1 CHEMICAL FRACTIONATION OF INORGANIC CONSTITUENTS IN ENTRAINED

2 FLOW GASIFICATION OF SLURRY FROM STRAW PYROLYSIS

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Highlights:

- 21 1. Significant amount of K is volatized during gasification at 1400 °C
- 22 2. The amount of K is depending on the chemical composition and phase distribution of the gasified
- 23 straw char
- 3. The reaction of the ash components in the straw char is delayed by the vaporization of the
- 25 surrounding liquid phase
- 4. The non-equilibrium conditions lead to different reaction pathways at the edge and in the center of
- the char particles

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ABSTRACT:

Pressurized entrained-flow gasification (PEFG) of straw biomass is currently being studied as a potentially sustainable and economically viable process to produce fuels and other vital chemicals. In the process chain the gasification is integrated and straw is converted via pyrolysis into a bioslurry consisting of a liquid, tar-rich phase and char. Afterwards, the slurry is gasified into a tar-free, low-methane syngas which is a basic reactant for the synthesis of biofuels. At the high temperatures over 1200 °C the ash constituents of the char in the bioslurry melt and flow down the inner wall as slag. The slag viscosity has to be in a certain range to form a protective layer at the reactor wall and to guarantee a continuous removing. For this reason, the composition of the molten ash at the reactor wall has to be well known. Due to several fractionation processes in the gasifier the composition of the slag at the reactor wall does not correspond directly with the slurry ash. Therefore, experiments were conducted to identify depletion and enrichment processes in the gasifier. Finally, the composition of the slag at the reactor wall is obtained and can be used for the adjustment of the viscosity.

Keywords: gasification, ash fractionation, slag, viscosity, biochar, straw

1 INTRODUCTION

Biofuels are a CO₂-neutral alternative to conventional fuels which are based on fossil energy carriers. The synthesis of biofuels is achieved by Biomass-to-Liquid (BtL) technologies. Low-grade biogenic feedstocks are upgraded to high-grade synthetic fuels with high energy density. An example of a BtL-process is the bioliq process developed at the Karlsruhe Institute of Technology [1]. The basic resource is residual straw from agriculture. Because of its low energy density, straw has to be converted into an intermediate energy carrier. The technique used is fast pyrolysis at temperatures around 500 °C under inert-atmospheric conditions. This process results in solid char, liquid and aqueous condensates and non-condensables. The mixed condensates and char form a slurry that is used for feeding the high temperature pressurized entrained flow gasifier in the following process step [1-3].

In the gasifier temperatures over 1200 °C and pressures of 40-80 bars are dominant. The resulting tarfree and low methane syngas is used as the basic reactant for synthetic biofuels and chemicals [4-6]. The high temperatures cause the ash constituents of the slurry to melt. The formed slag layer flows down the reactor wall, is separated in a quench at the outflow and removed continuously. The slag also protects the refractory material at the reactor wall against corrosion. Therefore, the viscosity of the slag at the reactor wall has to be in a certain range to prevent stalling of the protective layer at low viscosities and avoid blockages on the reactor outlet at high viscosities [7]. In the literature a typical viscosity range of 10-25 Pa*s is suggested for slag inside entrained flow gasifiers [8].

The viscosity is primary influenced by temperature and chemical composition [9-12]. For reliable calculation of the slag viscosity, these two factors have to be well known. The temperature at the gasifier wall can be calculated by CFD and is subject of other studies [13-15]. However, the chemical composition of the slag at the reactor wall is not necessarily identical with the chemical compositions of the feedstock ash. Varying chemical composition of different straw feedstocks influence the reaction chains concerning the release and fate of volatile inorganic species [16], which in turn influence the composition and thus the viscosity of the resulting slag. Furthermore, fractionation processes occur during the gasification of the slurry in the reactor flame. Thus, knowledge about the fractionation of inorganic constituents related to the gasifier conditions and different straw compositions are mandatory for predicting the composition of slag inside the reactor, based on which its viscosity can be calculated. Therefore, the aim of the present investigation was to identify the relevant fractionation processes depending on ash composition. The results will later be used to develop a model for predicting slag composition and viscosity based on the ash composition of the fuel and the process parameters.

2 MATERIALS AND METHODS

2.1 Materials

Table 1

The char samples investigated in this study are obtained from four types of straw collected in 2002, namely Spanish oats (Avena sativa, denominated as H2), Spanish winter barley (Hordeum vulgare, H5), Spanish carinata (Brassica carinata, H7), and Danish wheat (Triticum aestivum, H10). The straw samples, comprising hulls, stems, and leafs, were shredded (<5 cm), homogenized and stored under dry conditions at room temperature. For the present investigations, samples were pyrolysed at 550 °C in inert atmosphere (N2) in a first step. The chemical analyses of the chars were performed by the Central Division of Chemical Analysis (Forschungszentrum Jülich). Major elements (C, H, N, and S) were analyzed by a CHNS analyzer (Tab. 1). Moisture and ash content were determined by mass loss during drying at 105 °C and ashing in air at 550 °C, respectively. The oxygen was calculated by balancing to 100%. The amount of chlorine was determined by ion chromatography. The inorganic fraction was analyzed by inductively coupled plasma—optical emission spectroscopy (ICP—OES). The normalized ash composition is shown in Figure 1.

Figure 1

- Each char sample has a characteristic chemical composition illustrating the wide diversity of straw concerning the distribution of inorganic constituents. The sample H2 is characterized by high SiO₂ and CaO contents. H5 is enriched in K₂O and Cl. In H7, CaO und K₂O are the main components. H10 is dominated by SiO₂ and K₂O.
- Alkaline metals (K, Na) influence the viscosity of the molten ash and volatize under gasification conditions. In the straw samples, K is the most abundant alkaline metal (Fig.1). The distribution between the accompanying components Si, Ca, Cl will influence the release behavior of K significantly [17-19]. Therefore, the study will mainly focus on the content of these four elements.

2.2 Sample Preparation and Gasification

- The char from the previous pyrolysis step was grinded to a size <1 mm. According to the process conditions of the bioliq $^{\odot}$ -process a model fuel consisting of 20 mass $^{\%}$ pyrolysis char as solid phase and 80 mass $^{\%}$ ethylenglycol ($C_2H_6O_2$) as liquid phase was created [20]. 200 mg of slurry were filled into a Pt-sample boat in each experiment.
- Several investigations used the high-temperature furnace reactor combined with the MBMS instrument [19, 21, 22]. The experimental setup in this study is based on these previous investigations

(Fig. 2)

Figure 2

The gasification experiments are conducted in an electrically heated furnace including a high-density alumina tube to prevent reactions of the tube walls with the released species. The sample boat was fixed on a ceramic rod, inserted into the preheated furnace and kept there for varying retention times from 10 s to 50 s. The temperature was adjusted to 1400 °C in the reaction zone for all samples investigated in this study. Furthermore, a continuous gas flow of 20% CO₂ and 80% He was realized corresponding with 0.6 L/min CO₂ and 2.4 L/min He.

2.3 Analysis of the gasified char

The chemical analysis of the gasified solids was conducted by the Central Division of Chemical Analysis (Forschungszentrum Jülich). Changes of the chemical composition of each sample between the initial char and the residual char after gasification at varying retention times were analyzed and presented. The results were normalized by the content of Ca, which is abundant in all samples and released in negligible amounts into the gas.

released in negligible amounts into the gas.

Additionally, mineral phases of the initia

Additionally, mineral phases of the initial and gasified char were characterized by powder X-Ray-Diffraction (XRD). The high content of amorphous carbon in the sample leads to a high background in the XRD-diffractrograms and mineral phases with minor X-ray reflexes (e.g. silicates) were overlaid. Therefore, identification was restricted to semi-qualitative results, which describe the relative amount of the main mineral phases. Although the conventional oxidation of ash for analysis would minimize

the background in the spectra, the method is not suitable in this case because it may change the phase

170 content.

Scanning electron microscopy (SEM) was conducted at a voltage of 7kV. The integrated EDX-detector enabled the local determination of the chemical composition. The distribution of inorganic components in the char was observed and differentiated between the retention times of the samples in the furnace. The samples were dried at 250 °C and coated with Ir before SEM analysis.

2.4 Gas Analysis (MBMS)

 MBMS-analysis was used to determine the composition of the evolving gas phase during the gasification process. The inlet of the MBMS device is directly connected to the outlet of the alumina tube furnace. The gas phase enters the first vacuum chamber of the instrument through a 0.3 mm nozzle. After the two following vacuum chambers a pressure of 100 nPa is reached and the gas phase is analyzed by mass spectrometry. The detailed setup of the MBMS is described in Bläsing et al. [17, 18]. In this study, CO/CO₂ and K as main fragment of potassium species was qualitatively and semi-quantitatively measured. The reduction of CO₂ by the glycol and carbon of the char during the reaction

results in the formation of CO [23]. Thus, the start and the end of reaction of the glycol and char can be identified. The kinetics of the reactions of the ash components are recorded by the volatilized K in the gas phase.

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3 THEORY

- The viscosity of biomass slags is strongly dependent on the incorporation of alkaline metals (Na, K) 191 [24]. Because alkaline metals are volatile at high temperatures, alkali fractionation is an influencing 192 process to the slag viscosity [25-30]. In straw biomass, K is the main alkali metal in the organic matrix 193 and in inorganic salts such as KCl or K₂CO₃ [16, 31-34]. Previous studies estimate that <20% of K are 194 195 released at temperatures <700 °C as KOH or in combination with tars from organic bonded K in the straw matrix [35, 36]. Inorganic salts are stable below 700 °C [26, 31, 32]. In the temperature range of 196 197 700-900 °C evaporated K is attributed to the decomposition of KCl, K₂CO₃ and remaining organicbonded K in the char [31, 37]. When temperatures exceed 900 °C, KOH is formed by the reaction of 198
- K_2CO_3 with steam and released [32, 37].
- The remaining K at these temperatures is retained in silicates. Some silicates are already present in the raw straw or formed by ion exchange during the decomposition of K-salts at lower temperatures. The
- surrounding organic matrix can limit the formation of new inorganic compounds in char during
- 203 gasification [26, 32].
- A part of S is volatized at temperatures <700 °C. Another part remains in the char by forming K₂SO₄
- 205 [32]. At temperatures >1000 °C sulfates start to evaporate. At reducing conditions, the decomposition
- 206 temperature shifts below 500 °C [37, 38].
- Due to the abundance of KCl in straw biomass the release pathway of Cl also influences the
- fractionation of K. While 40-60% Cl evaporates at temperatures below 700 °C [26, 31, 32], <20% K is
- released in this temperature regime. The released Cl can originate from oxygen-rich, functional groups
- of organic and inorganic compounds and forms HCl in the gas phase. At temperature above 700 °C the
- remaining amount of Cl is released from decomposing KCl [26, 32]. The reaction pathways are
- 212 influenced by the composition of the surrounding gas atmosphere, the residence times and
- composition, the particle size and porosity of the ash particles [16, 31, 37].
- The hot gas phase can be online monitored by using molecular beam mass spectrometry (MBMS).
- Recent studies focus on the different release of inorganics from coal, straw, and blends under
- combustion or gasification related atmospheres [17, 18, 39, 40]. Devolatilization and char reactions
- were distinguished at several times. Devolatilization is characterized by relatively short-term,
- 218 intensive reactions and is related to pyrolysis or gasification conditions in these investigations. Char
- reactions take place after the devolatilization with lower intensity in the MS-spectra and are even in
- 220 gasification related experiments, where oxygen is the gasification agent, similar to combustion-like
- 221 conditions under experimental conditions [18]. To minimize oxygen excess during char reactions,

investigations that are more recent used CO₂ as gasification agent as it is very common in investigations on char conversion kinetics [23].

Whereas most recent studies investigate the conversion of solid straw or coal in combustion or inert pyrolysis environment at temperatures up to 1000 °C [26, 31, 34, 35], the studies in gasification environment were not focused on straw biomass [16, 41, 42]. The actual study fills this gap by investigating the release behavior of a suspension consisting of solid straw char and liquid glycol under gasification related conditions. The wide diversity of straw composition is considered by different chemical composition of ash-forming components such as Si, K and Ca. The reaction pathways and kinetics of the release were investigated by varying residence times under constant temperatures around 1400 °C. The gained qualitative information of this study will provide knowledge about the chemical fractionation processes in the conversion zone of an entrained-flow gasifier. This knowledge will be used to develop and improve a flowsheet model of the gasification process, which will provide a calculated chemical composition of the slag at the gasifier wall. This chemical composition will be used for further viscosity determinations.

4 RESULTS

4.1 Release Behavior

The four investigated straw samples (H2, H5, H7 and H10) show similar behavior during the release experiments. The glycol evaporates and reacts immediately with CO₂ when the sample enters the high temperature zone of the furnace. The reaction leads to a steep decrease of CO₂ and an increase of H₂O in the MBMS-spectrum during the first 10 s (Fig. 3). After ca. 15 s the amount of CO₂ increases in the gas because most of the glycol and its containing carbon have reacted. Since the char reacts much slower with CO₂ than the glycol, the CO₂ concentration during char gasification is higher. As long as the volatile part of the char, which contains the hydrogen, is converted, the H₂O concentration remains high. After ca. 30 s the reactivity of the residual char decreases slowly, therefore CO₂ further increases and H₂O decreases slowly. At the end of the measurement, the CO₂ concentration has not reached the inlet concentration because the char is not completely converted at that time.

After ca. 12 s the first part of potassium is released, shown by an increase of K-content in the gas, indicating the beginning reactions of the solid char, which contains the potassium. Thus, the release of K starts 2 s after the vaporization and reaction of glycol characterized by the decrease of CO₂ in the gas. Consequently, it can be assumed, that the surrounding glycol delays the reaction of the inorganic

sample.

components. After 20 s a second significant amount of K is released during char gasification of the

Figure 3

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4.2 Char Characterization

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SEM-images confirm the presence of glycol after 10 s in the remaining char. The structures shown in Fig. 4 were caused by the suspension of glycol and char in the slurry. The glycol was evaporated during sample preparation directly before the SEM-analysis. The agglomerated inorganic particles remain from the initial structures and are observed in the SEM-images. After residence times longer than 10 s all glycol has reacted during the gasification and the suspension structures disappear.

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Figure 4

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The main ash components (CaO, K_2O and SiO_2) of the investigated chars are displayed in Fig. 5. All four samples show a depletion of K_2O after reaction at 1400 °C. The amount of released K changes between the residence times pointing out different reactions at certain stages of the gasification. A significant depletion of K takes already place in all samples after 10 s of gasification by evaporating the K from the sample surface. The MBMS spectrum confirms this result showing a strong K-release after 10-15 s (Fig. 6).

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Figure 5

Figure 6

Figure 7 shows the mineral phases present in the chars after varies residence times determined 280 qualitatively by XRD. It gives information on the association of ash components and their reactions 281 during gasification. The samples H2 and H7 are characterized by high Ca-contents (Fig. 5). The Ca is 282 primary associated with CaCO₃ in the two samples, which is confirmed by the results in the overview 283 of the XRD-results (Fig. 7). The decomposition of CaCO₃ starts after 10 s and results in the 284 intermediate formation of CaO. At the same time, the atmosphere leads to the reduction of sulfates, 285 such as K₂SO₄. The oxygen of these sulfates is reduced and the sulfur is retained in sulfides, such as 286 CaS with the Ca of the CaO. 287 Relevant amounts of K-phases were not identified by XRD in the initial char of H2 and H7 and might 288 be incorporated in the C-matrix or in mineral phase with weak reflexes (e.g. silicates), which are 289 overlaid by the x-ray amorphous carbon. The main difference between H2 and H7 is the higher K-290 content in H7, which cannot be associated with Si, Cl or Ca (ratio in Tab. 1). K in H7 might be 291 incorporated in K₂SO₄ and char matrix while K in H2 can occur in silicates and with Ca. Therefore, 292 H2 is characterized by a significant K-loss after 10 s in comparison with H7 (Fig. 5), which is also 293 visible in the MBMS-spectrum of the gas output (Fig. 6). 294

The K remaining in the samples H2 and H7 forms K-carbonates after 20-30 s (Fig. 7). The C originates from the char matrix and from the decomposing CaCO₃. The carbonates decompose with increasing retention time. The Ca remains as CaO or CaS in the char. The K is released continuously into the gas phase in H2. The low amount of Si in H7 cannot keep the K in the sample. Additionally, the higher amount of Ca is preferred to be incorporated in silicates and replace K from the stable phases [43]. Therefore, the strong K-release occurs after 40 during gasification of H7 and can be assigned to decomposition of intermediate formed K-carbonates (Fig. 5).

Figure 7

The untreated samples H5 and H10 contain significant amounts of K and Cl, which are incorporated in KCl (Fig. 7). In H5 the K/Cl ratio is lower than in H10, which indicates that more Cl is associated with KCl (Tab. 1). The vaporization of KCl in H5 results in a strong decrease of Cl and K in several stages after 10 s, 40 s and 50 s (Fig. 8). A similar decrease of K was also observed in analysis of H10 and is confirmed with its MBMS spectrum (Fig. 3). The K in H5 is also associated with K₂SO₄. The decomposition of this phase results in the formation of CaS, similar to the process in H2 and H7. H10 has the highest Si-content of the samples. The formation of K-carbonates was not observed for this sample, which may indicate the preferred incorporation of K in silicates instead of carbonates (Tab. 1). However, most K is still present as KCl until 30 s and is then released directly into the gas phase instead of being retained in the sample to form K-silicates (Fig. 7). Therefore, pure SiO₂ is still present in H10 after 40 s retention.

Figure 8

Most of the inorganic components are distributed in small aggregates in the char matrix containing different elements (Ca, Mg, Na, K, Si, S, etc.). The random distribution of these aggregates results in discontinuous reaction pathways during the gasification. Thus, the location of the inorganic particles has a significant influence to the reaction kinetics. A part of Si occurs in separate grains (Fig. 9A). Furthermore, the physical structure of the straw remains visible. The plant species of H5 and H10 incorporate high Si-amounts in layers around their structure which can be seen in Fig. 9A/B after 40 s retention time. EDX analysis of these could show an incorporation of Ca, K and Na in these Si-layers with increasing residence times. The presence of pure SiO₂ in XRD and SEM after retention of 40 s indicates, that this incorporation is not complete after long residence times and might be inhibited caused by the surrounding C-matrix.

Figure 9

5 DISCUSSION

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5.1 Devolatilization and char reactions

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According to Bläsing et al. [18] the gasification process can be divided into devolatilization and char reactions. Devolatilization is the dominating process after 10 s retention time in this study. Glycol and volatile components like K at the char surface are instantly evaporated which is shown in the MBMS-spectra. The endothermal evaporation of the surrounding glycol results in a cooling of the solid particles and delays the reactions in the char particles. After 20 s char reactions have a stronger influence on the reaction pathways of the samples and the gas phase composition. Mineral phases in the char decompose and form new phases which incorporate and release volatiles. The diffusion of the elements can be suppressed by the surrounding matrix [38]. Therefore, the release pathways are highly depended on the chemical composition and the distribution of inorganic components in the straw or char samples.

In the initial samples, K is incorporated in the identified phases KCl and K2SO4 and some

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5.2 Potassium

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overlaid and unidentified phases. The presence of Cl in H5 and H10 indicates the occurrence of KCl, which is a major source for release of K in these samples, although the time of KCl vaporization is different for the two samples. The mineral vaporizes at 700 °C [31, 32, 35, 36], which is achieved fast at the surface of the sample at temperatures of 1400 °C. Potential reaction partners cannot retain K in the char because the still existing C-Matrix inhibits the incorporation and the inorganic partners are randomly distributed. Therefore, a significant amount of K evaporates as KOH(g) or K(g) during the first 10 s [43, 44] and exceeds the amount of K predicted by the thermodynamic equilibrium calculation. Another part of K remains as KCl in the solid char until 30 s retention time (Fig. 7), although the salt should vaporize completely at the temperatures of 1400 °C. The temperature gradient delays inorganic reactions to the center of the single char particles and of the whole granulate. The effect is supported by the inhibition of inorganic diffusion processes by the surrounding C-matrix [31, 32, 37]. Kcarbonates are formed as intermediate products, which are characterized by higher vaporization temperatures than KCl. As a result, the carbonates occur at higher residence times up to 50 s and keep additional K in the sample. The formation of K-carbonates is connected with silica [26, 32]. In sample H10 with high amount of silica, the K tend to be incorporated in silicates instead of forming

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structure over long retention times.

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carbonates which correspond with the literature [32, 33, 38]. More stable silicates fix the K in their

5.3 Silica

Silicates might occur in the initial char, but identification by XRD is limited due to the background caused by the high C-content. The decomposition of salts and carbonates release K and Ca. A certain amount is incorporated in silicates and remains in the char at long residence times during the experiment [37, 45, 46]. Another silica phase is sand, which most likely entered the sample during sample collection. The grains were identified in the SEM.

The incorporation of K in the Si layer structures, which surround the char particles, was observed in this study according to Knudsen et. al. [32]. The significant amount of Si will result in the formation of silicates, which incorporate K, Ca and inorganic components by substitution [43, 47]. The concentration of Si in these layers at the edge of the char particles delays the formation of these silicates because of the longer migration paths. Additionally, K can be replaced by more stable Ca, which increases the release of K into the gas phase. The occurrence of Si in straw influences the release behavior significantly because available K prefers to be incorporated in high temperature-stable silicates.

5.4 Application to the condition in the bioliq®-gasifier

 The atomized fuel in the bioliq[®]-gasification has a reactive surface in the order of magnitudes higher than the samples of the present experiments. Therefore, reactions will proceed much faster in the real gasifier than in the present batch-type experiments. However, the residence times in the gasifier of 2-6 s are also significantly shorter [48]. Thus, the principal impact of the effects on the inorganic compounds will be similar.

The liquid phase is volatized in the zone of droplet evaporation in the gasifier. The endothermic reaction implies a cooling effect shown in the experiment and also appears in the gasifier causing an average droplet temperature in the gasifier around 400 °C [7]. Consequently, the delay of inorganic reactions is also present in the gasifier, although it appears in a smaller time scale.

After the volatilization of the ethylenglycol, a part of K is released from the char surface according to the K-release after 10 s in the experiments of this study. As a result, the volatized K is not available to influence the viscosity at the reactor wall, because the released K immediately leaves the reactor with the gas phase.

The C-matrix still existing after 40 s in the experiment shows the impact of the C-matrix to inorganic reactions during char gasification. The short residence times in the flame of the gasifier also imply an incomplete conversion of the char particles resulting in a C-matrix, which surrounds the particles in the gasifier for a certain time. According to the results of this study, the C-matrix will imply reducing conditions to the inorganic components and inhibits the diffusion of the inorganic components in the zones of the gasifier. Thus, intermediate products, e.g. K-carbonates, are formed during the gasification process and will reach the reactor wall influencing the chemical composition of the slag,

e.g. by bringing additional K into this system. Additionally, the carbonates can form molten salts at the reactor wall and influence the flow behavior of the slag, which is dominated by oxide melts containing silica [43]. The degree of char conversion is depending on the chemical composition, the distribution of the inorganics in the ash and surrounding the liquid phase.

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6 CONCLUSIONS

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Inorganic components are influencing the viscosity of the evolving slag at the inner wall of the gasifier significantly. They are present in the char of the feedstock and thus surrounded by a C-matrix. Due to the short residence time of the feedstock in entrained-flow gasification, this C-matrix remains present around the inorganic particles in the char over several zones in the gasifier. This study shows a delay of inorganic reactions caused by the C-matrix and the impact to the reaction condition in the char particles. Thus, the C reduces the inorganic oxides resulting in oxygen-free inorganic compounds like CaS. The solid char is surrounded by liquid ethylenglycol. Its vaporization is an endothermic reaction, taking place before the reaction of the char and cooling down the temperature at the char particles. Consequently, the presence of the liquid phase delays inorganic reactions in the char by 2 s in the experiments. In the gasifier, the delay is shorter but still has to be considered due to the shorter residence times. The amount of K will have a strong influence on the flow-behavior of the slag during the gasification process. The network modifier K is a main inorganic component in straw biomass and will decrease the viscosity of the slag in the entrained flow gasifier. In addition, the alkaline metal is partly released into the gas phase before reaching the gasifier wall, where the slag is formed. In this study, several stages of K-release were defined referring to the accompanying components e.g. Si and Ca and the conditions of reaction. The first release stage occurs after 10 s in all samples of this study and is assigned to the direct release of K from the samples surface. During this process, additional K is released because inorganic reaction partners are not available due to separation by the surrounding C-matrix. This effect is strong at the surface of the sample because temperature conditions change fast from room temperature to 1400 °C there and minimize the chance to reach reaction partners, which keep K in the char. The additionally released K is thus not available for influencing the viscosity at the gasifier wall. The following stages of release are depended on the bonding conditions of accompanying components and appear different in the four investigated samples. While for the sample H2 the K is released continuously after the surface reaction, the other three samples show a second significant release between 20 s and 40 s. The differences of the K-release are referred to the formation of different intermediate phases due to the varying chemical composition. Additionally, the temperature gradient

- 443 the formation the intermediate phases. The intermediate phases can incorporate additional K in the
- char, which upon reaching the gasifier wall increase the K-content of the slag. The limiting factor is
- the local availability of reaction partners, which is depended on the randomly distributed inorganic
- aggregates in the char samples.

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Table 1: Chemical composition und element ratios of the investigated char samples

Mass fraction (%)	H2	H5	H7	H10
Char components				
С	69.54	61.01	64.94	64.3
Н	3.21	3.57	3.55	2.95
N	1.78	0.78	0.93	1.1
Moisture	1.58	4.57	4.48	2.90
Ash	11.31	17.65	12.31	18.96
O (difference)	12.57	12.42	13.78	9.79
Σ	100	100	100	100
Ash components	I			
Cl	0.06	2.489	0.13	0.739
Al	0.0094	0.0053	0.00633	0.07
Ca	2.664	0.72	1.63	1.283
Fe	0.0234	0.00401	0.0061	0.0685
K	2.44	6.51	6.5	5.41
Mg	0.236	0.2493	0.135	0.218
Na	0.455	0.94	0.0772	0.0693
P	0.393	0.1008	0.361	0.2645
S	0.219	0.373	0.33	0.28
Si	0.972	1.613	0.032	3.7
	1	1	<u>'</u>	
Element ratios (molar	basis)			
K/Si	1.8	2.9	145.8	1.0
K/Cl	37.0	2.4	45.4	6.7
K/Ca	0.9	9.3	4.1	4.3
Ca/Si	1.9	0.3	35.6	0.2

Figure 10: Ash composition of the char samples (oxygen free composition calculated to 100 mass-%)

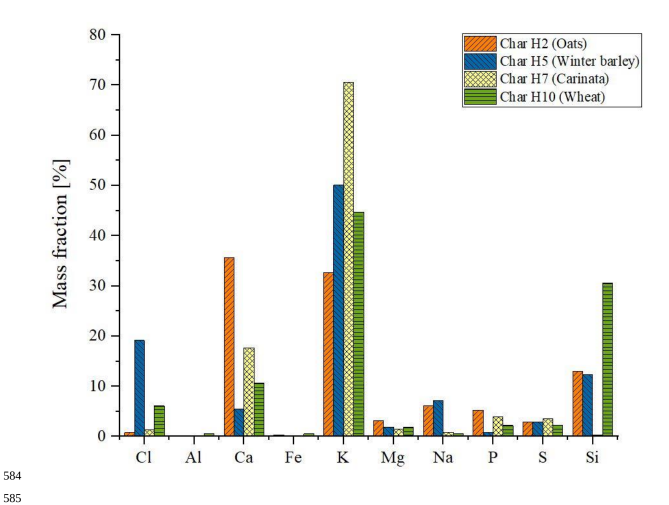


Figure 11: Schematic drawing of the MBMS instrument

588 589

Multiplier Detector addition heating additional 5-zone furnace Al₂O₃-rod Turbopump 3 Skimmer 10⁻⁶ 10⁻² 0 10° mpar mbar mbar gas inlet He, CO₂ Ionisation Molecular Al_2O_3 -Pt-boat Turbopumps 1+2 Beam tube with slurry Deflector

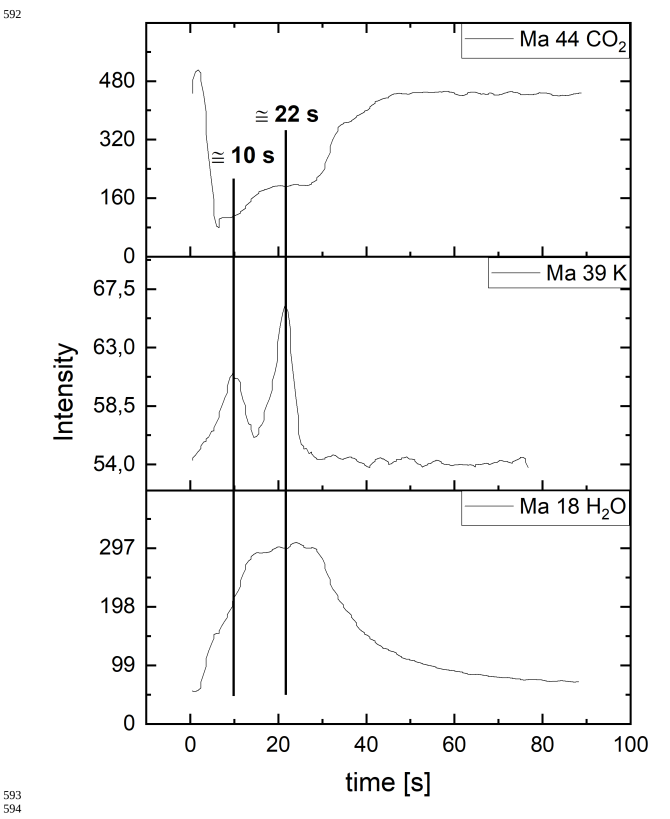


Figure 13: Structures of suspension consisting of remaining ash constituents in the char of H5 after 10s in SEM-images (BSE)

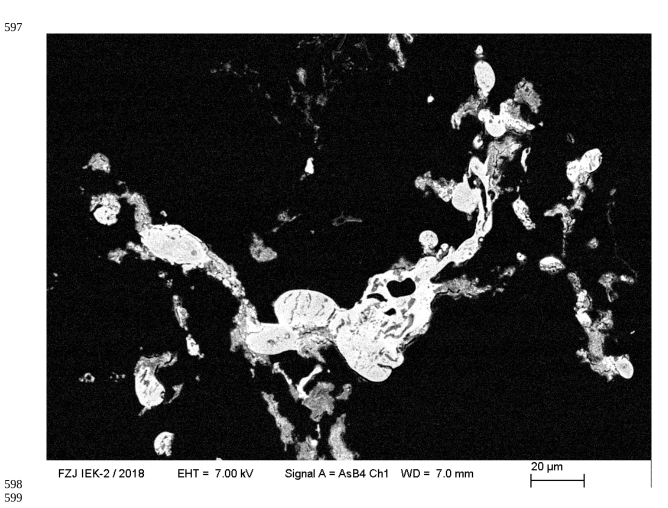
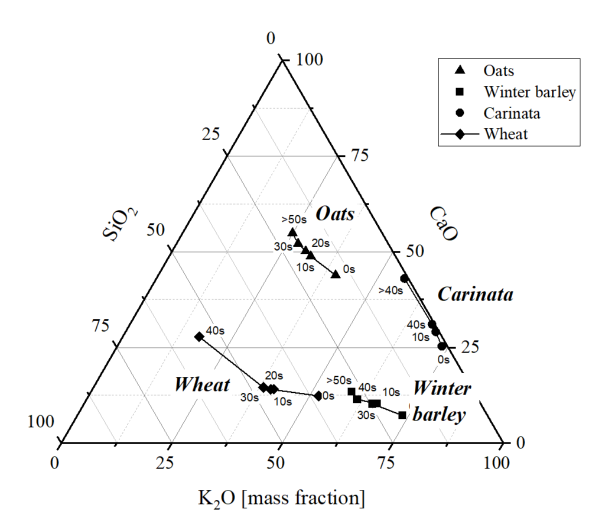


Figure 14: Concentration of the three main ash components in the char of the four straw samples H2, H5, H7 and H10 after various residence times





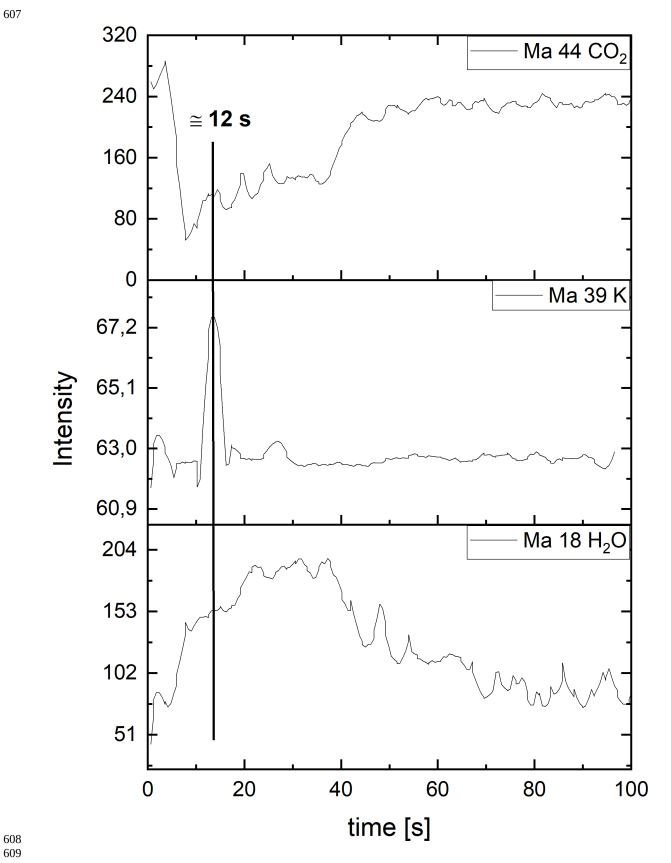


Figure 16: Phase distribution in the chars of the four straw samples H2, H5, H7, and H10 after varied residence times determined qualitatively by XRD for each sample

	0s	10s	20s	30s	40s	>50s
	CoCO			CaO		1
H2	CaCO ₃					CaS
			I		K(HCO ₃)	
		KC1				
Н5	K ₂ SO ₄		1		CaS	
	112004					K ₂ CO ₃
	K ₂ SO ₄					
H7		CaCO ₃			CaO	
		$K_2Ca(CO_3)_2$		CaS		
				K(HCO ₃)		
			SiO ₂			
H10		KCl				
					CaS	

Figure 17: Chemical composition of char H5 after various residence times and divided by the Cacontent

