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Pelletizing properties of torrefied spruce

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Abstract

Torrefaction is a thermo-chemical conversion process improving the handling, storage and combustion properties of wood. To save storage space and transportation costs, it can be compressed into fuel pellets of high physical and energetic density. The resulting pellets are relatively resistant to moisture uptake, microbiological decay and easy to comminute into small particles. The present study focused on the pelletizing properties of spruce torrefied at 250, 275 and 300 °C. The changes in composition were characterized by infrared spectroscopy and chemical analysis. The pelletizing properties were determined using a single pellet press and pellet stability was determined by compression testing. The bonding mechanism in the pellets was studied by fracture surface analysis using scanning electron microscopy. The composition of the wood changed drastically under torrefaction, with hemicellulloses being most sensitive to thermal degradation. The chemical changes had a negative impact, both on the pelletizing process and the pellet properties. Torrefaction resulted in higher friction in the press channel of the pellet press and low compression strength of the pellets. Fracture surface analysis revealed a cohesive failure mechanism due to strong inter-particle bonding in spruce pellets as a resulting from a plastic flow of the amorphous wood polymers, forming solid polymer bridges between adjacent particles. Fracture surfaces of pellets made from torrefied spruce possessed gaps and voids between adjacent particles due to a spring back effect after pelletization. They showed no signs of inter-particle polymer bridges indicating that bonding is likely limited to Van der Waals forces and mechanical fiber interlocking.

1. Introduction

The utilization of wood and agricultural biomass residues for sustainable heat and power production is an important part of future energy concepts [1]. One of the major challenges of biomass utilization for heat and power production is its unfavorable handling properties. Biomass is a bulky and
inhomogeneous material, making it both difficult and expensive to store and transport. Furthermore, it is
difficult to comminute into small particles and has a relatively low energy density (compared to fossil
fuels) and high moisture contents. An ancient process to improve the combustion properties of wood is
the manufacturing of charcoal, resulting in a product that burns at higher temperature, is easier to ignite,
and can be stored easier due to its better moisture resistance. Nevertheless, charcoal contains only 20-55
% of the raw material’s energy content, depending on how well the process is carried out [2].
Torrefaction is an advance of this process, in which wood is roasted under controlled conditions
(heating rate, temperature, time) in an inert atmosphere, retaining most of its energy [3]. The process
results in an attractive fuel, with improved heating value, low moisture content and ease of size
reduction [4]. Torrefaction is usually carried out between 230 to 300 °C [5] and removes moisture,
carbon dioxide and volatiles from the biomass. Volatiles such as hydroxymethylfurfural, furfural and
aldehydes are formed during the dehydration and decarboxylation reactions of the long polysaccharide
chains [6]. The oxygen-to-carbon ratio is lowered significantly and the energy density of the biomass is
increased, making it an ideal fuel for gasification processes where high oxygen contents are
disadvantageous [7]. The value of torrefied biomass can be further improved by mechanical
compression into pellets of high physical and energetic density [8]. The pelletization of biomass reduces
its handling costs and results in a solid fuel of standardized shape and size that can be fed automatically
in industrial and household size boilers being used for heat and power production. At present, there is a
solid interest for utilizing torrefied wood pellets in existing large-scale combined heat and power (CHP)
plants to replace coal with a biofuel without major changes in the power plant design [9].

The global production of fuel pellets from biomass was estimated to be about 13 x 10^9 kg in 2009 with
strong growth rates, and it is expected that Europe alone will reach a consumption of 50 million tons per
year by the year 2020 [10]. Consequently, the variety of raw materials used for pellet production has
greatly increased in recent years and is expected to further increase in the future. The production of
mixed biomass pellets produced from many different agricultural and forestry residues is increasing
Biomass composition has a great effect on the pellet quality [12] and the pelletizing process itself [13]. Few studies have been published on using thermally pre-treated biomass in the pelletizing process [14,15] but to the best of the authors knowledge none have so far studied in detail the impact of thermal degradation of the wood polymers on the pellet quality and integrity. In the present work, pellets were produced from Norway spruce and after torrefaction at 250, 275 and 300 °C. The chemical changes after torrefaction were investigated and their effect on the friction in the press channel of a pellet mill was studied using a laboratory scale single pellet press unit. The pellet stability was determined by compression testing and the internal bonding of the pellets was studied by fracture surface analysis using scanning electron microscopy (SEM).

2. Experimental

2.1 Materials

The raw material used in this study was Norway spruce (Picea abies K), harvested in southern Sweden (Skåne/Småland) during 2007. Wood stems were collected in autumn, debarked and comminuted into wood chips. The material was dried in a warehouse by free air circulation for four weeks and further chopped into particles < 5mm in diameter using a hammer mill (Model 55, Jensen and Sommer Aps, Denmark). The material was packed in paper bags permeable to air and moisture and stored for 24 month in a dry storage. The used samples had a particle size between 1-2.8 mm and the mass fraction of water was about 8.2 %.
2.2 Torrefaction

A lab scale torrefaction unit was constructed and built for this research similar to the one developed by Pimchuai et al. [5]. A metal box with a volume of about 2 L and two openings (5 mm in diameter) for nitrogen inlet and gas outlet was used as reactor. The box was installed in a programmable muffle furnace (S90, Lyngbyøven, Denmark) and connected to a nitrogen cylinder with pressure and flow regulator, water seal valve and fittings and pipes for gas inlet, outlet and temperature sensors. A temperature sensor (iron-constantan thermocouple) was installed in the center of the metal reactor and connected to a thermometer (52 KJ, John Fluke, USA) and a computer system controlling the heating of the oven. A torrefaction time of two hours and maximum temperature (\(T_{\text{max}}\)) of 250, 275 and 300 ºC have been chosen according to Pimchuai et al. [5].

About 450 g of wood particles were weighed in and sealed in the metal box. The box was put in the oven and heated at a rate of 2 ºC min\(^{-1}\) until \(T_{\text{max}}\) was reached and kept constant for 2 hours. Afterwards the oven was switched off and the samples were allowed to cool. Nitrogen was flushed through the box at a rate of 0.5 L min\(^{-1}\) until the samples were cooled to ambient conditions. Dry weight was determined before and after torrefaction and was used to calculate the mass loss. The torrefied samples were conditioned in climate chambers at 27 ºC at 65, 80 or 90 % relative humidity until a constant weight was reached. The samples were termed T250, T275 and T300 according to the \(T_{\text{max}}\) reached during torrefaction.

2.3 Attenuated total reflectance infrared spectroscopy (ATR-FTIR)

ATR-FTIR spectra of the pellet fracture surfaces were recorded at 30 ºC using a Fourier transform infrared spectrometer (Nicolet 6700 FT-IR, Thermo Electron Corporation, USA), equipped with a temperature-adjustable ATR accessory (Smart Golden Gate diamond ATR accessory, Thermo Electron
Corporation, USA). Samples were dried at 105 °C for 4 hours and stored in airtight containers until used for testing. A minimum of five measurements per sample was performed. To ensure good contact, all hard, solid samples were pressed against the diamond surface using a metal rod and consistent mechanical pressure. All spectra were obtained with 200 scans for the background (air), 100 scans for the sample and with a resolution of 4 cm⁻¹ from 500-4000 cm⁻¹. Spectra were normalized at around 760-790 nm where the spectra were free of any distinct IR bands.

2.4 Biomass characterization

A fiber analysis to study the biomass composition (cellulose, lignin, hemicelluloses and ash content) was conducted according to the procedure used by Davis et al. [16] and can be summarized as follows. About 10 g per sample were milled to pass through a screen with a mesh size of 0.84 mm and then vacuum-dried at 45 °C. Approximately 100 mg sample were hydrolyzed in 1 mL sulfuric acid (13.5 mol L⁻¹) for 1 hour at 30 °C. The samples were diluted to a sulfuric acid concentration of 0.75 mol L⁻¹ by adding distilled water. Fucose was added as an internal standard, and a secondary hydrolysis was performed for 1 hour at 121 °C. To control for sugar degradation during secondary hydrolysis, a standard mixture of sugars was hydrolyzed in parallel with each batch of samples. Material loss during primary hydrolysis was minimal and therefore ignored. The following secondary hydrolysis samples were immediately filtered through tared Gooch porcelain crucibles containing glass fiber filters (934-AH, Whatman, USA). The filtrate and three washes with 5 mL distilled water were collected in 100 mL volumetric flasks and brought to volume with water. The acid-insoluble residue (Klason lignin and insoluble degradation products) was washed for additional six times with 10 mL hot distilled water and its weight determined gravimetrically. Klason lignin values were corrected for ash content by gravimetric measurement following incubation of the lignin at 575 °C for > 3 hours. Sugar contents of the hydrolysates were determined by anion exchange high performance liquid chromatography using
pulsed amperometric detection. After filtration through 0.45 μm PTFE membranes, acid hydrolysates (sulfuric acid concentrations ranging between 0.2-0.75 mol L\(^{-1}\)) were injected with no further treatment. The chromatographic system consists of an autosampler (AS50, Dionex, USA) a quaternary gradient high pressure pump (GS50, Dionex, USA), and a pulsed amperometric detector (ED50, Dionex, USA). Sugar separation was achieved with guard and analytical columns (Carbo-Pac PA1, Dionex, USA) connected in series. Sugars were eluted with distilled water at a flow rate of 1.1 mL min\(^{-1}\) and a temperature of 18 °C. For detection, sodium hydroxide solution (0.3 mol L\(^{-1}\)) as added as a post-column reagent at a flow rate of ca. 0.3 mL min\(^{-1}\). Prior to each injection, the anion exchange columns were conditioned with 400 mL sodium acetate solutions 0.24 mol L\(^{-1}\) and then equilibrated with distilled water. Sugars were quantitated using an internal standard method. Results are reported in terms of percent of the original sample mass dry matter.

Moisture uptake was studied by spreading 5 g of each material on a tray and conditioning in climate chambers (USDA Forest products lab, USA) at 27 °C and a relative moisture content of 65, 80 and 90 %. Equilibrium moisture content was determined after the weight of the sample was constant for three days in a row. The moisture content was calculated based on weight loss after oven drying the material for 8 hours at 105 °C.

2.5 Pellet preparation and determination of pelletizing pressure P_x in the channel of the pellet press

The pellets were prepared as described in [12], using a single pellet press (invented and constructed at the workshop of the Technical University of Denmark, Denmark). The press consisted of a cylindrical die 7.8 mm in diameter, made of hardened steel, lagged with heating elements and thermal insulation. The temperature was controlled using a thermocouple connected to a control unit. The end of the die was closed using a removable backstop. Pressure was applied using a metal piston. The entire pellet press was mounted in a material test system (MTS 810 Material Test system, MT Systems Corporation,
USA) so that piston movement could be controlled and the force could be measured using a 100 kN load cell. The die was rinsed with acetone, and wiped clean using a paper towel before each use, and when changing raw materials. To simulate the pelletizing process within a commercial pellet mill, the pellet had to be built up in sequential layers [17]. The die was heated to 100 °C. Spruce and torrefied spruce particles equilibrated at 65 % relative humidity (see Figure 3 for moisture content) were loaded stepwise in amounts less than 0.25 g into the unit, and then compressed at a rate of 2 mm s⁻¹ until a maximum pressure of 200 MPa was reached. The pressure was released after five seconds, the piston removed, and more biomass was loaded and compressed until the pellet had a length of about 16 mm. This results in a layered structure, similar to pellets obtained by commercial units, although there are some differences. The most significant difference is that the lower part of the pellet is pressed repeatedly, and the upper layers are pressed fewer times, with the top layer being pressed only once. For determination of pelletizing pressure in the press channel of the pellet mill, $P_x$, the pellets were removed from the die by removing the backstop and pushing out the pellet at a rate of 2 mm s⁻¹. The applied maximum force was logged and $P_x$ was calculated based on the pellet surface area.

2.6. Determination of pellet strength

The internal strength of the manufactured pellets was analyzed by compression testing and determined as the force at break. Pellets 16 mm (± 1 mm) in length and between 7.9 and 8.2 mm in diameter were produced in the single pellet press, stored at a relative humidity of 50 % and 20 °C for three weeks, and tested under the same conditions. The pellets were placed on their side (the pellet’s cylindrical shape oriented horizontally) in the same material tester as was used for pellet preparation. Compression tests were performed using a disc shaped metal probe with a ball bearing of 50 mm in diameter and attached to a 100 kN load cell. The test was run at a compression rate of 20 mm min⁻¹ and was stopped after pellet failure. The average force at break and its standard deviation were calculated based on 5
replications per test condition.

2.7 Scanning electron microscopy (SEM)

SEM was used to study the bonding mechanism of the prepared biomass pellets by fracture surface analysis of failed pellets. The compression test resulted in total disintegration of the pellets, and therefore fracture surfaces were prepared by manually breaking a pellet into two parts. Care was taken to replicate the way each pellet was broken and that it took place in the same region. A tiny notch was cut in the center of the pellet using a razor blade, and the pellets were snapped into two. Care was taken to examine the fracture surface away from the notch. The two halves of the fractured pellet were attached to metal stubs using a conductive silver paste (Conductive silver paste plus, SPI Supplies, USA) that was carefully applied below and around the sample to prevent electric charging of the specimen. The upper surface was coated with a thin layer of gold using a sputter coater (Desk-1 sputter coater, Denton, USA). Electron micrographs were recorded using a scanning electron microscope (LEO EVO 40 SEM, Carl Zeiss, Germany) operated at 5-15 kV. Multiple samples were observed for each type of pellet and representative images were selected for each sample type.

2.8 Pellet density

The obtained pellets were stored at 50% relative humidity at 20 °C for one month. The unit density of the pellets was calculated by determination of the pellets weight and dimensions for at least 5 samples for pellets pressed at each condition.
3. Results and Discussion

Figure 1. Color change during the torrefaction process.

The torrefaction of spruce at 250, 275 and 300 °C resulted in three products of light brown, dark brown and black color as shown in Figure 1. The color change is mainly attributed to chemical changes of the lignin, i.e. the formation of chromaphoric groups, mainly the increase of carbonyl groups [18]. The loss of dry matter (anhydrous weight loss) through volatilization during torrefaction was 25 % at 250 °C, 34 % at 275 °C and 53 % at 300 °C.

<table>
<thead>
<tr>
<th>Norway Spruce</th>
<th>Cellulose</th>
<th>Hemicellulose</th>
<th>Acid insoluble fraction</th>
<th>Ash</th>
</tr>
</thead>
<tbody>
<tr>
<td>T250</td>
<td>43.6</td>
<td>6.6</td>
<td>43.2</td>
<td>0.1</td>
</tr>
<tr>
<td>T275</td>
<td>32.7</td>
<td>0.7</td>
<td>62.2</td>
<td>0.3</td>
</tr>
<tr>
<td>T300</td>
<td>0.2</td>
<td>0.0</td>
<td>99.0</td>
<td>0.6</td>
</tr>
</tbody>
</table>

*) In case of spruce the acid insoluble fraction is defined as Klasson lignin content
The chemical analysis of the torrefied spruce shows a strong decrease of hemicellulose and cellulose content and is attributed to the thermal degradation of the carbohydrate polymers into volatile compounds and the evaporation of water and carbon dioxide [19]. Hemicelluloses are well known to undergo a two step thermal degradation where light volatiles i.e. mono and polysaccharides, followed by their catalytic degradation into CO and CO₂ [20]. Lignin undergoes depolymerisation, mainly β-aryl-ether linkages and re-condensation reactions that lower its average molecular weight [6]. The relative increase in the acid-insoluble fraction with torrefaction temperature is likely due to both the volatilization of some of the carbohydrate fraction as well as the formation of acid-insoluble degradation products from them [21]. For example, cellulose can undergo scission reactions with aromatization and cross-linking, resulting in an insoluble solid [22]. The chemical changes occurring during torrefaction have been studied using ATR-FTIR-spectroscopy and the results are shown in Figure 2.

![ATR-FTIR spectra](image)

*Figure 2. ATR-FTIR spectra of dry spruce and torrefied spruce at 250, 275 and 300 °C. For better comparability, the scale of the y-axis was adjusted.*
The IR spectra of spruce and torrefied spruce at 250 and 275 °C have several features in common, while the spectra of spruce torrefied at 300 °C has very different characteristics. The broad band in the OH stretching vibration region at about 3600-3200 cm⁻¹ is due to intra- and inter molecular hydrogen bonds and bands characteristic of crystalline cellulose [23-25]. The intensity of these bands decreases with increasing torrefaction temperature and has almost disappeared at 300 °C. This indicates that the torrefied material contains less water and hydrogen bonding sites due to the degradation of hemicelluloses and cellulose, which is supported by the chemical analysis data (Table 1). A lignin vibration can be found at about 1269 cm⁻¹ (the aromatic C-O stretching of methoxyl and phenyl propane units) and at 1516 cm⁻¹ and 1508 cm⁻¹ (C=C aromatic ring vibrations) [23, 26, 27]. These bands were present in the raw material and after treatment at 250 and 275 °C at same intensity but has disappeared after treatment at 300 °C, this suggests that the lignin, to large extent, has been degraded at this high temperature. Vibrations at about 1735 cm⁻¹ are related to C=O stretching vibrations of the carboxylic acids of hemicelluloses (i.e. xyloglucan, arabinoglucuronoxylan and galactoglucomannan) [23, 28]. This vibration is only present in the raw material. The torrefied samples show a band at about 1700 cm⁻¹ which is attributed to a degradation product formed during the torrefaction. This indicates that hemicelluloses are degrading already at 250 °C, which agrees with the chemical analysis data (Table 1). The band at 1160 cm⁻¹ is celluloses antisymmetric stretching of C-O-C glycosidic linkages [29] its intensity is strongest for untreated spruce and decreases with increasing torrefaction temperature. The band is not present at 300 °C, which indicates that most cellulose has degraded at this temperature.

Torrefaction has been reported to have a strong effect on the mechanical stability and combustion properties of biomass [2,3,4].

Since the IR data has shown that most of the hydrogen-bonding hydroxyl groups have been removed during torrefaction, the moisture uptake of torrefied biomass is expected to decrease. The results
presented in Figure 3 show that the spruce absorbed twice as much water as it did after torrefaction at 250 °C. The ability to absorb water decreased further after the 275 and 300 °C treatment.

![Graph showing moisture content of spruce and torrefied spruce after three weeks storage at 27 °C and 65, 80 and 90 % relative humidity.](image)

**Figure 3.** Moisture content of spruce and torrefied spruce after three weeks storage at 27 °C and 65, 80 and 90 % relative humidity.

There are three different types of bound water in wood [30]. Non-freezing bound water is specifically bound to the hydroxyl groups of the wood polymers, especially the hemicelluloses. Freezing bound water is loosely bound to hydroxyl groups organized in clusters and can be found at high humidity in nanovoids and on the wall lining of macro- and microvoids. Free water is held in macro- and microvoids and bound by capillary forces but not to specific sorption sites. Since hemicelluloses are degraded during the heat treatment and most hydroxyl groups are removed, it is likely that water in torrefied wood is mainly bound as free water in macro- and micro-sized voids where it is held in place by capillary forces. This supports the observation, made during this study, that torrefied wood reaches equilibrium moisture content much faster than untreated wood.

The differences in composition and water content have also a strong effect on the pelletizing properties
for spruce and the torrefied spruce. The pelletizing pressure in the press channel of the pellet mill (\(P_x\)) is a crucial parameter in pelletizing processes in terms of process energy consumption and pellet quality [15, 31].

\[\text{Figure 4. Pressure (}\ P_x\ \text{during the pelletization of spruce and torrefied spruce.}\]

\(P_x\) increases drastically when comparing spruce and torrefied spruce (Figure 4). This increase is most likely attributed to the lack of water and low hemicelluloses content in the torrefied spruce. Water acts as a plasticizer, lowering the softening temperature of the wood polymers. In nature, hemicelluloses bind lignin and cellulose fibrils and provides flexibility in the plant cell wall [32]. Their degradation embrittles wood, making it easier to comminute into small particles [4, 33]. The degradation of the hemicelluloses, cellulose and the lignin are likely to affect important pelletizing parameters such as the friction coefficient and Poisson ratio which are directly correlated to \(P_x\) [15].

It is likely that extractives are removed during the torrefaction process. Extractives have been shown to play an important role during the pelletizing process [12] and are likely to act as lubricant lowering the
friction in the press channel.

Figure 5. Pellets made from spruce and torrefied spruce. From left to right: Spruce and torrefied spruce at 250, 275 and 300 °C.

The pellets were very different in their quality (Figure 5). No pellets could be made from spruce torrefied at 300 °C and even at 275 °C the pellets exhibits many defects. The pellet length (after one month storage) increased with an increasing degree of torrefaction from 19.3 ± 0.3 mm for spruce pellets to 23.9 ± 1.1 mm for spruce torrefied at 250 °C and 28.5 ± 1.2 mm for spruce torrefied at 275 °C. The unit density decreased with an increasing torrefaction temperature from 1090 ± 19 kg m⁻³ (untreated spruce) to 832 ± 39 kg m⁻³ for pellets made from spruce torrefied at 250 °C and to 698 ± 30 kg m⁻³ for pellets made from material torrefied at 275 °C. This attributes to a spring-back effect and is a sign of poor adhesion between the particles [34].
Figure 6. Compression strength of spruce and torrefied spruce pellets.

Figure 6 shows a strong decrease in pellet compression strength as a result of torrefaction. Pellets made from T300 were too weak to be tested. Yildiz et al. [33] have tested the compression strength of spruce that had been heat treated between 130 and 200 °C for 2 to 10 hours. They found that the compression strength of their samples decreased both with treatment time and temperature and concluded that the strength loss is connected to the degradation of hemicelluloses.

In an earlier work [12], the bonding and failure mechanisms in fuel pellets made from spruce, beech and straw was studied by means of fracture surface analysis using SEM. This method was applied to pellets made from torrefied spruce (Figure 7).
Figure 7. Low, medium, and high magnification, respectively, of a pellet fracture surface for spruce (a-c), spruce torrefied at 250 °C (d-f) and spruce torrefied at 275 °C (g-i).

The low and medium magnification images show more inter-particle gaps and voids with increasing torrefaction temperature, indicating poor adhesion between adjacent particles and/or spring back effects [12, 34, 35]. Images taken at higher magnification, (Figure 7c,f and i) provide a deeper insight into the bonding and failure mechanisms of the pellets. The failure surface of spruce pellet (Figure 7c) indicates a cohesive failure with a high energy absorption. Fiber ends and particles are sticking out of the surface and few voids are found. This suggests that lignin and hemicelluloses have exceeded their glass transition temperatures at the pelletization conditions (e.g., moisture content, temperature and pressure), allowing them sufficient mobility to flow into cracks and crevices and establish solid bridges between adjacent particles. The pellets from torrefied wood, T250 (Figure 7f) and T275 (Figure 7i), have flatter
failure surfaces than those of the spruce pellets, indicating a minimized polymeric flow. The amount of hydrogen binding sites decreases gradually with the torrefaction temperature and the torrefied biomass contains less moisture, thus both hydrogen bonding between polymer chains of adjacent particles and a polymeric flow of the lignin and hemicelluloses (forming solid bridges) can be assumed less likely to occur. According to Rumpf [36] who has studied the binding mechanisms in biomass granules and agglomerates it is likely that Van der Waals forces and fiber interlocking remain as the major forces keeping a pellet together. Since these forces are weak compared to covalent bonds and hydrogen bonding this could be an explanation both for the observed spring back effect (Figure 5) as well as for the low compression strength of pellets made from torrefied spruce (Figure 6).

It is well known that a combination of torrefaction time and temperature determine the degree of polymer degradation and the mechanical properties of heat treated wood [33, 35]. The mechanical properties of a pellet have been shown to depend on the one hand on the biomass composition i.e. moisture and extractives content and on the other hand on the processing conditions i.e. temperature and pressure [12, 37-40]. Since hemicelluloses and lignin are both subject to thermal degradation during torrefaction and the amount of available hydrogen bonding sites is reduced, it can be expected that the pellet strength of torrefied pellets is lower compared to pellets made from untreated spruce. Furthermore, the moisture content of the torrefied wood is lower which results in an increase in the glass transition temperatures of the remaining hemicelluloses and lignin [41]. This may reduce the inter-diffusion of the wood polymers between adjacent particles in a pellet and thus the formation of solid bridges between them. The resulting pellets are brittle and less stable than pellets made from untreated wood. To increase the mechanical properties of torrefied biomass pellets it is necessary to establish a better bonding between the particles. One possibility is to add an additive that compensates for the lost bonding sites and ideally binds between the hydrophobic surface of the biomass and remaining polar groups on the wood polymer surface such as short fatty acids or mono glycerids. Another option could be to adjust the torrefaction parameters so that less of the wood polymers are degraded to ensure
sufficient hydroxyl groups on the wood polymer chains to form strong inter particle bonds. This could be achieved by shorter treatment time, lower torrefaction temperature, different torrefaction medium (e.g. wet torrefaction) or surface increase (e.g. steam explosion conditions).

The development of a combined torrefaction pelletization concept requires that both processes are adjusted to each other. An interesting recent work reports improved mechanical stability and moisture resistance for pellets made from steam exploded Douglas fir [14] performed at temperatures between 200-220 °C for 5-10 min. The water vapor grants a fast heat transfer and the short treatment time and relatively low temperature (compared to our study) preserves sufficient amounts of amorphous polymers that can be plasticized during pelletization and form solid inter-particle bridges.

Future, studies have to investigate the optimal torrefaction conditions to obtain a raw material that both has the beneficial properties of torrefied biomass and also can be pelletized into stable pellets. Furthermore the impact of raw material composition on the torrefaction and pelletizing process has to be investigated.

4. Conclusions

The torrefaction of biomass degrades hemicelluloses, cellulose and lignin and removes moisture from the material. Both effects have a strong effect on the pelletizing properties of the biomass. The friction in the press channel of a pellet mill increases, resulting in high pelletizing pressures which increase the energy uptake of the mill and might result in a decrease of capacity and in worst case an overheating (risk of fire) of a blockage of the mills press channels.

Torrefaction breaks down the carbohydrates and therefore the ability to establish hydrogen bonds between polymer chains of adjacent particles can be assumed to be strongly reduced. The lack of
moisture increases the glass transition temperature of the remaining carbohydrate polymers and restricts the ability of polymeric flow and the establishment of solid bridges between particles. There are several options to overcome this lack of bonding capacity. For example, an additive with a high bonding capacity could be introduced after the torrefaction process that would compensate for the lost hydrogen bonding sites due to torrefaction.

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