates with the 497 highest volume. The aforementioned amount of lignin in the composite is too high for 498 499 the successful hydrogen bond formation between the lignin particles and the natural rubber matrix. As a result, the lignin particles can leach out from the samples during 500 the swelling ratio measurements, which makes the high solvent uptake possible. The 501 502 swelling ratio is related to the crosslink density. (Jacobet al., 2004; Park and Cho, 2003) suggested that the maximum interaction between the rubber chain and filler, 503 and the lowest crosslink density are characteristic for the composites with the highest 504 admixture of lignin. The obtained results showed that the lignin content caused a 505 decrease in the crosslink density (Table 6). Lignin in the natural rubber vulcanizates 506 lowers the number of polysulfidic crosslinks between the rubber chains if the disulfidic 507 and monosulfidic crosslinks remain constant (Faruket al., 2016b). The decreased 508

amount of polysulfides lowers the hardness, resilience, abrasion resistance and 509 tensile properties (Stelescu et al., 2010). The lignin particles mask some of the sites 510 on the natural rubber molecules, that otherwise would have been available for 511 crosslinking (Kumaran et al., 1978), and cause a decrease in the network density. 512 Due to the occurrence of masked crosslinking sites, the distance between the 513 network nodes increases. The larger distance between the networks leads to the 514 polymer chain tangling. The increased molecular weight between crosslinks (see 515 Table 6) is a proof of the increased complexity of natural rubber macromolecules 516 between the network nodes. 517

During the equilibrium swelling measurements the values of weight loss were 518 recorded over time (Figure 8). The samples containing 5 and 10 phr of lignin 519 displayed a lower weight change than the reference samples. The sample with a 5 520 phr lignin content showed the lowest weight loss. The two composites with the 521 highest lignin contents displayed a higher weight loss than the reference samples. 522 The performed measurement allowed for concluding that a low lignin content in the 523 composites containing glycerolysate as a plasticizer, resulted in a decrease in the 524 525 solvent (toluene) absorption.

526

527 **Figure 8**Weight loss during the swelling measurements performed on the lignin-528 rubber composites.

529

530 4. CONCLUSIONS

The softwood-lignin/natural rubber composites were successfully prepared 531 532 with the use of a novel plasticizer. Nowadays, the plasticizers mainly originate from the petrochemical resources. Glycerolysate used in this work is the product of the 533 chemical recycling of polyurethanes. The results of the presented research 534 demonstrated that the mechanical and thermal properties of softwood-lignin/natural 535 rubber composites obtained with the use of glycerolysate as a plasticizer are similar 536 to those of the lignin-containing composites prepared with other commercially 537 available plasticizing agents. In addition, the data showed that the composite 538 containing 5 phr of lignin displayed the best properties. Specimen WLI5G had the 539 540 highest values of hardness and abrasiveness, and superior tensile properties. The Scanning Electron Microscopy (SEM) confirmed a positive interaction between the 541 rubber chains and the lignin particles. The SEM images of composite WLI5G showed 542 543 good dispersion of lignin, without the lignin agglomerates. The prepared four composite formulas with increasing lignin content allowed the determination of 544 maximum lignin content in the composite that would not alter the material's positive 545 properties. The composite containing the highest amount of lignin (WLI40G) had the 546 biggest and most numerous agglomeratesand, consequently, the worst mechanical 547 548 (the lowest values of tensile strength, elongation after break, etc.) and thermal properties (e.g. the fastest weight loss). This observation has been confirmed by the 549 results of swelling measurements, where sample WLI40G displayed the highest 550 551 swelling ratio. The application of lignin that is the product of natural origin, and the use of the product of the chemical recycling of polyurethanes makes the described 552 procedure eco-friendly. Its economic advantages should also be considered. 553 Glycerolysate, used as a plasticizing agent, is less expensive than the commonly 554 employed plasticizers derived from petrochemicals. In summary, this work is a 555

556 promising development in the field of natural product utilization, and the application of 557 the products of chemical recycling in the natural rubber matrix.

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562

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712 Figure Captions

Figure 1 a) FTIR spectra of the composites, pure lignin, natural rubber and glycerolysate; b) FTIR spectra of the obtained specimens for the wavenumbers ranging from 1800 to 800 cm⁻¹.

Figure 2SEM images of a) reference samples, WLI0G; b) samples containing 5 phr
of lignin, WLI5G; c) samples with 10 phr of lignin, WLI10G; d) samples with 20 phr of
lignin, WLI20G; and e) samples with 40 phr of lignin, WLI40G.

Figure 3 The logarithm of storage modulus (Log E') of the lignin-rubber composites
plotted versus temperature.

Figure 4 Tan δ as a function of temperature of the lignin-rubber composites.

Figure 5 Thermogravimetric (TGA) curves for pure lignin, reference samples, and
lignin-rubber composites with different lignin content.

Figure 6 Differential thermogravimetric (DTG) curves for pure lignin, reference samples, and lignin-rubber composites with different lignin content.

726 **Figure 7** Stress-strain curve of the rubber composites.

Figure 8Weight loss during the swelling measurements performed on the lignin-rubber composites.

COMPONENT	QUANTITIES OF INGREDIENTS (phr)				
	WLI0G	WLI5G	WLI10G	WLI20G	WLI40G
Natural rubber	100	100	100	100	100
Stearic acid	3	3	3	3	3
Zinc oxide	5	5	5	5	5
Stabilizer AR	1.5	1.5	1.5	1.5	1.5
Accelerator T	0.5	0.5	0.5	0.5	0.5
Glycerolysate	2	2	2	2	2
Lignin (INDULIN AT)	0	5	10	20	40
Sulphur	3	3	3	3	3

Table 1 Composition of lignin filled natural rubber-based composites.

Composite	Alfa transition temperature		
	Τ _α [°C]		
WLI0G	-38		
WLI5G	-40		
WLI10G	-44		
WLI20G	-41		
WLI40G	-43		

Table 2 Alfa transition temperature [°C] of the lignin-rubber composites.

Table 3 Thermal degradation characteristics of the lignin-rubber composites and pure lignin, where $T_{5\%}$ is a temperature of 5% weight loss, $T_{50\%}$ is a temperature of 50% weight loss, $T_{80\%}$ is a temperature of 80% weight loss, and T_{max} is a temperature of the maximum rate of weight loss. Residue at 600°C [%].

	Thermal degradation characteristic [°C]				
Sample	T _{5%}	T 50%	T80%	T _{max}	Residue at
					600°C [%]
WLI0G	308	393	430	386	6
WLI5G	303	396	433	390	8
WLI10G	298	394	431	390	8
WLI20G	290	395	438	390	11
WLI40G	270	395	447	390	15
lignin	267	567	-	382	48

Table 4The effect of lignin content on the tensile properties of lignin/natural rubbercomposites.

COMPOSITE	POSITE Tensile strength E		Permanent elongation
	[MPa]	break [%]	after break [%]
WLI0G	16.25 <u>+</u> 1.1	717 <u>+</u> 12	18.1 ± 1.0
WLI5G	15.16 <u>+</u> 0.4	626 <u>+</u> 15	15.5 <u>+</u> 1.1
WLI10G	14.22 <u>+</u> 1.6	706 <u>+</u> 15	18.4 <u>+</u> 0.4
WLI20G	14.39 ± 0.7	733 ± 3	18.2 <u>+</u> 0.5
WLI40G	8.39 ± 0.7	682 <u>+</u> 26	21.9 ± 0.5

Table 5 The results of hardness, resilience, density and abrasiveness tests for theanalyzed composites.

COMPOSITE	HARDNESS	RESILIENCE	DENSITY [g/cm ³]	ABRASIVENESS
	[ºSh A]	[%]		[cm ³]
WLI0G	43.6 ± 0.3	58.7 ± 2.0	0.9724 ± 0.0003	0.0899 ± 0.0080
WLI5G	45.5 ± 0.5	57.6 ± 0.8	0.9846 ± 0.0021	0.0847 ± 0.0170
WLI10G	44.3 <u>+</u> 0.5	57.3 ± 0.9	0.9922 ± 0.0006	0.1179 ± 0.0143
WLI20G	41.2 <u>+</u> 0.3	52.0 ± 1.1	1.0099 ± 0.0004	0.1469 ± 0.0221
WLI40G	38.3 ± 0.4	38.7 ± 0.7	1.0420 ± 0.0009	0.1999 <u>+</u> 0.0487

Table 6 The results of equilibrium swelling measurements performed on the lignin-rubber composites.

COMPOSITE	SWELLING	VOLUME	MOLECULAR	CROSSLINK
	RATIO	FRACTION	WEIGHT	DENSITY
	SR [%]	V _{fr} [-]	BETWEEN	v x 10 ⁻³
		• " []	CROSSLINKS	[mol/cm ³]
			M _C [g/mol]	
WLI0G	290.8 ± 3.1	0.2374 ± 0.0026	191.6 ± 2.1	5.01 ± 0.05
WLI5G	280.9 ± 0.3	0.2371 ± 0.0003	194.2 ± 0.2	5.07 ± 0.01
WLI10G	278.4 <u>+</u> 0.9	0.2313 ± 0.0007	200.4 ± 0.6	4.95 ± 0.01
WLI20G	279.9 <u>+</u> 1.6	0.2156 ± 0.0011	218.2 ± 1.0	4.62 ± 0.02
WLI40G	308.2 ± 2.9	0.1771 ± 0.0018	268.6 ± 2.6	3.88 ± 0.04

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758

759 Figure 2



761 Figure 3



762

763

Figure 4



765 Figure 5





769 Figure 7





