Medical Peat Waste Upcycling to Carbonized Solid Fuel in the Torrefaction Process

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Abstract: Peat is the main type of peloid used in Polish cosmetic/healing spa facilities. Depending on treatment and origin, peat waste can be contaminated microbiologically, and as a result, it must be incinerated in medical waste incineration plants without energy recovery (local law). Such a situation leads to peat waste management costs increase. Therefore, in this work, we checked the possibility of peat waste upcycling to carbonized solid fuel (CSF) using torrefaction. Torrefaction is a thermal treatment process that removes microbiological contamination and improves the fuel properties of peat waste. In this work, the torrefaction conditions (temperature and time) on CSF quality were tested. Parallelly, peat decomposition kinetics using TGA and torrefaction kinetics with lifetime prediction using macro-TGA were determined. Furthermore, torrefaction theoretical mass and energy balance were determined. The results were compared with reference material (wood), and as a result, obtained data can be used to adjust currently used wood torrefaction technologies for peat torrefaction. The results show that torrefaction improves the high heating value of peat waste from 19.0 to 21.3 MJ kg⁻¹, peat main decomposition takes place at 200–550 °C following second reaction order (n = 2), with an activation energy of 33.34 kJ mol⁻¹, and pre-exponential factor of 4.40 × 10⁻¹ s⁻¹. Moreover, differential scanning calorimetry analysis revealed that peat torrefaction required slightly more energy than wood torrefaction, and macro-TGA showed that peat torrefaction has lower torrefaction constant reaction rates (k) than wood 1.05 × 10⁻⁵–3.15 × 10⁻⁵ vs. 1.43 × 10⁻⁵–7.25 × 10⁻⁵ s⁻¹.

Keywords: peloids; waste to energy; waste to carbon; circular economy; torrefied biomass; kinetics lifetime prediction; mass balance; energy balance; fuel properties

1. Introduction

Peloids are commonly used in spa treatment and cosmetology for cosmetic and healing purposes. Peloids refer to inorganic, organic, or a mixture matter that originated through geological and/or biological processes. Özay et al. [1] proposed to divide peloids by origin and composition into four groups: peat, sludge (bituminous or mineral), slime (sea or delta), and earth (rock or schist) [1]. The most used peloid in Poland is peat. Leśniak [2] shows that a typical SPA facility in Poland uses from 2.90 to 3.42 Mg of peat yearly. Assuming an average value of 3.16 Mg year⁻¹ and a current number of spa facilities (103), the annual peat waste (PW) production counts about 324 Mg [2]. The PW management depends on the specific situations in the country and its law regulations [3]. In general, waste peloids generated in beauty salons are safe, but ones from spa centers may be microbiologically contaminated, and in such a situation, PW must be considered as medical waste, requiring a special waste treatments method, e.g., waste incineration. However, other methods of PW recycling and upcycling should be developed. In this study, we check the validity of thermal upcycling of medical/cosmetic peat waste into high-quality solid fuel, free of possible infectious substances using the torrefaction process.
1.1. Peat Applications

Peat can be classified according to different factors such as fiber content or fiber decomposition, organic matter, and vegetation forming the organic content. In addition, ASTM D4427-13, “Standard Classification of Peat Samples by Laboratory Testing”, can be used for peat classification. There is also a special von Post scale, which classifies peat soils based on the degree of its decomposition at 10 points (H₁–H₁₀ with the interpretation from none to complete decomposition) [4]. For medical/cosmetic purposes, typically, peat with classes from H₆ to H₁₀ is used [5]. Each class corresponds to 10% of decomposition (degree of humification). Where H₁ means peat, which squeezed releases almost clear water, plants are identifiable, and no amorphous material is present, while H₁₀ means completely decomposed peat, which squeezed escape between fingers, without discernible plants structures [6]. Medical/cosmetic peat usage in Europe is dated to the 18th century, but external usage as poultices is dated to the early 19th century [7]. Peat in cosmetics is used, inter alia, for facial care treatments supporting the treatment of acne, and scars, and sun discoloration, as an antiaging treatment that improves the elasticity of the facial skin and so on [5]. On the other hand, in a healing spa (peat bath), peat is mainly used to overheat the body. It is possible thanks to high heat capacity and relatively flat cooling curve in comparison to water. As a result, baths improve blood circulation, effects myalgia, and chronic inflammation, and so on [7].

Using peat as fuel started earlier. Since the 12th-century, peat has become an important energy source in Europe. Nevertheless, currently using peat for energy production has ended in most countries [8]. Only six countries in the EU (Estonia, Finland, Ireland, Latvia, Lithuania, and Sweden) use peat for energy purposes excavating around 9.4 mln Mg per year of it. Among these countries, only 208 peat-fired energy plants exist, and only Finland uses peat in significant amounts, covering around 25% of its national heat needs [9]. Nevertheless, peat does not play a significant role in the energy sector of the EU (1.7%), and since 2010, its consumption has decreased. As a result, it is highly probable that peat’s role in the EU energy sector will further decrease [10]. On the other hand, Russia, which has a significant peat reserve (~30% of the world’s peat), is considered to go back to peat mining for energy uses as a more environmentally friendly alternative for coal and to diversify locals’ energy sources. The main peat advantage is its abundance and low-cost mining in comparison to coal. Nevertheless, peat is characterized by high moisture content and relatively high ash content. Therefore, before energy use, peat required seasoning and pretreatment [11–13].

Besides medical/cosmetic and energy purposes, peat is also widely used in horticulture and agriculture. Peat-specific properties allow its use as a soil improver, growing medium, compost ingredient, etc. For example, peat mining for agricultural usage is common in such countries as Germany and Poland [10], which do not use it for energy purposes. On the other hand, peat and peatlands represent a valuable global carbon store and are critical for preserving biodiversity [14]. Moreover, the analysis of the European Commission from 2018 shows that limiting the use of peatlands is effective for CO₂ emission mitigation [11]. Therefore, peat replacement materials are sought. The opportunity for such replacement material is looking in biochars and hydro-chars obtained from organic waste [15,16]. Since carbon materials alone or in mixtures with organic substances have the potential to substitutes (partially or full) peat for agricultural usages. For example, some biochars (depending on substrate and process conditions) have higher pH, increased surface area, excellent water, and nutrient retention properties in comparison to peat. Moreover, biochars may contain different forms of nitrogen and have a considerable amount of potassium [15]. That suggests that biochar from medical/cosmetic peat waste may partially replace fresh peat usage in agriculture/horticulture or other emerging applications such as odors mitigation [17].
1.2. Peat Thermal-Upgrading Methods

Some studies show that peat can be upgraded thermally by various methods, i.e., torrefaction [18], pyrolysis [19], gasification [20], or hydrothermal carbonization [21]. Each process favors different input substrates, process conditions, and end products. Torrefaction is typically used to improve the calorific value and storage properties (increased hydrophobicity) of biomass. It takes place at 200–320 °C in the absence of oxygen. Solids are the main process products [22]. The pyrolysis also takes place in the absence of oxygen but at temperatures over 320 °C. As a result, depending on pyrolysis conditions main products are liquid, gas, or solid; nevertheless, amounts of solids obtained from pyrolysis always are lower than from torrefaction. On the other hand, gasification takes place in a partially oxidative atmosphere and converts substrates mainly into gases. This process requires temperatures from ~800 to 1150 °C [23,24]. The main drawback of mentioned processes is the fact that materials need to be dry, or additional energy needs to be spent on water removal. Moreover, in the case of pyrolysis and gasification, high process temperatures required much more energy in comparison to torrefaction [25]. Partially, these drawbacks can be overcome by hydrothermal carbonization (HTC), which favors wet substrates, and low temperatures 150–320 °C. Nevertheless, HTC required sophisticated reactors that can stand high pressure up to 50 bars or more [26,27].

Some studies show the prospect of thermal peat upgrading to fuels (gases or solids) using the overmentioned processes [18–21,28]. Nevertheless, none of them focus on medical/cosmetic peat waste particularly. In addition, none of the studies alone provide comprehensive data of peat upcycling to carbonized solid fuel by torrefaction, including the effect of process conditions on CSF fuel qualities, torrefaction kinetics, energy, and mass balance of the whole process. Especially that peat is not homogenous material, and its composition varies significantly around the world. As a result, adopting data from different studies to make any conclusion may not be fully successful and precise.

The torrefaction is a well-known and world-wise used process for wood energy properties upgrading [29]. Besides energy densification in a solid product, torrefaction removes water, increases hydrophobic characteristics, decreases energy to grind, and decreases biodegradability [30]. As a result, torrefied material becomes easier to transport, manage, and storage [31]. Such positive effects are mainly due to hemicellulose decomposition. The hemicellulose is rich in hydroxyl groups (including carboxylic and phenolic groups), which can form a hydrogen bond with water increasing bonded water content [30,32]. When hemicellulose is decomposed because of depolymerization, demethoxylation, and bond cleavage reactions, these groups are removed as well. As a result, fuel becomes hydrophobic and more fungal-resistant [32]. In addition, changes in chemical composition (decrease in available nutrients and production of toxic chemicals) the resistance to biodegradation increases [30]. The end effect and properties of torrefied material depend on used feedstock, reactor type, and process conditions, i.e., temperature, time, heating rate, feedstock particle size, pressure, etc. [33].

1.3. Aim of the Study

In this study, we performed initial laboratory analyses on medical/cosmetic peat waste upcycling using torrefaction. The analyses covered the effects of process conditions on carbonized solid fuel quality, thermal decomposition kinetics, torrefaction kinetics with lifetime prediction, and theoretical mass and energy balance determination. The analyses were performed following our previous research related to wood biomass torrefaction. Therefore, obtained data are comparable to well-described wood material. As a result, obtained data may be used for adjusting the existing technology used for wood torrefaction to peat torrefaction.
2. Materials and Methods

2.1. Materials

The medical/cosmetic peat come from a peat bog “Kołobrzeg” owned by Uzdrowisko Kołobrzeg S.A. The peat bog is located around 3.5 km from Kołobrzeg, Poland. The peatland is classified as a low type [2,34], and the medical properties of the peat are confirmed by the National Institute of Public Health. The natural moisture content of peat obtained from the peat bog was 89.9%, and organic dry mass was 92.5% of dry mass. The peat had a degree of humification (according to the van Post scale) between H4 and H6 [2,35].

2.2. Methods

2.2.1. Torrefaction Process—CSF Production

The carbonized solid fuel (CSF) was produced using a muffle furnace (Snol 8.1/1100, Utena, Lithuania). The dry peat samples of ~20 g were placed at ceramic trays and put in the muffle furnace. To keep the inert atmosphere, CO2 was introduced into the furnace. The furnace was heated from room temperature to setpoint temperature (200–300 °C, intervals 20 °C). When the temperature inside reached the setpoint, it was kept for 20, 40, or 60 min. After this time, the furnace was left to self-cooling, and when the temperature dropped below 100 °C, samples were removed. For the CSF production process, mass yield (MY), energy densification ratio (EDr), and energy yield (EY) were determined using Equations (1)–(3).

\[ MY = \frac{m_{CSF}}{m_{raw}} \times 100 \]  
\[ EDr = \frac{HHV_{CSF}}{HHV_{raw}} \]  
\[ EY = MY \times EDr \]  

where:
MY—mass yield, %; \( m_{CSF} \)—mass of produced CSF, g; \( m_{raw} \)—a mass of substrate used for CSF production, g; EDr—energy densification ratio, -; HHV_{CSF}—high heating value of CSF, J × g\(^{-1}\); HHV_{raw}—high heating value of substrate used for CSF production, J × g\(^{-1}\); EY—energy yield, %.

2.2.2. Proximate Analysis and High Heating Value Determination

Proximate analysis and high heating value determination were performed for raw peat and produced CSF. The proximate analysis consisted of moisture content (MC), volatile matter (VM), ash content (AC), and fixed carbon (FC). Additionally, volatile solids (VS) and combustible parts (CP) were determined. The proximate analysis with additional parameters was determined according to methods presented in previous work [36]. The high heating value (HHV) was determined using a calorimeter (IKA, C200, Staufen, Germany). Each sample was tested in three repetitions. Used standards, methods, and equipment are listed in Appendix A, Table A1.

2.2.3. Statistical Analyses

CSF production data and results of the proximate analysis with HHV were subjected to regression analysis. The regression analysis aimed to provide useful empirical models for easy determination of CSF properties and CSF production process in function of torrefaction temperature and time. The regression was performed according to the methodology described in previous work [37]. In short, empirical data were subjected to four main regression equations (linear, second-order polynomial, factorial regression, and response surface regression). Then, models were compared using the determination coefficient \( R^2 \) and the Akaike information criterion (AIC). Then, models with the highest \( R^2 \) and the lowest AIC were chosen as the best fitted to the experimental data. Next, coefficients significance was checked. When some coefficients turned out to be not significant, it was removed from the model, and rest of the coefficients were recalculated again.
To check if there is an effect of process temperature and the treatment time on CSF properties, ANOVA analyses were performed at ($p < 0.05$). Then, to check between which groups the difference occurs, a Tukey post-hoc test was performed.

### 2.2.4. Thermal Decomposition Analysis

The thermal decomposition of dry peat and wood (as reference material) was studied using two different approaches and equipment, TG/DTG-DSC analyzer and macroTG/DTG analyzer. The mentioned abbreviation means TG—thermogravimetry; DTG—difference thermogravimetry; DSC—differential scanning calorimetry. The macro term refers to sample size. In standard TG, the sample is measured in mg, while in macro-TG, the sample mass is measured in g.

Firstly, samples were subjected to TG/DTG-DSC analysis, where samples were decomposed at a range of 30–800 °C. In this test, the temperature increased with a constant rate of 10 °C × min$^{-1}$. For this analysis, a sample of ~3.6 mg was subjected to a simultaneous thermal analyzer (Netzsch, 449 F1 Jupiter, Selb, Germany) in a nitrogen atmosphere to prevent Boudouard reaction at >700 °C that occur when CO$_2$ is used [38]. Experiment results showed material decomposition in the function of time and allowed for the determination of decomposition kinetics. Moreover, TG-DSC results were used for the calculation of the theoretical mass and energy balance of the torrefaction process (Section 2.2.5) according to previous work [39]. The decomposition kinetics were determined using the well-known Coats–Redfern method (CR) [40]. For the CR method, the kinetic triplet consists of activation energy (E$_a$), pre-exponential factor (A), and reaction order (n). In this study, kinetic triplets were determined for the whole process as well as decomposition peaks. The full methodology of kinetic triplet determination is described elsewhere [36].

Secondly, dry peat samples were subjected for macroTG/DTG analysis at torrefaction temperature range 200–300 °C. For this test, stand-mounted tubular furnace (Czylok, RST 40 × 200/100, Jastrzębie-Zdrój, Poland) coupled with the laboratory balance (RADWAG, PS 750.3Y, Radom, Poland) was used. The samples of ~3 g were heated by 60 min at 200, 220, 240, 260, 280, and 300 °C in the CO$_2$ atmosphere to imitate conditions of setpoint temperatures of CSF production (Section 2.2.1). The results showed thermal decomposition at specific torrefaction temperature in function of time. Moreover, results were used for torrefaction kinetics determination for isothermal temperature according to previous work [41]. In short, for each torrefaction temperature, the constant reaction rate (k) was determined by experimental data fitting to the first-order reaction. Then, using determined k values, the Arrhenius plot, ln(k) vs. 1/T, was drawn. Next, from the obtained slope and intercept, the activation energy and pre-exponential factor for the torrefaction process were obtained. Then, using obtained kinetic triplet, the prediction of the lifetime of material exposed to isothermal conditions may be performed for any temperatures in the range of 200–300 °C. In this study, kinetic triplet was used for the prediction of the temperatures, which was used for kinetic determination. The result of wood subjected to this method is presented in previous work [41].

### 2.2.5. Theoretical Mass and Energy Balance of the Torrefaction Process

Theoretical mass and energy balance of the torrefaction process were calculated for peat and wood (as reference material) using data from some of the analyses. The balances were calculated to produce 1 g of CSF. The balance calculations include the mass of substrate needed to produce 1 g pf CSF; external energy needed to heat this mass of substrate to the setpoint temperature (DSC results) (including mass change during the heating); the energy contained in 1 g of CSF (HHV results); a mass of gas produced during the production of 1 g of CSF, and energy contained in this gas. To calculate the mentioned values, results of MY, HHV, and TG/DSC were used. The graph of balances determination is shown in Figure 1. The green squares represent the order of preformed calculations, the gray squares represent experimental/calculated data used for balances determination, and the blue squares stand for input and output data.
The calculations were performed in four steps. First (I), knowing MY of CSF production, a mass of substrate needed for 1 g of CSF was calculated. Then, knowing the HHV of a substrate, energy contained in the input substrate was calculated. Secondly (II), the results of DSC were used to determine the amount of energy needed to heat a proper amount of substrate to the setpoint temperature. Here, DSC results were given by thermal analyzer software right away in mW per mg. Therefore, by multiplication DSC results by time, the external energy provided to the sample was calculated. Thirdly (III), the energy contained in CSF was determined by experimental HHV determination. Fourthly (IV), the mass and energy contained in the gas produced alongside the production of 1 g of CSF were determined. The mass of gas was calculated as the difference between the used substrate and the mass of obtained CSF, while the energy was calculated as the difference between energy contained in substrate and CSF increased by external energy provided to heat substrate to the setpoint temperature. Here it was assumed that CSF is completely cooled before it is removed from the reactor, and this energy goes to the gas phase (Figure 1).

To keep calculations as simple as possible, the following assumptions were adopted:
(i) Processed material is completely dry, MC = 0%;
(ii) Only external energy is used to provide heat for the process;
(iii) The process has 100% efficiency, no heat losses;
(iv) All external energy after the process goes to gaseous products.

3. Results and Discussion

3.1. CSF Production and Proximate Analysis

All raw data generated in this study were given in Supplementary Materials. The result of the raw peat and CSF properties were summarized in Table 1. The 3D models with equations presenting occurring changes in the function of torrefaction temperature and time are presented in Appendix A (Figure A1a–i).

The raw peat used in this study had 84.8% of moisture content, and its dry mass was characterized by 59.3%, 26.3%, and 14.4% content of VM, FC, and AC, respectively. Moreover, the average high heating value was 19,013 J × g⁻¹. The peat composition varies in a wide range. For example, Canadian peat has only 36% of MC, but almost 70% of VM, around 30% of FC, and only 2% of AC [42], while Bangladesh peat was characterized by 20–50% of MC, 7–48% of VM and ash content up to 90% [43]. Therefore, direct data comparison needs to be performed carefully.
Table 1. Results of CSF production and CSF properties.

| Temp. | Time | MY, % | EDr, - | EY, % | VM, % | AC, % | FC, % | VS, % | CP, % | HHV, J × g⁻¹ |
|-------|------|-------|--------|-------|-------|-------|-------|-------|-------|-------------|
| 0     | 0    | -     | -      | -     | 59.3  | 14.4  | 26.3  | 83.0  | 85.6  | 19,013      |
| 200   | 20   | 97.6  | 1.03   | 100.6 | 56.0  | 14.1  | 29.9  | 83.1  | 85.9  | 19,598      |
|       | 40   | 97.3  | 1.05   | 102.1 | 55.8  | 14.2  | 30.0  | 82.9  | 85.8  | 19,941      |
|       | 60   | 97.1  | 1.04   | 101.4 | 57.2  | 14.6  | 28.2  | 82.7  | 85.4  | 19,841      |
| 220   | 20   | 98.4  | 1.03   | 101.5 | 56.8  | 14.1  | 29.0  | 83.2  | 85.9  | 19,619      |
|       | 40   | 95.4  | 1.07   | 101.7 | 56.3  | 14.3  | 29.4  | 82.8  | 85.7  | 20,272      |
|       | 60   | 91.2  | 1.06   | 97.0  | 55.0  | 15.2  | 29.7  | 81.9  | 84.8  | 20,217      |
| 240   | 20   | 96.7  | 1.02   | 98.9  | 55.3  | 14.2  | 30.5  | 83.0  | 85.8  | 19,444      |
|       | 40   | 90.7  | 1.07   | 97.0  | 54.0  | 14.8  | 31.2  | 82.2  | 85.2  | 20,341      |
|       | 60   | 87.7  | 1.07   | 93.7  | 53.0  | 15.1  | 31.9  | 81.9  | 84.9  | 20,318      |
| 260   | 20   | 93.1  | 1.05   | 97.5  | 54.3  | 14.7  | 31.0  | 82.4  | 85.3  | 19,922      |
|       | 40   | 83.4  | 1.10   | 91.3  | 49.3  | 16.9  | 33.8  | 80.0  | 83.1  | 20,836      |
|       | 60   | 83.4  | 1.13   | 93.9  | 49.1  | 17.0  | 33.9  | 79.8  | 83.0  | 21,419      |
| 280   | 20   | 81.1  | 1.08   | 87.4  | 47.5  | 16.8  | 35.7  | 79.9  | 83.2  | 20,495      |
|       | 40   | 79.9  | 1.12   | 89.6  | 44.6  | 17.7  | 37.7  | 78.8  | 82.3  | 21,324      |
|       | 60   | 77.6  | 1.11   | 86.2  | 45.6  | 17.6  | 36.8  | 79.0  | 82.4  | 21,124      |
| 300   | 20   | 81.8  | 1.11   | 91.0  | 44.7  | 17.3  | 38.0  | 79.4  | 82.7  | 21,163      |
|       | 40   | 73.4  | 1.12   | 82.2  | 40.3  | 18.5  | 41.2  | 77.7  | 81.5  | 21,315      |
|       | 60   | 69.1  | 1.10   | 75.9  | 38.1  | 20.3  | 41.6  | 75.8  | 79.7  | 20,873      |

The results showed that increasing process temperature and time causes the decrease in peat MY from 97.6% to 69.1% at 200 °C and 20 min and 300 °C and 60 min, respectively (Figure A1a). The opposite trend was observed for EDr, where, with the increase in temperature and process time, the EDr values increased from 1.03 for CSF produced at 200 °C and 20 min to 1.12 for CSF produced at 300 °C and 40 min (Figure A1b). Moreover, with increasing temperature, the EY decreased from 100.6% to 75.9% (Figure A1c) for CSF produced at 200 °C and 20 min and 300 °C and 60 min, respectively. The obtained results have a typical tendency of lignocellulosic materials. For comparison, the wood torrefied at the same conditions is characterized by MY of 92–55%, 1.05–1.20 of EDr, and 96–66% of EY [44]. Since peat showed to has smaller mass losses, comparable EDr, and higher EY than wood, it suggests that it is a suitable material for torrefaction. Weight losses are an inevitable element of each torrefaction, and mass losses are related to water removal and partial devolatilization. When during devolatilization, more oxygen than carbon leaves processed material, its high heating value increases, and as a result increase in EDr is observed. The EDr presents how much energy is in the CSF when compared to not torrefied material, and when EDr is higher than 1, the CSF contains more energy than not torrefied material. Typically, torrefaction leads to EDr increase [45] but some materials, mostly not lignocellulosic, have the opposite trend, e.g., sewage sludge [46]. In this study, EY was over 100% at 200–220 °C (Table 1), which is a result of the inaccuracy of empirical measurements. Theoretically, EY cannot be higher than 100% because it would mean that the produced CSF is more energy than in substrate used for its production. As was mentioned above, during torrefaction, devolatilization always occurs, and they take away some chemical energy and move it to gas and liquid fractions.

Devolatilization is a process where large molecules are thermally decomposed by temperature into smaller ones, and when molecules are small enough, they are released from the material by convection forces. During this phenomenon occurs, inter alia, decarbonization, dehydrogenation, deoxygenation [47]. As a result, volatile matter decreases, and ash and fixed carbon content increase. Typical biomass consists of cellulose, hemicellulose, and lignin, and these components affect the properties of biomass and torrefied biomass. The hemicellulose (xylan) consists of 78.57% of VM, 2.11% of AC, and 19.33% of FC. The cellulose consists of 95.21% of VM and only 4.79% of FC. Ash is not present in cellulose. The lignin has the lowest VM of 54.61% but the highest FC of 29.25% and AC of 16.15 [48].
Since in the torrefaction temperatures, mainly hemicellulose decomposes, its properties have the highest effect on torrefied material properties.

The torrefaction process resulted in a decrease in VM from 59.3% for dry unprocessed peat to 38.1% for CSF produced at 300 °C and 60 min (Table 1). At the same conditions, AC increased by ~6% from 14.4% to 20.3%, and FC by ~15% from 26.3% to 41.6%. These changes are characterized by smooth trends, where the temperature has much more effect on observed change than the time (Figure A1d–f). In addition, increasing temperature resulted in a decrease in VS and CP from 83% to 75.8% and from 85.6% to 79.7%, respectively (Figure A1g–h). For comparison, the energetic non-woody biomass is characterized by VM of 70–94%, AC of 1.3–17%, and FC of 2.4–17.2% [49,50], while wood biomass has much homogeneous composition, 82–84%, 15.5–16.4, and 0.3–0.8%, respectively [51]. That means the torrefied peat has less VM and more FC than typical wooden biomasses, but it also has much more AC. Peat, torrefied at 300 °C, becomes like lignite (VM of 39.8%, AC of 8.5%, FC of 51.6%) [52].

3.2. Process Kinetics

3.2.1. TG/DTG and DSC Results

In Figure 2, the results of TG/DTG and DSC analysis are presented. The TG/DTG curves show that the main decomposition of peat and wood starts at similar temperatures, around 200–220 °C; nevertheless, peat turns out to be more resistant for thermal degradation and is characterized by a much lower mass losses in comparison to wood. It is obvious, as peat is more decomposed material than wood. The used peat has a degree of humification (according to the van Post scale) between H4 and H6 [35], meaning that it may be characterized by moderate decomposition. DTG results also showed that peat main decomposition occurred at 200–550 °C and was characterized by two peaks, the first maximum at ~325 °C and the second at ~450 °C. In contrast, wood main decomposition occurred at ~200–380 °C with a maximum at ~330 °C. DSC results are presented in Figure 2b. For both materials, DSC curves are similar up to 300 °C. First, they have one endothermal reaction at ~100 °C and then one exothermal at ~300 °C. At temperatures higher than 300 °C, the peat curve increases more than a woods curve. Moreover, during peat decomposition, one exothermic reaction occurs at 600 °C, while for wood, two endothermic reactions at 478 and 730 °C may be seen.

Figure 2. Thermal analysis results of (a) TG/DTG, (b) DSC.
The observed differences are related to the chemical composition and degree of humification of tested materials. The wood is made of five elements: lignin (18–35%), carbohydrates such as cellulose (40–50%), hemicelluloses (25–35%), organic extractives, and minerals (ash) [53], while peat is an organic matter mixture consists of partially decomposed plants grown in bogs, swamps, and marshes [54]. The organic matter mixture may vary significantly depending on peat age, depth, groundwater level, and so on [55]. For example, Iglovikov et al. [55] analyzed peat soil that consists of bitumens (6–10%), hemicellulose (14–18%), cellulose (1–1.5%), humic acids (28–34%), fulvic acid (15.5–19%), lignin (5–7%), and other organic and inorganic compounds. The study of Brown et al. [56] showed that cellulose and lignin content vary from 14.9% to 46.7% and from 1.1% to 5.2%, respectively.

Typically, the thermal decomposition of hemicellulose, cellulose, and lignin in an inert atmosphere takes place at 220–315, 290–400, and 100–900 °C, respectively [57,58]. Hemicellulose decomposes with two characteristic peaks at ~245 and ~296 °C, while cellulose decomposes at one stage with a maximum of ~345 °C. This means that the decomposition of cellulose occurs intensively in a shorter time than hemicellulose decomposition [48]. This suggests that visible in Figure 2a wood main decomposition at ~200–380 °C with a maximum at ~330 °C is related to hemicellulose and cellulose decomposition, wherein hemicellulose peak is not visible (lack of left shoulder). Some cellulose presence in the peat sample is also visible in the first main decomposition peak at ~325 °C. Nevertheless, lower mass losses in the peat sample at these temperatures suggest that there is a much lower amount of cellulose than in the wood. On the other hand, the studies [55,56] showed much different cellulose concentrations in the peats (1.5% vs. 46.7%). Therefore, there is a need to take caution to state that this decomposition was mainly related to cellulose. Nevertheless, continuous mass losses in both samples at temperatures higher than 400 °C are mainly related to lignin decomposition.

Because peat has a more complex organic matrix, its thermal degradation cannot be described only by three components such as typical biomass. The bitumens that may be constituents up to 10% of the peat [55] decompose in two stages [59]. The study of Murugan et al. [59] shows that the first stage starts at 80 °C and ends at 360 °C with a maximum peak at 295 °C, while the second stage starts at 360 °C and end at 540 °C with a maximum at 445 °C [59]. On the other hand, Santos et al. [60] analyzed humic and fulvic acid thermal degradation. The humic acid was characterized by multiple decomposition peaks at 280, 500, 575, 620, and 800 °C. In contrast, fulvic acid showed to be more resistant to thermal decomposition than humic acids. Fulvic acid was characterized by continuous mass loss with small decomposition peaks at 200, 310, and 545 °C, and one big decomposition peak at 920 °C [60]. These results suggest that bitumens and humic acids were partially responsible for the main decomposition of peat alongside cellulose and hemicellulose at 200–400 °C, and they are the main reason for visible right shoulder at 400–500 °C (Figure 2a). Moreover, Spedding [54] suggests that up to 110 °C free and absorbed water is evaporated from the peat. Above 110 °C, polymerizations of waxes and resins start and lead to a decrease in peat hydrophilic character. At temperatures higher than 160 °C, gases such as CO and CO and colloidal bound water start to be released. From 200 to 320 °C, cellulose, pentosans, and pectin decompose. Since these components contain bound oxygen in their structure, it results in additional water releasing during thermal decomposition, and next, at 280 °C, tars start to be generated alongside other gases.

3.2.2. Peat Thermal Decomposition Kinetics by Coats–Redfern

Table 2 contains kinetic data of peat thermal conversion. The analysis was performed in a nitrogen environment at a heating rate of 10 °C × min⁻¹ using the Coats–Redfern method. The kinetic triplets were calculated for the whole process 30–800 °C and the decomposition peaks. The first peak in both samples is related to water and light volatile organic constituents’ removal [61]. The second peak is related to the main decomposition of compounds that were described in the previous paragraphs. The kinetics analysis revealed that for peat and wood, the first decomposition related to water evaporation was charac-
terized by a reaction order of 3, with an activation energy of 27.11 and 6.67 kJ × mol$^{-1}$, respectively. The analysis also showed that the main decomposition of peat had second reaction order with an activation energy of 33.34 kJ × mol$^{-1}$, while wood main decomposition had a 1.68 reaction order with an activation energy of 65.47 kJ × mol$^{-1}$. The overall whole decomposition process of peat turns out to have n of 1.33 with an Ea of 21.1 kJ × mol$^{-1}$, while wood has n of 1.68 with 65.47 kJ × mol$^{-1}$. All kinetic triplets had a high determination coefficient over 0.9. Only the first peat peak had a slightly lower value. The high value of determination coefficients means that the order of reactions and the activation energy were calculated properly.

### Table 2. Kinetic triplets determined at $\beta = 10 ^{°}C \times min^{-1}$ using the Coats–Redfern method.

| Material, - | Note, - | Temperature Range, $^{°}C$ | $n_r$ - | $E_a$, kJ × mol$^{-1}$ | $A$, s$^{-1}$ | $R^2$ - |
|------------|---------|-----------------------------|---------|------------------------|-------------|---------|
| Peat       | Whole process | 30–800 | 1.33   | 21.10 | $1.97 \times 10^{-2}$ | 0.97   |
| Peat       | First peak   | 30–200 | 3.00   | 27.11 | $1.92 \times 10^{-1}$ | 0.89   |
| Peat       | Main decomposition peak | 200–550 | 2.00   | 33.34 | $4.40 \times 10^{-1}$ | 0.99   |
| Wood       | Whole process | 30–800 | 1.54   | 27.91 | $2.08 \times 10^{-1}$ | 0.91   |
| Wood       | First peak   | 30–200 | 3.00   | 6.67  | $6.86 \times 10^{-5}$ | 0.99   |
| Wood       | Main decomposition peak | 215–430 | 1.68   | 65.47 | $9.05 \times 10^{2}$ | 0.98   |

The provided data cannot be directly compared with other kinetic studies because the peat composition is a complex matrix, and there are many different approaches for kinetics determination. For example, Chen et al. [62] analyzed decomposition kinetics of forest peat under nitrogen atmosphere and described it using reactions of four fractions: moisture, hemicellulose, cellulose, and lignin. The moisture removal had Ea of 61 kJ × mol$^{-1}$ and n of 2.18. The hemicellulose, cellulose, and lignin had Ea of 90, 121, and 65 kJ × mol$^{-1}$, respectively, while the reaction orders were 0.96, 0.85, and 2.31, respectively [62]. On the other hand, Wen et al. [63] analyzed peat moss decomposition and distinguished six stages related to observed decomposition peaks. For $\beta = 10 ^{°}C \times min^{-1}$ and CR method only stages 2–4 were determined at 136–309, 309–379, 379–611, and 611–652 $^{°}C$. For these stages, the Ea changed from 71.3 to 1018.8 kJ × mol$^{-1}$, while the reaction order changed from 0.6 to 2.6 [63]. Similarly, it is with wood. For example, Fraga et al. [64] analyzed the decomposition of wood using the CR method considering only a first-order reaction model (n = 1). The result showed that wood decomposes in two stages. The first stage occurred at 250–390 $^{°}C$, while the second one was at 390–612 $^{°}C$. The Ea for these stages were 190 and 117 kJ × mol$^{-1}$, respectively [64]. Nevertheless, despite the differences in the observed values, and approaches, the results suggest relatively suitable predictions of conversion in these works.

#### 3.2.3. Peat Torrefaction Kinetics by an Isothermal Method

To compare the kinetics of tested peat at torrefaction temperature range, an isothermal experiment using macro-TGA (sample mass of 3 g) was performed according to previous work [41]. The dry peat was heated at temperatures of 200–300 $^{°}C$ (interval 20 $^{°}C$) in 60 min, and then TG data were subjected for constant reaction rates with assumptions of the first-order decomposition. As a result, constant reaction rates (k) for specific temperatures were determined. The higher k, the higher mass losses are observed, or the same mass loss is obtained at a shorter time. Moreover, determined Ea and A can be used to calculate k for any torrefaction temperature. It was found that k values changed from $1.05 \times 10^{-5}$ to $3.15 \times 10^{-5}$ s$^{-1}$ and that Ea and A were $22.02$ kJ × mol$^{-1}$ and $3.41 \times 10^{-5}$ s$^{-1}$, respectively (Table 3). The experimental mass loss in specific temperatures can be seen in Figure A2a, while mass loss predicted using calculated k values in Figure A2b. In the previous work [39], we summarized and recalculated serval waste material torrefaction kinetics by the same method. The graphic of k values at specific temperatures for wood refuse-derived fuel (RDF), sewage sludge, and digestate is shown in Figure A3. The result shows that peat decomposition is much slower than wood but faster than sewage sludge and digestate up to 260 $^{°}C$. Torrefaction at over 260 $^{°}C$ is slower for peat than sewage sludge and
digestate (Figure A3). Meanwhile, RDF torrefaction is slower than peat at all torrefaction temperatures (Figure A3). These results suggest that the thermal decomposition of peat is comparable with other waste materials but differs from the torrefaction of wood.

Table 3. Torrefaction kinetic triplets for isothermal conditions \((T = 200–300 \degree C, t = 60 \text{ min})\) considering a first-order reaction model.

| Temperature, \(\degree C\) | \(k, \text{s}^{-1}\) | \(E_a, \text{kJ} \times \text{mol}^{-1}\) | \(A, \text{s}^{-1}\) | \(R^2, -\) | Reference |
|---------------------------|-----------------|-------------------|-----------------|--------|-----------|
| Peat                      |                 |                   |                 |        |           |
| 200                       | 1.05 \times 10^{-5} |                   |                 |        |           |
| 220                       | 1.82 \times 10^{-5} |                   |                 |        |           |
| 240                       | 2.24 \times 10^{-5} |                 |                 | 22.02  |           |
| 260                       | 2.39 \times 10^{-5} |                 |                 | 3.41 \times 10^{-3} | 0.894   | This study |
| 280                       | 2.70 \times 10^{-5} |                 |                 |        |           |
| 300                       | 3.15 \times 10^{-5} |                 |                 |        |           |
| 220                       | 1.43 \times 10^{-5} |                 |                 |        |           |
| 240                       | 1.82 \times 10^{-5} |                 |                 |        |           |
| 260                       | 3.19 \times 10^{-5} |                 |                 |        |           |
| 280                       | 3.64 \times 10^{-5} |                 |                 |        |           |

| Wood                      |                 |                   |                 |        |           |
| 200                       | 1.43 \times 10^{-5} |                 |                 |        |           |
| 220                       | 3.19 \times 10^{-5} |                 |                 |        |           |
| 240                       | 4.15 \times 10^{-5} |                 |                 |        |           |
| 260                       | 5.37 \times 10^{-5} |                 |                 |        |           |
| 280                       | 7.25 \times 10^{-5} |                 |                 |        |           |
| 300                       | 9.31 \times 10^{-5} |                 |                 |        |           |

3.3. Theoretical Mass and Energy Balance

Table 4 consists of data regarding theoretical torrefaction energy and mass balance. The calculations were performed for peat and wood as reference material. The main goal of the calculations was to show the energy and mass fate during the production of 1 g of CSF at specific conditions. The first and second headings show torrefaction’s specific conditions for which calculations were performed. The third heading contains data of substrate mass used to produce 1 g of CSF at specific conditions. The fourth headings contain data of chemical energy contained in this substrate. The fifth heading is related to external energy provided to the substrate to perform the torrefaction process (here calculated using DSC results). The next one shows the energy contained in 1 g of CSF (here as HHV result). The seventh heading shows the mass of produced gas, and the last heading shows the energy contained in this gas.

The results show that with increasing temperature and time, the mass of substrate needed to produce 1 g of CSF increases. Demand for peat ranged from 1.025 to 1.446 g for 200 and 300 \(\degree C\), while for wood substrate, demand range from 1.042 to 1.831 g. The required mass of peat is lower than in the case of the wood for all torrefaction conditions. It means that to produce the same amount of CSF, less peat is needed than wood. This is the result of mass yield and decomposition differences between processed materials, as raw peat is more humified material than wood.

With increasing substrate mass, increase the energy of the substrate used to produce a unit of CSF. Up to 260 \(\degree C\), the differences in substrate chemical energy for peat and wood are negligible, but over 260 \(\degree C\), much more chemical energy in the wood is consumed to produce 1 g of CSF. The biggest difference occurs at 280 \(\degree C\) in 60 min (24,497 \(J_{\text{sub}} \times J^{-1}_{\text{CSF}}\) vs. 32,177 \(J_{\text{sub}} \times J^{-1}_{\text{CSF}}\) for peat and wood, respectively). This is a result of the high decomposition rate of wood at these temperatures.

The DSC results showed that the external energy required to heat the substrate to setpoint temperature is slightly higher for peat than for wood. The external energy for peat differs from 179 to 633 \(J \times J^{-1}_{\text{CSF}}\), while for wood differs from 136 to 578 \(J \times J^{-1}_{\text{CSF}}\). The differences probably result from a difference in substrates composition and its specific heat. The specific heat of peat and wood depends on temperature and moisture. Assuming that both materials were completely dried, the peat should have a higher specific heat than wood because it required more energy. Nevertheless, results of specific heat in the literature provide overlapping data for peat and wood. For example, specific heat for dry peat differs from 0.9 to 1.3 \(J \times (g \times K)^{-1}\) [65] and for dry wood from 1.1 to 1.45 \(J \times (g \times K)^{-1}\) [66,67].
| Temp. °C | Time, min | Peat Wood | Peat Wood | Peat Wood | Peat Wood | Peat Wood | Peat Wood | Peat Wood | Peat Wood | Peat Wood | Peat Wood |
|---------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| 200     | 20        | 1.025     | 1.042     | 19,491    | 19,068    | 179       | 136       | 19,598    | 18,551    | 0.025     | 0.042     | 71        | 654       |
|         | 40        | 1.027     | 1.071     | 19,533    | 19,596    | 179       | 136       | 19,941    | 18,960    | 0.027     | 0.071     | −229      | 773       |
|         | 60        | 1.030     | 1.088     | 19,576    | 19,898    | 179       | 136       | 19,841    | 19,178    | 0.030     | 0.088     | −87       | 856       |
|         | 20        | 1.016     | 1.063     | 19,326    | 19,440    | 224       | 185       | 19,619    | 18,753    | 0.016     | 0.063     | −70       | 871       |
| 220     | 40        | 1.049     | 1.125     | 19,939    | 20,589    | 224       | 185       | 20,272    | 19,312    | 0.049     | 0.125     | −109      | 1462      |
|         | 60        | 1.097     | 1.139     | 20,849    | 20,830    | 224       | 185       | 20,217    | 19,440    | 0.097     | 0.139     | 857       | 1575      |
|         | 20        | 1.034     | 1.144     | 19,666    | 20,931    | 289       | 251       | 19,444    | 19,496    | 0.034     | 0.144     | 511       | 1684      |
| 240     | 40        | 1.103     | 1.256     | 20,965    | 22,981    | 289       | 251       | 20,341    | 20,077    | 0.103     | 0.256     | 913       | 3155      |
|         | 60        | 1.141     | 1.290     | 21,685    | 23,604    | 289       | 251       | 20,318    | 20,371    | 0.141     | 0.290     | 1657      | 3484      |
| 260     | 20        | 1.074     | 1.133     | 20,425    | 20,720    | 379       | 336       | 19,922    | 19,456    | 0.074     | 0.133     | 883       | 1600      |
|         | 40        | 1.199     | 1.391     | 22,804    | 25,442    | 379       | 336       | 20,836    | 20,767    | 0.199     | 0.391     | 2347      | 5011      |
|         | 60        | 1.200     | 1.554     | 22,809    | 28,428    | 397       | 336       | 21,419    | 21,136    | 0.200     | 0.554     | 1770      | 7627      |
| 280     | 20        | 1.233     | 1.308     | 23,447    | 23,932    | 495       | 446       | 20,495    | 20,467    | 0.233     | 0.308     | 3447      | 3911      |
|         | 40        | 1.252     | 1.531     | 23,809    | 28,012    | 495       | 446       | 21,324    | 21,432    | 0.252     | 0.531     | 2980      | 7026      |
|         | 60        | 1.288     | 1.759     | 24,947    | 32,177    | 495       | 446       | 21,124    | 21,646    | 0.288     | 0.759     | 3868      | 10,976    |
| 300     | 20        | 1.223     | 1.621     | 23,248    | 29,653    | 633       | 578       | 21,163    | 21,457    | 0.223     | 0.621     | 2718      | 8773      |
|         | 40        | 1.363     | 1.829     | 25,918    | 33,454    | 633       | 578       | 21,315    | 22,034    | 0.363     | 0.829     | 5236      | 11,998    |
|         | 60        | 1.446     | 1.831     | 27,498    | 33,493    | 633       | 578       | 20,873    | 22,002    | 0.446     | 0.831     | 7257      | 12,069    |

*Value determined using DSC analysis result; **value determined using calorimetric analysis result (HHV); ***value is the sum of chemical energy contained in gas and heat from external energy, if no external energy stays in CSF.
The results showed that for both materials, the energy contained in CSF increases with process temperature, which is mainly a result of oxygen removal in the devolatilization process [47]. As a result of progressing mass loss during torrefaction at higher temperatures, an increase in gas production is visible. The gas production for peat ranges from 0.025 to 0.446 g of gas per 1 g of CSF, while for wood, from 0.042 to 0.831 g of gas is produced per 1 g of CSF made from wood. Although, gas constitutes a significant amount as a by-product, especially at 300 °C. It contains much less energy than CSF. Nevertheless, when gas energy is higher than the external energy needed to produce CSF, the process may be self-sufficient (assuming gas combustion). In this study, the results suggest that the production of CSF from wood can be self-sufficient at any torrefaction conditions, while peat requires temperatures higher than 240 °C. However, cautions should be kept with that result interpretation because data show theoretical calculations for dry material and lack of energy losses related to the use of specific torrefaction technology. The peat used in this study had around 85% moisture content, and it can be easily calculated by using latent heat of evaporation (2257 J × g⁻¹H₂O at 100 °C). For water evaporation of 1% from any material, there is a demand for 22.57 J of additional energy (assuming dryer efficiency at 100%). It means that for the presented case (peat with 85% of moisture content) for each torrefaction condition, over 1918.5 J per processed gram of fresh peat should be added. As a result, any torrefaction conditions do not provide a self-sufficient process.

For the above, presented results are the initial step and base for future calculations when specific conditions and technology (moisture of processed substrate, drying technology, e.g., mechanical, conventional, solar, mixed drying, torrefaction process, e.g., batch or continuous reactors) will be considered. Furthermore, because the calculations were made with wood materials as a reference, currently available technology for wood torrefaction/pyrolysis can be easier recalculated and redesigned (if needed) for peat.

This paper presents the initial test on medical waste peat torrefaction. We intended to check the influence of temperature and residence time on the basic fuel properties of CSF. However, here we intend to draw the possible applications and needs for additional characterization of CSF. We recommend that the next experimental work should also cover more detailed characterization of CSF following techniques FTIR, XRD, SEM, TEM, and BET, to check other possible applications of CSF, i.e., soil additive, adsorbent.

4. Conclusions
In this work, for the first time, medical/cosmetic peat waste was torrefied, and the effect of torrefaction temperature and time on CSF quality was determined. Parallelly, decomposition of peat waste was determined according to the CR method using TGA equipment (temperature range 30–800 °C) and torrefaction kinetics using macro-TGA equipment (temperature range 200–300 °C) were determined. In addition, macro-TGA was used for lifetime prediction. Moreover, theoretical torrefaction mass and energy balances of peat waste were determined. As a result, the full process of peat torrefaction with its effects was presented. The main findings of performed analyses were:

- Torrefaction improves fuel properties of peat, HHV increase from 19.0 to 21.3 MJ × kg⁻¹;
- Peat main decomposition takes place at 200–550 °C following second reaction order (n = 2), with an activation energy of 33.34 kJ × mol⁻¹ and pre-exponential factor of 4.40 × 10⁻¹ s⁻¹;
- Macro-TGA shows that peat decomposition at torrefaction temperature is comparable with other waste materials but differs from wood, for which the constant reaction rates (k) are much lower;
- Theoretically, torrefaction of dry peat can be self-sufficient at temperatures higher than 240 °C, but future calculations with specific technology need to be performed to find real conditions of self-sufficient peat processing.

Since peat has already been used worldwide, technologies of its use exist. Therefore, there should be no problem finding a place for CSF made from medical/cosmetic peat.
waste. Moreover, CSF has similar energetic properties to lignite, and as a result, it can be considered as co-fuel to the lignite incineration plant or for gasification for energy or hydrogen purposes. In addition, since biochars are used as partial replacements in agriculture, it may be an additional place for biochar made from peat. Whether medical peat waste will be used for energetic uses or agricultural one, the work provides useful information about its processing and main properties.

Another problem, which has not been arisen here, but should be further studied, is that the distribution of peat waste is very dispersed; therefore, small-scale torrefaction systems should be designed and tested.

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Abbreviations

A  Pre-exponential factor
AC  Ash content
AIC  Akaike information criterion
ASTM  American Society for Testing and Materials
CSF  Carbonized solid fuel
CP  Combustible parts
CR  Coats–Redfern method
DSC  Differential scanning calorimetry
Ea  Activation energy
EDr  Energy densification ratio
EU  The European Union
EY  Energy yield
FC  Fixed carbon
HHV  High heating value
HTC  Hydrothermal carbonization
MC  Moisture content
MY  Mass yield
n  Reaction order
PW  Peat waste
R²  Determination coefficient
SPA facility  A place devoted to a renewal of mind, body, and spirit by different methods
TGA  Thermogravimetric analysis
VM  Volatile matter
VS  Volatile solids
Appendix A

Table A1. Standards and methods used for fuel properties determination.

| Properties                  | Standard/Method               | Equipment                                                                 | Ref. |
|-----------------------------|------------------------------|---------------------------------------------------------------------------|------|
| Moisture content (MC)       | PN-EN 14346:2011             | Dryer (WAMED, model KBC-65W, Warsaw, Poland)                             |      |
| Volatile matter (VM)        | TGA method                   | TGA—tubular furnace (Czylok, RST 40 × 200/100, Jastrzębie-Zdrój, Poland) coupled with a laboratory balance (RADWAG, PS 750.3Y, Warsaw, Poland) | [36] |
| Ash content (AC)            | PN-Z-15008-04:1993          | Muffle furnace (Snol 8.1/1100, Utena, Lithuania)                           |      |
| Fixed carbon (FC)           | by difference (calculation)  | Muffle furnace (Snol 8.1/1100, Utena, Lithuania)                           |      |
| Volatile solids (VS)        | PN-EN 15169:2011             | Muffle furnace (Snol 8.1/1100, Utena, Lithuania)                           |      |
| Combustible parts (CP)      | PN-Z-15008-04:1993          | Muffle furnace (Snol 8.1/1100, Utena, Lithuania)                           |      |
| High heating value (HHV)    | PN-Z-15008-04:1993          | Calorimeter (IKA, C200, Staufen, Germany)                                 |      |

MY(T,t) = 71.8242 + 0.400445 × T − 0.00121217 × T^2 − 0.000727896 × T × t
R^2 = 0.94, AIC = 311.04

VM(T,t) = 0.543524 × T − 0.00129347 × T^2 + 0.273621 × t
− 0.00137518 × T × t
R^2 = 0.96, AIC = 238.01

Figure A1. Cont.
Figure A1. Torrefaction temperature and time effect on properties of CSF produced from medical/cosmetic peat waste, the equations boundary, $T = 200–300 \, ^\circ C$, $t = 20–60 \, \text{min}$. (a) mass yield, (b) energy densification ratio, (c) energy yield, (d) volatile matter, (e) ash content, (f) fixed carbon, (g) volatile solids, (h) combustible parts, (i) high heating value.
Figure A2. Results of macro-TGA, the effect of isothermal temperatures in time (a) experimental data, (b) lifetime prediction.

Figure A3. The constant reaction rates (k) in function of temperature for different materials at specific temperatures [39].

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