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Chopped basalt fibres: A new perspective in reinforcing poly(lactic acid) to produce injection moulded engineering composites from renewable and natural resources

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Abstract. This paper focuses on the reinforcing of Poly(lactic acid) with chopped basalt fibres by using silane treated and untreated basalt fibres. Composite materials with 5–10–15–20–30–40 wt% basalt fibre contents were prepared from silane sized basalt fibres using extrusion, and injection moulding, while composites with 5–10–15 wt% basalt fibre contents were also prepared by using untreated basalt fibres as control. The properties of the injection moulded composites were extensively examined by using quasi-static (tensile, three-point bending) and dynamic mechanical tests (notched and unnotched Charpy impact tests), dynamic mechanical analysis (DMA), differential scanning calorimetry (DSC), heat deflection temperature (HDT) analysis, dimensional stability test, as well as melt flow index (MFI) analysis and scanning electron microscopic (SEM) observations. It was found that silane treated chopped basalt fibres are much more effective in reinforcing Poly(lactic acid) than natural fibres; although basalt fibres are not biodegradable but they are still considered as natural (can be found in nature in the form of volcanic rocks) and biologically inert. It is demonstrated in this paper that by using basalt fibre reinforcement, a renewable and natural resource based composite can be produced by injection moulding with excellent mechanical properties suitable even for engineering applications. Finally it was shown that by using adequate drying of the materials, composites with higher mechanical properties can be achieved compared to literature data.

Keywords: biodegradable polymers, polymer composites, poly(lactic acid), chopped basalt fibres, injection moulding

1. Introduction

In the 21st century, due to the increasing environmental consciousness renewable resource based and inherently biodegradable polymers gain more and more role. These materials can be fully synthetized from biomass or previously not utilized agricultural feedstock and during their biodegradation harmless materials like water, carbon-dioxide and humus are produced, where the latter can be used to grow agricultural plants, so thus it can be easily seen that these materials fit into the life cycle of nature and also fit into the train of thoughts of sustainable development [1–3]. It is also believed that in the near future biodegradable polymers will be used for producing plastic products with short term applications, however nowadays in most cases for these applications still unnecessarily durable petrol-based plastics are used, what causes more and more waste management problems due to their not complete recyclability.

One of the most promising renewable resources to produce biopolymers is starch. Starch can be found all over the world in abundant amounts, because starch containing agricultural plants like corn, wheat, potato are grown extensively in a lot of countries. In the last few decades much research has

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been made to make starch processable with conventional plastic processing equipment, and as a result thermoplastic starch (TPS) was developed and extensively researched [4–6]. Although TPS is cheap, its numerous drawbacks like low mechanical properties, high shrinkage, and water solubility retarded its widespread usage. Nowadays TPS together with Poly(ε-caprolactone) (PCL) is used as a compound and commercialised under trade name Mater-Bi (Novamont, Italy). Another possible way to utilise starch is its fermentation in the presence of sugar into lactic acid, what can be further polymerised into Poly(lactic acid) (PLA). In the last decade PLA was in the focus of interest, and it became the most promising biopolymers of all due to its excellent mechanical properties, easy processing by conventional equipment, low shrinkage, reasonable price and harmless biodegradation capability [7–9]. Although PLA is already commercialised and more and more PLA products are on the market, these products are still mainly related to packaging industry, even so the high strength and stiffness of PLA suggest it to be used as the matrix of a durable engineering composite material, but its low heat deflection temperature (HDT) and low impact strength still retards its usage as a high performance composite.

Lots of efforts have been made to further reinforce PLA with typically natural plant fibres [10–16] or even with glass fibres [17] to make a biocomposite for applications with higher demand then packaging. In most cases by adding natural plant fibres to PLA higher increase of mechanical properties occurred when solvent casting [10, 11], pultrusion [12], extrusion followed by compression [13], film-stacking [14, 15], compression followed by injection moulding [16] processes were used compared to the conventional and most productive extrusion followed by injection moulding process [17, 18]. This is probably due to the reduced length of the natural fibres in the latter technology. Furthermore, the increase of HDT due to the incorporation of various fibres is only a few times experienced. Iwatake et al. [10] demonstrated that by using solvent casting method and micro-fibrillated cellulose (MFC) fibres a cellulose fibre network interconnected by hydrogen bonds was developed which resisted the applied stress independently of the softening of PLA during dynamic mechanical analysis (DMA) tests. By using the above mentioned processes to increase HDT and impact strength it is crucial to have strong connection between the fibres and the PLA, and the adequate quantity and quality (according to stiffness and length) of reinforcing fibres is also important, which will prevent the well-known high stiffness loss of amorphous PLA products above glass transition temperature ($T_g$). However, by adding natural plant fibres to reinforce PLA, the real breakthrough did not occur because in order to gain significant reinforcing effect especially in the desired impact strength low productivity processes like solvent casting or film stacking method have to be used but to commercialise PLA based composites and make complicated 3D shaped, and accurate parts highly productive production technologies like injection moulding should be applied.

Besides natural fibres, a promising alternative may be the usage of basalt fibres, which is a novel reinforcement for composites. Although basalt fibres are not biodegradable but still considered as natural, because they can be produced by using basalt (volcanic) rocks, which can be found in nature and virtually in every country around the globe. Moreover, basalt is biologically inert, and the weathering of basalt rocks increase the mineral content of soil which further strengthens its natural character. Basalt fibres are typically produced by two diferent technologies: Junkers method and spinneret technology [19]. The shorter basalt fibres are prepared by the Junkers method and the continuous fibres are prepared by the spinneret technology [20]. Its mechanical properties and chemical composition is similar to the glass fibre, and it can be used between −200 and 600°C without the significant loss of mechanical properties [21, 22]. Basalt fibres are biologically inert as well as environmentally friendly and can be used in very aggressive environments too [23, 24]. The surface of the basalt fibres can be modified easily, thus they can be used as reinforcement fibres for composites [25, 26]. Deák et al. [27] investigated the influence of different coupling agents on the mechanical and thermomechanical properties of basalt fibre reinforced Polyamide 6 (PA6). The tensile, flexural and impact strengths were improved in the highest level by the type of 3-glycidoxypropyltrimethoxysilane sizing. The suitability of that sizing material in a wide temperature range was proven by DMA tests. Basalt fibres seem to be a good alternative to reinforce PLA, however, the number of papers in basalt
fibre reinforced PLA is very limited [28–30]. Liu et al. [28] demonstrated that basalt fibres with the adequate sizing can improve tensile, flexural and impact strength of PLA more significant as plant fibres. It was also found that by using 20 wt% of basalt fibres and 20 wt% of ethylene-acylate-glycidyl methacrylate copolymer (EAGMA) the impact strength can be further increased up to 34 kJ/m² from the original value of PLA of 19 kJ/m² (unnotched Charpy test results). Finally the authors stated that the fibres are strongly bonded to the PLA because no gap was found at the fibre-matrix interface. Kurniawan et al. [29] used atmospheric pressure glow discharge plasma polymerization as surface treatment of basalt fibres and analysed the adhesion between the fibres and PLA in hot pressed composites. It was found that by increasing the plasma polymerisation time the tensile strength of the composites increased compared to PLA when the reaction time was higher than 3 minutes. Although the plasma polymerised fibres seem to be well wetted by the PLA, the tensile strength was lower compared to results from Liu et al. [28]. Finally Chen et al. [30] successfully produced PLA based scaffolds reinforced with basalt fibres. They proved that the basalt fibres retard degradation rate of the scaffold, thus most likely decrease inflammatory responses caused by acidification. Moreover, basalt fibres did not notably affect osteoblast viability and growth which means that basalt fibre reinforced PLA scaffolds can be potentially used in hard tissue repair.

According to these results basalt fibres seem to be a good selection to reinforce PLA, because PLA based composites with higher mechanical properties can be achieved compared to natural plant fibre reinforcement. However, the widespread analysis of the various mechanical, thermal, flow and dimensional stability properties of injection moulded basalt fibre reinforced PLA based composites can be only partly found in the literature. Thus, this paper focuses on the comprehensive usability analysis of chopped basalt fibres with and without silane treatment as potential reinforcing fibre for injection moulded, renewable and natural resource based PLA composites as high performance, durable engineering materials.

2. Materials, processing and experimental

Semi-crystalline PLA injection moulding grade resin was used for the measurements (type AI1001 from eSUN, Shenzhen, China, with a D-Lactide content of 4%). It was dried at 120°C for 6 hours prior to extrusion. A somewhat high drying temperature was chosen, because according to our previous research [31] simply by using adequate drying, significantly higher adhesion could be developed between the phases, because the residual moisture could induce degradation and it could also weaken the adhesion between the fillers or fibres and the matrix material. Chopped basalt fibres produced with Junkers method from Kameny Vek (Dubna, Russia) were obtained with the company’s own silane treatment type KV-12 developed especially for strong adhesion with polyesters. The average diameter of the fibres was 13 µm, while the initial length was 10 mm. The fibres were also dried along with the PLA resin to remove residual moisture. Some of the basalt fibres were heated to 600°C to remove silane sizing. The PLA pellets and the treated and untreated (removed silane) basalt fibres were dry blended and extruded by using a LabTech Scientific twin screw extruder (Bangkok, Thailand) (screw diameter = 26 mm, L/D = 40) equipped with a 2 mm hole diameter die. A temperature profile of 175–180–185–190°C (from the hopper to the die) was used with a screw rotational speed of 30 rpm. 5–10–15–20–30–40 wt% basalt fibre composites were prepared with silane treatment (abbreviated as 5BS, 10BS, 15BS, 20BS, 30BS, 40BS) and 5–10–15 wt% basalt fibre composites with removed treatment for control (abbreviated as 5BN, 10BN, 15BN). The extrudates were pelletized and annealed (crytallised) prior to injection moulding to avoid processing problems (pellet sticking to the screw) caused by cold crystallisation as reported in our previous research [32]. The pellets were injection moulded with an Arburg Allrounder 370S 700–250 injection moulding machine (Lossburg, Germany) equipped with a 30 mm diameter, L/D = 25 screw. The applied injection moulding parameters can be seen in Table 1.

ISO standard dumbbell, three-point bending and Charpy specimens with a cross-section of 4 × 10 mm, and 80 × 80 mm area, 2 mm thick flat specimens were injection moulded for the tests. Tensile and three-point bending tests were performed by using a Zwick Z020 universal testing machine equipped with Zwick BZ 020/TN2S force measuring cell with a force limit of 4 kN using a cross-head speed of 5 mm/min. Charpy tests were
performed with a Ceast Resil Impactor impact testing machine equipped with a 15 J impact energy hammer and a DAS8000 data collector unit. Both notched (2 mm deep notch) and unnotched specimens were also prepared for the impact tests; 15 and 6.21 J impact energy was used for the unnotched and notched specimens respectively. All tests were executed at room temperature and at a relative humidity of 60%. 6 specimens were tested for each measurement.

Melt flow index (MFI) measurements were made by using a Ceast 7027.0000 type melt flow indexer at 190°C with a load mass of 2.16 kg. Differential Scanning Calorimetry (DSC) was made on a TA Q2000 type calorimeter by using 3–6 mg of the samples. Heat/cool/heat scans were registered from 0 to 180°C with a heating and cooling rate of 5°C/min and nitrogen gas flow was used. Crystallinity was calculated according to the theoretical enthalpy of fusion for 100% crystalline PLA and α [-] is the mass fraction of basalt fibres. Dynamic Mechanical Analysis (DMA) was performed on a TA Q800 tester by using the injection moulded three-point bending specimens and dual cantilever. The dual cantilever was used in order to obtain storage modulus information above glass transition temperature ($T_g$). Amplitude of 20 µm with a span length of 35 mm and a frequency of 1 Hz were used from 0 to 160°C at a heating rate of 2°C/min.

Heat Deflection Temperature (HDT) was measured by using a Ceast 6505/000 HDT analyser. Injection moulded specimens with a cross-section of 4 × 10 mm and support distance of 100 mm were used. The specific load was 0.45 MPa and the heating rate was 2°C/min. The HDT value is obtained when the deflection reached 0.33 mm.

Dimensional stability of the injection moulded, 2 mm thick flat specimens were analysed by placing them into a heated oven and measuring the flow directional dimensional changes due to increasing temperature. The specimens were placed on a flat glass plate so as to get only dimensional changes and no deflection. The oven was heated to 40, 50, 60, 80, 100 and 120°C for 30 minutes and the dimensions of the specimens were recorded after cooling them to room temperature. These dimensions were compared with the dimensions recorded at room temperature prior to heating by using Equation (2):

$$D = \frac{L_b - L_a}{L_b} \cdot 100$$  \hspace{1cm} (2)

where $D$ [%] is the dimensional change, $L_b$ [mm] and $L_a$ [mm] are the in-flow length of the specimens measured before and after the 30 minute heating process respectively.

The deflection of the specimens was also measured by using the flat specimens placed in the same oven, but in a cantilever to be loaded by their own weight.

Finally, scanning electron microscopy (SEM) was performed by a Jeol JSM 6380LA type electron microscope. The fracture surfaces of the tensile specimens were used for the observations. Prior to observation, Au/Pd alloy was sputtered onto the surface to avoid electrostatic charging.

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**Table 1. Injection moulding parameters**

<table>
<thead>
<tr>
<th>Injection moulding parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume</td>
<td>50 cm$^3$ (flat specimen)</td>
</tr>
<tr>
<td></td>
<td>44 cm$^3$ (tensile specimen)</td>
</tr>
<tr>
<td></td>
<td>42 cm$^3$ (bending specimen)</td>
</tr>
<tr>
<td>Injection pressure</td>
<td>Variable according to basalt fibre content</td>
</tr>
<tr>
<td>Switch-over point</td>
<td>12 cm$^3$</td>
</tr>
<tr>
<td>Injection rate</td>
<td>50 cm$^3$/s</td>
</tr>
<tr>
<td>Holding pressure</td>
<td>600 bar</td>
</tr>
<tr>
<td>Holding time</td>
<td>20 s</td>
</tr>
<tr>
<td>Residual cooling time</td>
<td>30 s</td>
</tr>
<tr>
<td>Screw rotational speed</td>
<td>15 m/min</td>
</tr>
<tr>
<td>Backpressure</td>
<td>30 bar</td>
</tr>
<tr>
<td>Decompression volume</td>
<td>5 cm$^3$</td>
</tr>
<tr>
<td>Decompression rate</td>
<td>5 cm$^3$/s</td>
</tr>
<tr>
<td>Temperature of the first zone</td>
<td>165°C</td>
</tr>
<tr>
<td>Temperature of the second zone</td>
<td>175°C</td>
</tr>
<tr>
<td>Temperature of the third zone</td>
<td>180°C</td>
</tr>
<tr>
<td>Temperature of the fourth zone</td>
<td>185°C</td>
</tr>
<tr>
<td>Temperature of the fifth zone</td>
<td>190°C</td>
</tr>
<tr>
<td>Temperature of the mould</td>
<td>20°C</td>
</tr>
</tbody>
</table>
3. Results and discussion

3.1. Melt flow index analysis

Prior to injection moulding, the melt flow index of the basalt fibre reinforced PLA composites was analysed (Figure 1).

It is found that by the incorporation of basalt fibres the MFI values decreased, what was expected, at the same time above 15 wt% the MFI values did not change essentially. This could be explained by the increased MFI of the PLA phase itself which compensated the MFI decreasing effect of the fibres. The increased MFI of PLA was possibly due to increased shear caused by the higher amount of fibres and due to higher residence times caused by lower output of the extruder during melt blending. It is also worth to note that by using silane treated basalt fibres the MFI values decreased more compared to untreated basalt fibre composites. This difference could be related to the adhesion between the fibres and the matrix and thus it suggest better compatibility between silane treated basalt fibres and PLA than between untreated basalt fibres and PLA presumably because the basalt fibres are distributed more uniformly – due to the silane treatment – in the melt, which thus becomes more viscous.

3.2. Mechanical properties

The mechanical properties of the injection moulded specimens were analysed by using quasi-static tensile and three-point bending tests and notched and unnotched Charpy impact tests. The tensile strength and the tensile modulus results can be seen in (Figure 2, Figure 3). Note that standard deviation is presented in all of the diagrams in the entire paper, however, in most cases it is so low, that the markings overlie it.

It is obvious according to Figure 2 and Figure 3 that silane treated basalt fibre composites have superior mechanical properties (higher strength, the same order of magnitude of modulus) compared to the untreated basalt fibre composites, thus by using the silane treated basalt fibres, the composites reached a tensile strength of 123 MPa, a tensile modulus of over 8 GPa at the cost of minimal (0.6%) reduction of strain (measured but not presented). It was found that by increasing basalt fibre content both tensile strength and tensile modulus increased linearly in the 0–30 wt% basalt fibre content range. Moreover, it is also worth to note that there is low difference between the strength and modulus of 30 and 40 wt% silane treated basalt fibre reinforced PLA composites thus compared to 30 wt% basalt fibre content by adding another 10 wt% did not have significant reinforcing effect which suggest that at 40 wt% major fibre brakeage occurred probably caused by the fibre-fibre friction due to the increased amount of fibres.

The three-point bending tests show similar result compared to tensile tests (Figure 4, Figure 5).

The composites reached a flexural strength of 185 MPa and a flexural modulus of over 12 GPa.
Correspondingly to tensile properties, the flexural properties were also linearly related to the basalt fibre content in the range of 0–30 wt% basalt, and the possible fibre breakage can also be observed in the flexural strength of the 40 wt% basalt fibre content composite, as it is almost the same as the flexural strength of the 30 wt% basalt fibre content composite. Finally, the notched (Figure 6) and unnotched (Figure 7) Charpy impact tests revealed that the critically low impact strength of PLA, especially for engineering applications can be significantly increased by the incorporation of basalt fibres.

By incorporating 30 wt% basalt fibres the notched and unnotched Charpy impact strength of PLA of 2.7 and 23.0 kJ/m² increased to 9.3 (by 244%) and 38.3 kJ/m² (by 67%) respectively. Charpy impact test results as well as tensile and bending test results also confirm that above 30 wt% basalt fibre content no significant reinforcing effect can be achieved due to probably increased fibre breakage.

All of the above mentioned mechanical properties increased more or less linearly to the basalt fibre content in the range of 0–30 wt% basalt fibre content range, which is demonstrated by using linear regression. The coefficients for the regression can be found in Table 2.

The mechanical properties of the injection moulded composites are better compared to results from Liu et al. [28] by 12, 25 and 53% according to tensile, flexural and Charpy impact strength respectively at same basalt fibre content (40 wt%) and also better compared to result from Kurniawan et al. [29] by 24 and 143% according to tensile strength and tensile modulus at same basalt fibre content (25 wt% (note that 25 wt% basalt fibre content composite was not fabricated in this research, but due to the highly linear relationship, the properties were determined by interpolation)). The reason for the better properties can be found in the higher drying temperature.
perature of PLA prior to processing as it is published in our former research [31]. Moreover, these results indicate that the basalt fibre reinforced PLA could be used as durable engineering composite materials, because the developed composite has similar mechanical properties to Polyamide 6 (PA6) reinforced with 15 wt% glass fibre. A further advantage of the developed composite is, that the above mentioned high mechanical properties can not only be achieved by using long fibres and slow technologies like film-stacking to produce only flat-like products (as is was mentioned in the introduction in the case of plant fibre reinforced biocomposites), but by injection moulding, which is the most versatile thermoplastic processing technology to produce complex, 3D shaped engineering parts with high dimensional accuracy and low cycle times.

3.3. Scanning electron microscopy
Scanning electron microscope (SEM) observations were made on the fracture surface of tensile specimens to evaluate the adhesion between the phases (Figure 8, Figure 9). On the fracture surface of the 15 wt% basalt fibre reinforced PLA composites without treatment basalt fibres with smooth surface can be observed, which represents inadequate fibre wetting, and moreover a gap between the matrix and the root of the fibres can also be seen in the untreated basalt fibre reinforced composites, which indicates weak adhesion as it could already be seen in the mechanical properties (Figure 8a). On the fracture surface of the composite with the same amount of basalt fibres but with silane treatment, much stronger adhesion can be observed which is denoted by the absence of gap between the fibres

**Figure 8.** Cross-section of untreated (a) and treated (b), 15 wt% basalt fibre reinforced PLA composite

**Figure 9.** Cross-section of 20 wt% (a) and 40 wt% (b) silane treated basalt fibre reinforced PLA composite
and the matrix, and by the wetting of the fibre by PLA (Figure 8b).
For the 20, 30 and 40 wt% silane treated basalt fibre reinforced composites (Figure 9), also strong adhesion was found.

On the surface of the silane treated fibres a clear PLA layer can be found which represents excellent adhesion. These results are again better compared to literature data (Liu et al. [28]), where the surface of the basalt fibres were smooth and poorly wetted in spite of the reinforcing effect. Good quality wetting was achieved by Kurniawan et al. [29] by using plasma treatment of the basalt fibres, however only basalt fibre reinforced PLA composites were produced with lower mechanical properties compared to our results.

3.4. Differential scanning calorimetry

Differential scanning calorimetry (DSC) was performed by using 3–6 mg samples taken out from the middle of the cross section of the injection moulded specimens. The cooling DSC scans registered at a cooling rate of 5°C/min are presented in Figure 10.

It can be seen that pure PLA only slightly crystallises and the specimens become transparent during cooling from melt what is a common phenomena according to its very slow crystallisation kinetics, especially when using high cooling rate technologies like injection moulding, thus the material of the pure PLA specimens are to be further referred in the paper as amorphous PLA. At the same time the silane treated and not treated basalt fibres both had nucleating effect which is represented by the higher and higher exothermic peak found around 100°C with increasing basalt fibre content. This statement is supported by the quantitative evaluation of the DSC results (Table 3). The crystallinity of the injection moulded samples was calculated by the first heating scan and by using Equation (1), while the crystallinity of the samples cooled at 5°C/min was calculated by using the second heating scan.

It is found that at and above 20 wt% basalt fibre content the nucleating effect of the fibres was enough to develop significant and at the same time the maximum possible crystallinity for PLA, which is around 40–45%. However, this statement is only valid for crystallinity values obtained from the cooling and second heating DSC scan where the cooling and heating rate was set to 5°C/min, while the crystallinity of the injection moulded specimens obtained from the first heating scan was naturally much lower, but still significant; around 19% due to the high cooling rates during processing. Nevertheless, the nucleating effect of basalt fibres suggest that by using further nucleating agents like talc and/or lower cooling rates during processing presumably basalt fibre reinforced semi-crystalline PLA can be produced which has an enormous advantage according to its crystallinity compared to amorphous PLA what is represented in the much higher heat deflec-

Table 3. Crystallinity of basalt fibre reinforced PLA composites

<table>
<thead>
<tr>
<th>Material</th>
<th>Crystallinity of the injection moulded samples [%]</th>
<th>Crystallinity of the samples cooled at 5°C/min [%]</th>
<th>Crystallisation peak temperature [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amorphous PLA</td>
<td>1.1</td>
<td>3.0</td>
<td></td>
</tr>
<tr>
<td>5BN</td>
<td>8.5</td>
<td>21.6</td>
<td>96</td>
</tr>
<tr>
<td>5BS</td>
<td>4.0</td>
<td>25.6</td>
<td>99</td>
</tr>
<tr>
<td>10BN</td>
<td>10.6</td>
<td>20.9</td>
<td>96</td>
</tr>
<tr>
<td>10BS</td>
<td>14.0</td>
<td>24.4</td>
<td>97</td>
</tr>
<tr>
<td>15BN</td>
<td>17.2</td>
<td>23.1</td>
<td>97</td>
</tr>
<tr>
<td>15BS</td>
<td>17.5</td>
<td>21.1</td>
<td>97</td>
</tr>
<tr>
<td>20BS</td>
<td>18.7</td>
<td>42.9</td>
<td>100</td>
</tr>
<tr>
<td>30BS</td>
<td>19.0</td>
<td>44.2</td>
<td>100</td>
</tr>
<tr>
<td>40BS</td>
<td>18.5</td>
<td>44.8</td>
<td>100</td>
</tr>
</tbody>
</table>
tion temperature and thus it would be much more suitable for engineering applications.

### 3.5. Dynamic mechanical analysis

Dynamic mechanical analyses (DMA) were performed to examine the usability of the developed composites in the temperature range of 0–160°C and especially in the critical range above $T_g$. Due to the slow crystallisation of PLA, it remains mostly amorphous during injection moulding, thus the dramatic drop in storage modulus at $T_g$ and the increase in storage modulus caused by cold-crystallisation in the temperature range of 80–120°C can be observed (Figure 11).

Naturally the basalt fibres increase the storage modulus; the silane treated basalt fibres increased it more significantly than the untreated ones due to the better adhesion proved previously by the SEM and mechanical test results. It can also be seen that the nucleating effect of basalt fibres at and above 20 wt% basalt fibre content observed by DSC was not enough to develop significant crystallinity in PLA during injection moulding to avoid the dramatic drop in storage modulus at $T_g$, thus this high storage modulus loss still can be observed for all of the basalt fibre reinforced composites (Figure 12).

For further analysis the DMA curve of semi-crystalline PLA (C-PLA) was also registered (semi-crystalline PLA sample was made by annealing at 120°C for 1 hour to gain maximum possible crystallinity). By using 40 wt% silane treated basalt fibres, the storage modulus above $T_g$ was only reduced to the same order of magnitude as the storage modulus of semi-crystalline PLA (Figure 13), which is important because semi-crystalline PLA has a HDT value of 127°C and considered as heat resistant compared to amorphous PLA, which has a HDT value of 54°C.

However, semi-crystalline PLA still has, but only slightly higher storage modulus in the range of 58–76°C than the 40 wt% basalt reinforced composite. The storage modulus values of PLA, semi-crystalline PLA and the basalt fibre reinforced PLA composites at 40, 120°C and the most critical and lowest storage modulus values between 70–80°C (above $T_g$ but before cold-crystallisation) can be seen in Table 4.

It can be seen that by the incorporation of basalt fibres, the storage modulus increased linearly to reinforcement content and reached an around three-times higher modulus at 40°C and ten-times higher modulus at 120°C. By using linear regression it was found that the storage modulus at 40°C increased with a gradient of 114.0 MPa/basalt wt% content

![Figure 11. Storage modulus of amorphous PLA (PLA), and 5–10–15 wt% basalt fibre reinforced composites with and without silane treatment](image1)

![Figure 12. Storage modulus of amorphous PLA (PLA) and 5–10–15–20–30–40 wt% basalt fibre reinforced composites with silane treatment](image2)

![Figure 13. Storage modulus of amorphous PLA (PLA), semi-crystalline PLA (C-PLA) and 20–30–40 wt% basalt fibre reinforced composites with silane treatment](image3)
(\(R^2 = 0.99\)), while the storage modulus at 120°C increased with a gradient of 32.9 MPa/basalt wt% content (\(R^2 = 0.99\)) according to silane treated basalt fibres.

### 3.6. Dimensional stability

Dimensional stability of the basalt fibre reinforced PLA composites was examined by placing them into a heated oven and measuring the changes in flow directional (longitudinal) dimensions after heating by using Equation (2). Naturally, it is found that basalt fibres restrain deformation, and it can be observed that the main deformation occurs above \(T_g\), which is caused by the specific volume change due to cold-crystallisation (Figure 14).

It is again found that basalt fibres with silane treatment are much more effective than untreated basalt fibres, this time in reducing deformation. With increasing silane treated basalt fibre content, the longitudinal deformation of PLA of 5.9% was reduced to 2.6, 1.6, 1.1, 0.7, 0.5 and 0.5% according to 5–10–15–20–30–40 wt% basalt fibre content respectively (Figure 15).

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**Table 4.** Storage modulus values of amorphous PLA, semi-crystalline PLA and basalt fibre reinforced PLA composites

<table>
<thead>
<tr>
<th>Material</th>
<th>Storage modulus at 40°C [MPa]</th>
<th>Storage modulus at 120°C [MPa]</th>
<th>Lowest storage modulus between 70–80°C [MPa]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amorphous PLA</td>
<td>2390</td>
<td>140</td>
<td>5 (at 74°C)</td>
</tr>
<tr>
<td>Semi-crystalline PLA</td>
<td>2490</td>
<td>130</td>
<td>250 (at 70°C)</td>
</tr>
<tr>
<td>5BN</td>
<td>2900</td>
<td>190</td>
<td>7 (at 73°C)</td>
</tr>
<tr>
<td>5BS</td>
<td>2920</td>
<td>290</td>
<td>12 (at 72°C)</td>
</tr>
<tr>
<td>10BN</td>
<td>3130</td>
<td>250</td>
<td>9 (at 72°C)</td>
</tr>
<tr>
<td>10BS</td>
<td>3380</td>
<td>420</td>
<td>18 (at 73°C)</td>
</tr>
<tr>
<td>15BN</td>
<td>3600</td>
<td>360</td>
<td>14 (at 73°C)</td>
</tr>
<tr>
<td>15BS</td>
<td>4000</td>
<td>590</td>
<td>31 (at 73°C)</td>
</tr>
<tr>
<td>20BS</td>
<td>4500</td>
<td>780</td>
<td>45 (at 72°C)</td>
</tr>
<tr>
<td>30BS</td>
<td>5900</td>
<td>1200</td>
<td>109 (at 72°C)</td>
</tr>
<tr>
<td>40BS</td>
<td>7030</td>
<td>1430</td>
<td>166 (at 71°C)</td>
</tr>
</tbody>
</table>

The dimensional stability of the specimens was also analysed with another method by placing them in a cantilever and registering the deflection caused by the increasing heat and the own weight of the specimens. It was observed that the deflection of the pure PLA was significantly reduced (Figure 16), thus by incorporating basalt fibres, the dimensional stability of the specimens highly increased.

It is again demonstrated that silane treated basalt fibres were much more effective than untreated fibres in reducing the deflection of the specimens. By using the same amount of basalt fibres, silane...
treated basalt fibres reduced the deflection of PLA of more than 40 to 17 mm, while untreated basalt only reduced the deflection to 32 mm. 20, 30 and 40 wt% silane treated basalt reduced the deflection to 9, 5 and 5 mm respectively as it can be observed in Figure 17 and Figure 18.

As it can be seen, the PLA, 15BN and 15BS composites were highly deformed, while the deflection of the 30BS and 40BS composites was minimal; these latter can be considered as dimensionally stable up to 120°C.

3.7. Heat deflection temperature analysis

For further analysis of the dimensional stability standardised heat deflection temperature (HDT) tests were performed. The standardised HDT analysis with a support distance of 100 mm only allows 0.33 mm deflection of the specimens which puts up for much stricter requirements than our dimensional stability test discussed previously. Due to these strict requirements, HDT was only increased by from 54 to 59°C by using 40 wt% silane treated basalt fibres into PLA, a composite was fabricated with a tensile and flexural strength of 124 and 185 MPa, a tensile and flexural modulus of more than 8 and 12 GPa and a Charpy impact strength of 9.5 kJ/m² (notched) and 38.4 kJ/m² (unnotched) respectively. By using scanning electron microscopy strong adhesion was observed between the phases, and excellent wetting of the silane treated basalt fibres was seen, which is again much better compared to literature results. It was found by using differential scanning calorimetry that basalt fibres have nucleating ability on PLA, however, semi-crystalline PLA samples with the possible maximum crystallinity of 40–45% could only be made by using low cooling rate (5°C/min) and at least 20 wt% basalt fibre content. By using injection moulding, and thus high cooling rate, the crystallinity of the basalt fibre reinforced PLA composites reached a lower, but still significant value of 19%. Besides nucleating ability of the basalt fibres, they effectively reduced the deformation and deflection caused by increasing temperature making it a more or less dimensionally stable composite up to 120°C. According to the DMA results, by incorporating the basalt fibres the storage modulus at 40 and at 120°C became three-times higher and ten-times higher compared to the storage modulus of amorphous PLA respectively. Moreover the storage modulus of the 40 wt% basalt fibre reinforced PLA was in the same order of magnitude of a semi-crystalline PLA.

4. Conclusions

In our work the usability of a novel natural fibre, the volcanic rock based basalt fibre was examined as a potential reinforcement for the most promising renewable resource based biodegradable polymer, the poly(lactic acid) (PLA) as well as the comprehensive analysis of basalt fibre reinforced PLA injection moulded composites was performed. 5–10–15–20–30–40 wt% silane treated and 5–10–15 wt% untreated basalt fibre reinforced composites were produced by using extrusion and injection moulding. According to our previous results on starch filled PLA, high drying temperature was used (120°C) to remove the smallest remnants of moisture, which could typically highly deteriorate the adhesion between the phases. As a result silane treated basalt fibre reinforced PLA composites were produced with higher mechanical properties compared to literature. By incorporating 40 wt% silane treated basalt fibres into PLA, a composite was fabricated with a tensile and flexural strength of 124 and 185 MPa, a tensile and flexural modulus of more than 8 and 12 GPa and a Charpy impact strength of 9.5 kJ/m² (notched) and 38.4 kJ/m² (unnotched) respectively. By using scanning electron microscopy strong adhesion was observed between the phases, and excellent wetting of the silane treated basalt fibres was seen, which is again much better compared to literature results. It was found by using differential scanning calorimetry that basalt fibres have nucleating ability on PLA, however, semi-crystalline PLA samples with the possible maximum crystallinity of 40–45% could only be made by using low cooling rate (5°C/min) and at least 20 wt% basalt fibre content. By using injection moulding, and thus high cooling rate, the crystallinity of the basalt fibre reinforced PLA composites reached a lower, but still significant value of 19%. Besides nucleating ability of the basalt fibres, they effectively reduced the deformation and deflection caused by increasing temperature making it a more or less dimensionally stable composite up to 120°C. According to the DMA results, by incorporating the basalt fibres the storage modulus at 40 and at 120°C became three-times higher and ten-times higher compared to the storage modulus of amorphous PLA respectively. Moreover the storage modulus of the 40 wt% basalt fibre reinforced PLA was in the same order of magnitude of a semi-crystalline PLA.
in the critical temperature range of 60–80°C (where amorphous PLA products distort), which is a relevant result, because semi-crystalline PLA has a heat deflection temperature (HDT) of more than 120°C and considered as heat stable. At the same time, the storage modulus of semi-crystalline PLA (250 MPa) was still somewhat higher than 40 wt% basalt fibre reinforced PLA (166 MPa) at 70°C and as a result, by using basalt fibres the standard HDT value only increased from 54 to 59°C. This could be related to the strict HDT test condition requirement which allows only very small deflection, although the significant improvement in dimensional stability in the case of 30 and 40 wt% basalt fibre reinforced composites compared to PLA was demonstrated. If the stiffness of the basalt fibre reinforced PLA composite can be kept above a certain level and it could pass over the mentioned critical temperature range undeformed, then in the higher temperature range cold-crystallisation could ensure a HDT value of an at least 120°C. Nevertheless, probably by using further nucleating agents like tale and/or slow cooling rate during processing basalt fibre reinforced semi-crystalline PLA composites with a heat deflection temperature of more than 120°C can be produced, which is much more suitable for engineering applications. Finally it was demonstrated that it is possible to produce basalt fibre reinforced PLA composites from renewable and natural resources with excellent mechanical properties not only by using low productivity technologies like film-stacking method not to damage the fibres but by using productive extrusion and injection moulding technologies suitable for manufacturing highly accurate, complex, 3D shaped and durable engineering parts with low cycle times.

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