Potassium and soot interaction in fast biomass pyrolysis at high temperatures

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ABSTRACT

This study aims to investigate the interaction between potassium and carbonaceous matrix of soot produced from wood and herbaceous biomass pyrolysis at high heating rates at 1250°C in a drop tube reactor. The influence of soot carbon chemistry and potassium content in the original biomass on the CO2 reactivity was studied by thermogravimetric analysis. The XPS results showed that potassium incorporation with oxygen-containing surface groups in the soot matrix did not occur during high temperature pyrolysis. The potassium was mostly found as water-soluble salts such as KCl, KOH, KHCO3 and K2CO3 in herbaceous biomass soot. The low ash-containing pinewood soot was less reactive than the potassium rich herbaceous biomass soot, indicating a dominating role of potassium on the soot reactivity. However, the catalytic effect of potassium on the reactivity remained the same after a certain potassium amount was incorporated in the soot matrix during pyrolysis. Raman spectroscopy results showed that the carbon chemistry of biomass soot also affected the CO2 reactivity. The less reactive pinewood soot was more graphitic than herbaceous biomass soot samples with the disordered carbon structure.

1. Introduction

Entrained flow gasification (EFG) is a promising technology, which generates high quality syngas and small amounts of tars due to the high operating temperatures. However, the high soot yields lead to intensive gas cleaning and can cause a possible plant shut down. The reduction of soot formation increases the overall production system efficiency and improves the economic feasibility and reliability of the gasification plant. Moreover, the alkali metals released from high ash-containing lignocellulosic materials may form molten ash material, and thus, lead to deposition of slag upon the reactor wall [1,2]. Continuous extraction of the ash slag from the reactor is required to prevent outlet blockages and to ensure steady syngas production.

The released alkali metal ions in the biomass pyrolysis participate in soot formation and conversion reactions. It is known that alkali metal ions enhance CO2 reactivity of graphite and carbon black [3] and suppress soot formation [4]. The alkali metal ions can form a fly ash or be bonded to oxygen-containing surface groups in the soot matrix [5,6]. The fly ash particles contain K2S, KOH and K2CO3 [6]. The interaction of K2CO3 with the low ash-containing coal was investigated under CO2 gasification condition in a thermogravimetric analyzer by Kopycsinski et al. [7]. The pyrolysis results showed that CO was released from K2CO3, which was bound to a carbon surface. In the next step, the oxygen from the carbonate reacts further with the surface carbon and forms CO which leaves the reduced potassium complex of an unknown stoichiometry. In the last step, the transfer of the potassium cluster occurs. During fast pyrolysis, the elemental potassium is unlikely to be released in gas phase at low temperatures. However, little is known about the interaction between the carbonaceous matrix of soot and alkali. In catalytic gasification, the concentration of intercalated species is small, and it decreases with increasing temperature [8,9]. The elemental potassium, bonded to the soot matrix in phenolate groups remained inaccessible for the gaseous reactants, leading to the low catalytic activity [10]. The potassium-carbon complexes form series of stable and metastable phases with different physicochemical properties [11–14]. The K-C60 complex was investigated using 13C and 39K solid state NMR [15]. According to the 13C NMR spectra of intercalated graphite (K1C60) it exists as a single phase characterized by a resonance...
at 174 ppm at temperatures above 420 K [13]. At lower temperatures, two resonances of intercalated graphite at 187 ppm (K 3C60) and 174 ppm (KδC60,δ≤1) were identified by 13C NMR [13]. X-ray photoelectron spectrum of potassium doped double-walled carbon nanotubes showed two peak components at the binding energy of 293.3 eV and 296 eV (K 2p3/2 and K 2p1/2 of the K-oxides and K-cations) [16]. The X-ray photoelectron bands at 294.6 and 293.4 eV were assigned to ionic potassium intercalated within the graphite layers and to oxidized potassium at the surface of graphite [17]. The modeling using density functional theory method (DFT) confirmed the existence of an electron transfer from soot to gaseous oxygen through the active K* sites [18]. In this DFT model, soot structure was represented by graphene layers and potassium was modeled perpendicular to the plane of the soot model and near the zigzag and armchair edges of the soot. The DFT model for the C-C armchair face showed that the potassium intercalation in phenolate groups (C-O-K) inhibits the gasification reaction, confirming previous experimental observations [3,19].

This work aims to investigate the interaction between potassium and carbonaceous soot matrix in detail. The specific objectives of this study were to: (1) investigate the interaction of potassium with the carbonaceous matrix of soot produced at high temperatures (1250°C) in a drop tube reactor, and (2) understand the formation mechanism of potassium-carbon complexes in fast pyrolysis and their effect on the soot reactivity using Raman spectroscopy, X-ray photoelectron spectroscopy (XPS), 13C solid state NMR, X-ray diffraction (XRD), and thermogravimetric analysis (TGA).

2. Materials and methods

2.1. Raw biomass characterisation

Pinewood, beechwood, wheat straw and alfalfa straw were chosen for the fast pyrolysis study in a drop tube reactor (DTF). Fuel selection was based on the differences in the ash composition and plant cell compounds (cellulose, hemicellulose, lignin, extractives), as shown in the supplemental material (Table S-1). The ultimate and proximate analysis of pinewood, beechwood, wheat straw, leached wheat straw and alfalfa straw is shown in Table 1.

2.2. Pyrolysis with drop tube furnace

Soot and char samples were obtained from pyrolysis experiments in a drop tube reactor (DTF). Soot samples were generated at 1250°C to obtain a maximal soot yield. The DTF setup and operating conditions were described in detail by Trubetskaya et al. [20]. The reactor consists of an alumina tube (internal diameter: 54 mm, tube length: 1.06 m) heated by four heating elements with independent temperature control. Gas flow rate into the reactor is regulated by mass flow controllers (EL-flow Select, Bronkhorst High-Tech B.V.). The experiments were conducted by feeding 5 g of biomass at a rate of 0.2 g min⁻¹. Both primary (0.18 l min⁻¹ measured at 20°C and 101.3 KPa) and secondary (4.8 l min⁻¹ measured at 20°C and 101.3 KPa) feed gases were N₂. The residence time of fuel particles was estimated to be about 1 s, taking into account density changes during pyrolysis. Soot particles passing the cyclone (cut size 2.5 μm) were captured from the product gas flow by a grade QM-A quartz filter with a diameter of 50 mm (Whatman, GE Healthcare Life Science).

2.3. Solid residue characterisation

2.3.1. Ash compositional analysis

The ash compositional analysis was performed by ICP-OES (DIN EN 15290). Prior to the analysis, soot samples were pre-heated in oxygen at 10°C min⁻¹ up to 550°C and kept at that temperature for 7 h.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Proximate and ultimate analysis.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuel</td>
<td>Pinewood</td>
</tr>
<tr>
<td>Moisture, (wt% as received)</td>
<td>5.1</td>
</tr>
<tr>
<td>Ash (550°C, wt % dry basis)</td>
<td>0.3</td>
</tr>
<tr>
<td>Volatiles, (wt% dry basis)</td>
<td>86.6</td>
</tr>
<tr>
<td>HHV, (MJ kg⁻¹)</td>
<td>21.6</td>
</tr>
<tr>
<td>LHV, (MJ kg⁻¹)</td>
<td>20.2</td>
</tr>
</tbody>
</table>

2.3.2. Potassium ion selective electrode

The potassium ion-selective membrane electrode Orion 93–19 (Thermo Scientific, USA) was used to determine the amount of leached potassium in deionized water. The electrode potential was measured at room temperature using the Benchtop Meter Orion 720A (Thermo Scientific, USA). Five-points calibration was conducted using potassium chloride solutions with ionic strengths of 10⁻⁰, 10⁻¹, 10⁻², 10⁻³, 0.01 and 0.1 M.

2.3.3. X-ray diffraction

The XRD analysis of the soot matter was conducted using a Bruker D8 AXS X-ray diffractometer with (Cu-Kα, λ = 1.54056 Å and Cu-Kz2 radiation, λ = 1.54439 Å) operating in Bragg–Brentano (reflection) mode, using a secondary graphite monochromator, and a scintillation detector, in the range 5 to 80 degrees for 12.5 h. Soot samples were placed in small cups which were rotated during the XRD analysis. The multiple fitting of XRD pattern, crystallite size analysis and peak searches were done using PowderPlot software and Crystallographica Search-Match software (Version 3.1,0.0). The instrumental reflection broadening was subtracted from the experimental pattern by the PowderPlot software.

2.3.4. Raman spectroscopy

Raman spectroscopy was performed using an iNvia Raman microscope (Renishaw, UK) operating with a 514 nm laser line at a power of 30 mW. The measurements were performed in static mode with a centre at 1600 cm⁻¹ resulting in a 960–2200 cm⁻¹ spectral region. The laser power was set to 100% in the software and roughly 30% in the hardware. 1 s exposure time was used in normal confocality mode. A 20x lens and 8–15 μm step size (X and Y directions) were used for mapping, to generate 100–200 spectra/image for each soot sample. Deconvolution of the Raman spectra were conducted using the peak fit pro tool in the OriginPro software (OriginLab, USA) by combination of five Gaussian-shaped bands (D4, D3, D2, D1, and G) following Sadezky et al. [21] and as described in the supplemental material (Section S-10.1). The mean crystal size in the a-direction (Lₐ) with the fitting constants C₀ = −12.6 nm and C₁ = 0.033, which are valid for the laser
wavelength from 400 to 700 nm, is given by [22]:

\[
\lambda_\text{d} = \frac{C_\text{d} + C_\text{f}/A_\text{d}/A_\text{f}}
\]

2.3.5. Thermogravimetric analysis

The reactivity of soot samples in 20% volume fraction CO2 was determined using a thermogravimetric instrument Q5000 (TA Instrument, USA) by loading ca. 8 mg of sample in an Al2O3 crucible and heating from 30 to 1200°C in CO2 at a constant heating rate of 10°C min⁻¹. The kinetic parameters of soot samples were derived by the integral method presented by Coats and Redfern [23]. The reactivities of biomass soot samples were compared using reaction rates calculated from the derived kinetic parameters (A and E_a) at a fixed gasification temperature of 1000°C.

2.3.6. ¹³C solid state NMR spectroscopy

Solid-state NMR analysis was carried out on a Nuclear Magnetic Resonance spectrometer, Avance 400 (Bruker, Germany) (9.4 T) operating at Larmor frequencies of 400.13 and 100.58 MHz for ¹H and ¹³C, respectively. All experiments were conducted using a double resonance probe equipped with 4 mm (o.d.) rotors. Samples were analyzed without any additional preparation at room temperature by single-pulse (SP) magic angle spinning (MAS) as well as cross polarization (CP) MAS [24] utilizing high-power ¹H two-pulse phase-modulated decoupling (TPPM) [25] during acquisition and employing a spinning rate of 9 kHz. The ¹³C CP/MAS spectra were recorded using a recycle delay of 8 s, a contact time of 1 ms, an acquisition time of 45.9 ms and 4096 scans, whereas the ¹³C SP/MAS spectra were recorded using a recycle delay of 128 s, an acquisition time of 45.9 ms and 1080 scans. All ¹³C NMR spectra were referenced to the carbonyl resonance in an external sample of α-glycine at 176.5 ppm [26].

2.3.7. X-ray photoelectron spectroscopy

The XPS spectra were collected with an electron spectrometer AXIS Ultra DLD (Kratos Analytical Ltd, UK) using monochromated Al Kα source operated at 120 W. Analyser pass energy of 160 eV for acquiring survey spectra and a pass energy of 20 eV for individual photoelectron lines were used. The surface potential of dielectric potassium benzoate was stabilized by the spectrometer charge neutralization system. The C 1s component for benzene ring (sp² carbon) of potassium benzoate sample was set to 284.7 ppm [27]. The binding energy scale of the spectrometer was calibrated using Au 4f7/2 line of metallic gold at 84.0 eV. Processing of the spectra was accomplished with the Vision 2 software (Kratos Analytical Ltd, UK).

3. Results

3.1. Solubility of potassium in char and soot from herbaceous biomass

Preliminary investigations showed that the ash content of pinewood and beechwood soot samples remained unchanged after the washing in deionized water [20]. However, the removal of water-soluble potassium from herbaceous biomass soot led to a lower reactivity in CO2 gasification. Fig. S-7 illustrates that the washing of soot and char from alfalfa straw and wheat straw does not remove all potassium found in any of the soot samples formed under fast pyrolysis conditions. The relatively small fraction of potassium remaining in the washed soot from alfalfa and wheat straw suggests the formation of the potassium-carbon complexes or remaining non-soluble potassium fraction.

3.2. Reactivity of biomass soot

Fig. 1 shows differential weight loss curves (DTG) for the 20% volume fraction CO2 gasification of soot samples. The DTG curves show a single broad peak in CO2 gasification, indicating a heterogeneous soot mixture with respect to the composition [28]. The CO2 gasification of all soot except pine and washed soot from wheat straw took place at nearly the same temperature range from 830 to 900°C. The additional DTG peaks in the CO2 gasification of alfalfa straw soot were related to the reactivity of heavy hydrocarbon compounds [29]. The maximum reaction rate of pinewood soot was about 200°C higher than for the other biomasses, and thus, the pinewood soot was clearly less reactive. The washing of wheat straw soot led to a lower reactivity. The maximum reaction rate of washed soot of wheat straw in the CO2 gasification was shifted to temperatures about 100°C higher than for the soot from pyrolysis of leached wheat straw and non-treated wheat straw. The calculated tmax/tlag ratio showed that the gasification reaction in the TG analysis was influenced only by chemical kinetic limitations, as shown in the supplemental material (Table S-3).

3.3. ¹³C solid state NMR spectroscopy

By ¹³C solid-state MAS NMR it was demonstrated that all soot samples were characterized by a predominantly graphene-like structure [30], as shown in the supplemental material (Figure S-9). Only in alfalfa straw soot an additional presence of a carboxylate carbon was detected. The chemical shift of this agrees well with that of the potassium-calcium carbonate butschliite [31-33].

3.4. X-ray photoelectron spectroscopy

The XPS spectra of pinewood and beechwood soot mainly exhibit the carbon and oxygen lines with some traces of sodium, whereas herbaceous biomass soot samples additionally showed K 2p and Cl 2p lines, as shown in the supplemental material (Table S-4). The K and Cl atomic concentrations for the alfalfa straw soot were 3.5 and 1.2 at.%, whereas the K and Cl atomic ratio in pure KCl is 1:1, respectively. The remaining potassium (2.3 at.%) might have been bounded to the soot matrix as KOH and/or K2CO3 which was not detected (or fitted) in C 1s spectrum due to small amounts. The present XPS results strongly suggest that potassium species were incorporated in straw soot particles mostly as water-soluble KCl, K2CO3 and KOH.

3.5. Raman spectroscopy

Raman spectroscopy was carried out to examine primary differences
in the carbon structure of soot samples. The calculated integrated peak area ratio ($A_D/A_G$) in supplementary Table S-5 showed that the beechwood and leached wheat straw soot samples obtained the highest amount of carboxylates (0.2 and 0.3), whereas the $A_D/A_G$ ratio of wheat straw soot was the lowest (0.04). The relative $A_D/A_G$ ratio of pine wood soot is lower than for beechwood soot due to the low content of acetyl groups in hemicellulosic fraction of softwood [34]. All soot samples based on the $A_D/A_G$ ratios exhibited a common structure of amorphous carbon and nano-crystalline graphite, as discussed by Ferrari and Robertson [35]. In addition, the alfalfa straw soot contained a higher fraction of distorted small PAH clusters within the amorphous carbon than other soot samples, as observed experimentally by Aboud et al. [36]. The average extensions of graphene stacks ($L_g$) from the Raman bands in beechwood, wheat straw, alfalfa straw and leached wheat straw soot were lower than those of pine wood and washed soot from wheat straw. The size of one aromatic ring is 2.5 Å [37], and therefore, the size of PAHs ($L_g = 21–26$ Å) is equivalent to the size of 8–10 aromatic rings.

3.6. X-ray diffraction

The XRD analysis did not show any significant differences between woody and herbaceous soot samples in terms of graphitization, as shown in the supplemental material (Figure S-41). The XRD analysis of soot indicated formation of turbostratic or random layer lattice structures. The additional reflections, detected by the XRD measurements of wheat straw and alfalfa straw soot, were attributed to KCl and KHCO₃.

4. Discussion

The thermogravimetric experiments demonstrated significant differences in CO₂ reactivity for soot from pyrolysis of wood and herbaceous biomass. The CO₂ gasification of beechwood, wheat straw, alfalfa straw and leached wheat straw soot prepared at 1250 °C took place at nearly the same temperature range, whereas the maximum reaction rate of pine wood soot and washed soot from wheat straw was shifted to higher temperatures, indicating a lower reactivity. The activity of soot samples can be affected by the differences in carbon structure and alkali content [6,38,39]. Lápurna et al. [40] studied soot samples generated from the combustion of biodiesel and diesel fuels using thermogravimetric analysis and Raman spectroscopy and the authors found that the impact of soot nanostructure on the oxidation reactivity is significant. The biodiesel soot has a higher initial degree of graphitization with the higher curvature of the carbon fringes of an average particle size which increases the probability of collapsing into smaller fringes, and thus, enhances the reactivity. Trubetskaya et al. [41] studied the impact of lignocellulosic compounds and monolignols on the biomass soot reactivity and showed that the high content of extractives and lignin in the pine wood and beechwood could lead to the lower reactivity of woody soot. The previous studies also showed that the maximal reaction rates of extractives and lignin soot samples were less reactive than the cellulose and hemicellulosic soot, emphasizing a dominating role of differences in lignocellulosic composition on the soot reactivity. Interestingly, the authors found that the CO₂ reactivities of softwood and wheat straw lignin soot were similar beside the fact that the low ash-containing softwood lignin soot was expected to be less reactive than the Na⁺ rich wheat straw lignin soot.

In the present study, the Raman spectroscopy analysis showed that the alfalfa straw soot consists of small PAH clusters within the amorphous carbon with high curvatures of carbon fringes forming higher specific surface area, leading to the higher reactivity. In addition, the alfalfa and wheat straw soot samples obtained the highest $A_D/A_G$ ratio. Thus, more defective graphene sheets could form smaller carbon segments with a higher specific surface area resulting in the high CO₂ reactivity. This indicates that the carbon structure has an influence on the observed differences in soot reactivity. In addition, the previous Raman spectroscopy analysis indicated that the differences in carbon structure of soot from holocelluloses, lignin and monolignols were small [41]. However, the ash-forming elements are known to influence both the shape and nanostructure of pine wood soot particles during the entrained flow gasification of biomass [6].

In the present work, the Raman spectroscopy results suggested that the released alkali metal ions in the biomass pyrolysis had an influence on the soot carbon structure, leading to the formation of defects in the carbon matrix. Previous results showed that the soot CO₂ reactivity depends mainly on the alkali content in the original fuel and less on the soot nanostructure and soot particle size [42]. For the Ca and Na rich biodiesel soot, higher oxidation reactivities were observed than for the low ash-containing farmesan and diesel soot samples [43]. The authors also indicated that Ca, Na, and P compounds had a dominant role on the soot reactivity compared to the differences in nanostructure of biodiesel and diesel soot samples. Gustafsson et al. [44] observed that Ca was a dominant ash-forming element in wood pellet combustion. In the present study, the ash content in the original beechwood (1.4%) was higher than in the pinewood (0.3%), leading to the higher K release to the gas phase and so more K incorporation in the beechwood soot particles compared to the pinewood pyrolysis. The ash content in the original alfalfa straw (7.2%) was also higher than in the non-treated wheat straw (4.1%) and leached wheat straw (2.1%). The inorganic elements in all herbaceous biomass soot samples were mainly K, Cl, S, and Si. Previous equilibrium calculations and X-ray diffraction suggestions suggested an incorporation of water-soluble KCl in the alfalfa, non-treated and leached wheat straw soot samples [45]. The present results showed that 50% more of potassium was condensed onto the leached wheat straw soot compared to the non-treated wheat straw soot where potassium was probably retained as a silicate in the char. The lower Cl content in the leached wheat straw might indicate that potassium was released in the form of KOH, possibly forming K₂CO₃ [46]. Based on the XRD and XPS analysis results it was concluded that the potassium species were incorporated in the alfalfa and wheat straw soot particles mostly as KOH, KCl, KHCO₃ and K₂CO₃. The present results showed that potassium compounds represent major alkali metal ions incorporated into the biomass soot matrix. In addition, the XPS results showed that the elemental potassium was not bonded to the soot matrix in phenolate groups and was mainly adsorbed as potassium hydroxide and/or potassium carbonate on the herbaceous soot particle surfaces which is probably due to the short contact time in high-temperature biomass pyrolysis. The XPS results also indicated that the alkali metal compounds were not detected for pinewood soot. The water-soluble salts affect the biomass soot reactivity [42]. Soot from alfalfa straw, wheat straw, beechwood and leached wheat straw was 14 times more reactive than pine wood soot and washed soot from wheat straw, due to the stronger catalytic effect of ash compounds. Significantly smaller differences in CO₂ reactivity were observed for the pinewood soot and washed soot from wheat straw (2 times), as shown in the supplemental material (Table S-2). The amount and composition of the ash forming matter in the lignocellulosic materials lead to the differences in soot reactivity. The potassium content of leached wheat straw soot was lower than for alfalfa straw and wheat straw soot, and higher than for beechwood soot. However, the differences in reactivity of soot from beechwood, wheat straw, alfalfa straw and leached wheat straw were small, as shown in Fig. 2. The remaining potassium compounds in the washed soot from wheat straw led to a slightly higher reactivity compared to the pine wood soot. Thus, the present study indicated that the water-soluble salts determine the catalytic gasification rate because electron donor–acceptor (EDA) complexes and C-O-K groups on the soot surface were not formed. The high reactivity of potassium rich herbaceous soot samples was related to the reduction-oxidation cycles of K₂CO₃, elemental potassium, and K₂O according to Eqs. (2)-(4) [47,48]:

$$K_2CO_3 + 2C \rightarrow 2K + 3CO_2$$

(2)
Fig. 2. Maximal reaction rate at 1000°C (results from Table S-2) versus potassium content in soot from pine, aspen, pine straw, alfalfa straw, leached wheat straw and washed soot from wheat straw (results from Figs. S-5-S-7). The potassium content is shown in wt% relative to the soot (g g⁻¹ on dry basis).

\[
2K + CO_2 \rightarrow K_2O + CO \quad (3)
\]

\[
K_2O + CO_2 \rightarrow K_2CO_3 \quad (4)
\]

In addition, the catalytic effect of potassium on the reactivity remained similar after a certain concentration of potassium is reached in the soot particles, and thus, the changes in reactivity of soot from pyrolysis of alfalfa straw, wheat straw and leached wheat straw were small. The removal of potassium from the original wheat straw does not affect the soot reactivity. In contrast, the estimated activation energy of wheat straw containing potassium was 102% higher than that of wheat straw without potassium, indicating that the removal of potassium leads to a lower soot reactivity. Fig. 2 shows that the reactivity of soot samples generated from dry feedstocks with potassium concentration > 0.3 wt% changed only slightly. The contact between alkali metals and carbon is an essential parameter in the catalytic gasification of biomass that can be improved by mobility of the alkali metals [49]. The present results showed that the K₂CO₃ had a good contact with the soot carbonaceous matrix, leading to the higher CO₂ reactivity with the increased content of potassium until saturation was reached.

5. Conclusion

The novelty of this work relies on the fact that potassium is unlikely to be bonded to oxygen-containing surface groups in the soot matrix during high temperature pyrolysis. The present results showed that potassium was deposited on the soot surface as water-soluble alkali such as KCl, KOH, KHCO₃ and K₂CO₃. The thermogravimetric analysis showed that the differences in CO₂ reactivity are large for soot from wood and herbaceous biomass pyrolysis. The CO₂ reactivity of soot depends mainly on the potassium content in the original fuel and less on the carbon chemistry. However, the catalytic effect of potassium on the reactivity remains the same after a certain potassium amount was deposited on the surface of soot particles during pyrolysis. The low ash containing pine straw soot was less reactive with the more ordered graphene structure than other soot samples. The carbon structure of alfalfa and wheat straw soot appeared more graphene disordered and contributed to the greater reactivity in CO₂ gasification. The present work emphasized a significant influence of both potassium and carbon structure on the biomass soot reactivity.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at http://dx.doi.org/10.1016/j.fuel.2018.03.140.

References


