



Article

# Experimental Parameter Study on Synthesis Gas Production by Steam-Oxygen Fluidized Bed Gasification of Sewage Sludge

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Featured Application: The experimental data can be applied for designing or modeling of fluidized bed sewage sludge gasifiers for synthesis gas production.

**Abstract:** The conversion of biogenic residues to fuels and chemicals via gasification and synthesis processes is a promising pathway to replace fossil carbon. In this study, the focus is set on sewage sludge gasification for syngas production. Experiments were carried out in a 20 kW fuel input bubbling fluidized bed facility with steam and oxygen as gasification agent. In-situ produced sewage sludge ash was used as bed material. The sensitivity of the key operation parameters gasifier temperature, oxygen ratio, steam to carbon ratio, and the space velocity on the syngas composition ( $H_2$ , CO,  $CO_2$ ,  $CH_4$ ,  $C_xH_y$ ,  $H_2S$ , COS,  $NH_3$ , and tars) was determined. The results show that the produced syngas has high  $H_2$  and CO concentrations of up to 0.37 m<sup>3</sup> m<sup>-3</sup> and 0.18 m<sup>3</sup> m<sup>-3</sup>, respectively, and is thus suitable for synthesis of fuels and chemicals. By adjusting the steam to carbon ratio, the syngas'  $H_2$  to CO ratio can be purposely tailored by the water gas shift reaction for various synthesis products, e.g., synthetic natural gas ( $H_2/CO = 3$ ) or Fischer–Tropsch products ( $H_2/CO = 2$ ). Also, the composition and yields of fly ash and bed ash are presented. Through the gasification process, the cadmium and mercury contents of the bed ash were drastically reduced. The ash is suitable as secondary raw material for phosphorous or phosphate fertilizer production. Overall, a broad database was generated that can be used for process simulation and process design.

**Keywords:** gasification; sewage sludge; syngas; sulfur; tar; ammonia; biofuel; synthesis stoichiometry; operation parameters; nutrient recovery; circular economy



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# 1. Introduction

The mitigation of climate change requires the substitution of fossil carbon in carbonaceous fuels and goods by renewable carbon sources. Renewable carbon sources are, for instance, biomasses, such as wood or energy crops. However, the current available biomass feedstock cannot satisfy the massive amount of fossil carbon that is currently utilized worldwide. Therefore, also the utilization of carbon from available biogenic residues and waste streams has to be considered in order to replace as much fossil carbon as possible. Sewage sludge is a not avoidable residue with a quantity of over  $10 \times 10^9$  kg a<sup>-1</sup> on dry basis in Europe [1], over  $12 \times 10^9$  kg a<sup>-1</sup> in the USA [2] (1999 only  $7 \times 10^9$  kg a<sup>-1</sup> [3]), and over  $6 \times 10^9$  kg a<sup>-1</sup> in China with a steep increase [4]. Besides the utilization of the energy and carbon contents of sewage sludge, the recovery of phosphorous (e.g., from sewage sludge ash) to close the nutrient cycle and the safe recycling or disposal of hazardous components such as heavy metals are also of great importance [5]. Therefore, a change from the currently mostly practiced sewage sludge disposal methods, e.g., land-"use", landfill, and co-incineration in coal-fired power plants or cement plants, to mono-treatment recycling methods is required in order to fulfill the goals of a circular economy and climate change mitigation. For instance in Germany, the need of phosphorous recovery has already

been put into regulation [6]. The utilization of sewage sludge as fuel is also economically favorable as high disposal fees are attainable.

A conversion process is needed to practically replace fossil carbon with renewable carbon in established products such as transport fuels, chemicals or plastics. A promising and very flexible conversion process considering feedstock quality and product choice is the thermochemical gasification that generates a high calorific syngas which can be further converted to the desired product in a downstream catalytic synthesis [7]. Such a gasification process is the steam-oxygen fluidized bed gasification, that produces a nitrogen-lean and thus high calorific syngas, that is rich in hydrogen and carbon monoxide, and thus can be used for synthesis of fuels (e.g., synthetic natural gas, dimethyl ether, kerosene) and chemicals (methanol, plastic monomers) [8]. Also, the recovery of phosphorous can be achieved well from gasification-derived sewage sludge ash [9–11].

As gasification agent, a mixture of steam and oxygen is used to provide the necessary heat for the endothermic gasification process through partial fuel oxidation. Therefore only a single fluidized bed reactor is needed, which simplifies the process layout in contrast to the allothermal indirect dual fluidized bed (DFB) steam gasification [12,13]. The needed oxygen can be generated by an on-site power-to-gas facility that produces oxygen as by-product from water electrolysis, by a state of the art cryogenic air separation unit (ASU) with acceptable energy consumption of 720 kJ per kg  $O_2$  [14,15] or tentatively in future by air separation with membranes. The sewage sludge has to be dried in order to be utilized in fluidized bed gasification. To minimize the energy consumption for drying, a combination of mechanical dewatering (e.g., centrifuge) to a dry matter fraction of around 0.25 kg kg<sup>-1</sup>, solar drying, and thermal drying using low temperature heat will finally obtain a dry matter fraction of around 0.9 kg kg<sup>-1</sup> that can be used for fluidized bed gasification [16].

In the steam-oxygen gasification, the gasifier is operated at temperatures of about 850  $^{\circ}$ C which enables the thermochemical decomposition of fuel into permanent gases, tars, char and ash. The so produced char and tars are then gasified or reformed by reaction with oxygen and steam. The raw syngas is comprised of H<sub>2</sub>O, H<sub>2</sub>, CO, CO<sub>2</sub>, CH<sub>4</sub>, light hydrocarbons (mostly C<sub>2</sub>H<sub>4</sub>), and tars (larger hydrocarbons, e.g., aromatics and polyaromatics).

Steam-oxygen gasification of wood and biogenic residues has been studied by other researches [15,17–24], however there is not as much data available as on other gasification processes. Results from these literature references are summarized in previous work of this author [14], but also in the results section of this paper these references are used for a comparison of results where applicable. Also, semi-commercial demonstration plants for the steam-oxygen fluidized bed gasification of biomass have already been in operation with a thermal input of 18 MW [25] and 100 MW [26,27]. The technical results of these endeavors were encouraging, underlining the potential of the technology. However, for sewage sludge, little fuel data were available. This is why this paper is needed, and constitutes one core goal of this paper.

The product gas contains beside the desired syngas species, tars and other impurities. Especially for sewage sludge, a high H<sub>2</sub>S and NH<sub>3</sub> concentration is observed in the syngas which is not tolerated by downstream synthesis catalysts [28]. Therefore, gas cleaning is required before synthesis. Reducing the costs of gas cleaning is important for the economic feasibility of the process [29]. To design and develop the entire process chain consisting of gasification, gas cleaning, and synthesis, detailed information on the product gas quality and composition, including impurities such as tars, sulfur-species, and ammonia, is needed.

The reduction of tars and impurities by primary measures such as active bed materials is considered economically and technically favorable compared to secondary measures [30,31]. A CaO-containing bed material, e.g., achieved by addition of limestone (CaCO<sub>3</sub>) or dolomite is catalytically active for cracking and steam reforming of tars [28,32,33] and acts as sorbent to capture  $H_2S$  (CaO +  $H_2S \leftrightarrow CaS + H_2O$ ) and COS (CaO + COS  $\leftrightarrow$  CaS + CO<sub>2</sub>) [31,34,35]. This work delivers experimental data on the application of this sulfur capture for sewage sludge gasification and its temperature dependency.

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This study continues the author's previous research on steam-oxygen gasification wherein sewage sludge gasification was compared to straw and wood gasification [14] and the influence of limestone as bed additive for sewage sludge gasification was investigated [36]. In this paper, a deeper look is cast on the effect of operation conditions on the gasification performance. These insights are important to validate process models and to support process design.

#### 2. Materials and Methods

#### 2.1. Fuel and Bed Material

The sewage sludge used in this work originated mainly from the municipal waste water treatment plant Häldenmühle in Marbach am Neckar in south-west Germany. A minor fraction of the sludge (less than 0.2 kg kg<sup>-1</sup>) came from the municipal wastewater treatment plants "Beilstein" and "Oberes Bottwartal" that are also located in south-west Germany. Those treatment plants delivered the sludge to the drying facility of Bioenergie Bottwartal GmbH&Co.KG, where the sludge was dried thermally with hot flue gases from a biogas CHP. After drying, the fuel had particle sizes of 5–10 mm, which might already be suitable for application in commercial fluidized beds. To suit the used experimental facility, the dried sewage sludge was crushed with a beater mill using a 2 mm sieve. The sewage sludge, gasified in the experiments, had high N, S, Cl, and ash contents. The ash summed up to almost half of the fuels dry mass. Therefore, the ash itself was used as bed material.

The proximate and elemental analysis is given in Table 1. The composition of this sewage sludge lies within the ranges reported by [37], wherein information on organic and non-organic components of typical sewage sludge is also given.

**Table 1.** Sewage sludge proximate and elemental analysis; au: as used (raw), daf: dry ash free, wf: water free, fc: fixed carbon, vm: volatile matter.

Proximate Analysis in kg $kg^{-1}$				Elemental Analysis in kg ${ m kg^{-1}}$					
У <b>Н2О</b>	Yash	Уfс	Уvm	γс	γн	УO	γn	Υs	УСІ
au	wf				d	af			
0.065	0.476	0.082	0.918	0.510	0.069	0.320	0.075	0.024	0.002

For some experiments, limestone was used as bed additive without pre-calcination. The calcination took then place in-situ in the gasifier releasing CO<sub>2</sub>. The limestone type "Messinghausener Sand 0.3–0.7" was obtained from LHOIST Germany/Rheinkalk GmbH with origin Messinghausen in Germany and consisted of the sieve fraction 0.3 mm–0.7 mm.

Table 2 shows the mineral composition of sewage sludge ash and limestone. The major elements are Si, Ca, P, Al, and Fe. Heavy metal concentrations in the dry sewage sludge are presented in Table 9.

Table 2. Main elements of sewage sludge ash and limestone.

			•	Element	al Analy	sis in kg	$ m kg^{-1}$			
	¥A12O3	УСаО	¥Fe2O3	Ук20	УMgO	VNa2O	¥P2O5	YSO3	\SiO2	YCO2
Sewage Sludge Ash Limestone	0.129 0.001	0.223 0.526	0.103 0.001	0.017	0.028 0.009	0.004 0.001	0.162	0.045	0.292 0.057	0.405

The particle sizes of the used solids are shown in Table 3. The dried sewage sludge had a wide range in particle sizes. During the gasification process, the sewage sludge particles seemed to release their volatile components with only a slight decrease in particle size, as the comparison of the particle sizes of sewage sludge and its ash showed. The limestone reduced its particle size during its residence time in the gasifier due to calcination and abrasion and was finer than the sewage sludge ash. Comprehensive particle size distributions on these materials are given in [14].

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	d <sub>p,10</sub>	d <sub>p,50</sub>	d <sub>p,90</sub>
_		in μm	
Sewage Sludge	200	1010	2100
Sewage Sludge Bed Ash	170	520	1700
Limestone	300	650	1200
Calcined Limestone	165	370	640

**Table 3.** Particle sizes of fuel and bed materials.

## 2.2. Experimental Facility

A 20 kW fuel input fluidized bed facility was used in this work. It is shown schematically in Figure 1. The bubbling fluidized bed reactor is 3.5 m high and has an internal diameter of 0.15 m in the fluidized bed region and 0.20 m in the freeboard above. The facility is equipped with several thermocouples and pressure transducers. Electrical heating allowed to control and adjust the temperature inside the gasifier and to compensate unwanted heat losses that are inherent to a research scale facility. Preheated steam was injected through six bubble cap nozzles to fluidize the bed. Oxygen could due to technical reasons not be mixed with the steam like in an industrial plant but was introduced through two additional bubble cap nozzles. For safety reasons  $N_2$  was added to the oxygen stream to reduce the  $O_2$  concentration to 0.75 m<sup>3</sup> m<sup>-3</sup>. The mass flow of  $N_2$  and  $O_2$  in these gas feeds was controlled by automatic mass flow controllers.

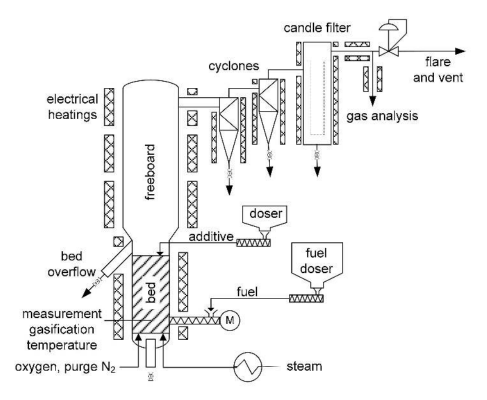


Figure 1. 20 kW fluidized bed gasification facility.

Dried sewage sludge was dosed gravimetrically with a screw feeder into the lower part of the fluidized bed, for which a small amount of nitrogen purge gas was also required. As a result of the high ash content in sewage sludge, the bed material was continuously removed during the gasification process by an overflow port to maintain a constant bed height.

For some experiments limestone was used as additive, which was dosed gravimetrically by a second doser through a port above the fluidized bed. A double ball-valve lock was used for pressure sealing.

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After the gasifier, the product gas is cleaned from particles by two cyclones and a candle filter, which are heated to  $400\,^{\circ}\text{C}$  to prevent tar condensation. The gas passed a pressure control valve before it was combusted in a flare.

## 2.3. Analysis Methods

Three slipstreams of the product gas were extracted for gas measurements after the candle filter of the facility.  $H_2$ , CO,  $CO_2$ , and  $CH_4$  were measured continuously in a combined NDIR and thermal conductivity gas analyzer (ABB AO2020), whereas hydrocarbons from  $C_2$  to  $C_4$  ( $C_xH_y$ ),  $H_2S$  and COS were semi-continuously (every 3 min) analyzed with a Varian CP-4900 Micro-GC. The slipstream for these devices was lead from the facility through a heated filter and a heated hose (both 180 °C) directly into 4 chilled impinger bottles, wherein the gas passed through an equal mixture of isopropanol and a 0.30 kg kg $^{-1}$  sulfuric acid aquatic solution for tar removal. Afterwards the gas was lead through an impinger bottle with a 0.15 kg kg $^{-1}$  sulfuric acid aquatic solution at ambient temperature to capture evaporated isopropanol. By these means, tars were removed to prevent damage to the analytical equipment, but due to the low pH value of the solution,  $H_2S$  and COS remained in the gas.

The continuous online gas measurement was also used to measure the dry volume flow of the syngas by a dilution method. After the candle filter, the syngas was diluted by a known  $N_2$ -stream controlled by a mass flow controller. The permanent gas concentrations were measured before and after the dilution. The syngas volume flow was calculated from the concentration changes.

The second sample line was used for wet chemical  $NH_3$  measurement according to a modified VDI 3838 [38] guideline: An impinger bottle was used to absorb the gaseous ammonia in a 1 mol  $L^{-1}$   $H_2SO_4$ -solution. For tar and humidity removal an isopropanol impinger bottle was arranged before the absorption solution. To avoid  $NH_3$  loss in the isopropanol bottle, the pH value was increased by adding NaOH. The samples were analyzed with UV-Vis spectroscopy.

The third sample line was used for tar measurement, where a heated filter and heated hose at  $350\,^{\circ}\text{C}$  were used. Tars were measured by extractive sampling and analysis according to the tar protocol CEN/TS 15439 [39]. For this, a quantified volume of the product gas was directed through cooled isopropanol washing bottles, wherein the tars were solved and condensed. The collected isopropanol-tar samples were analyzed by two methods as follows.

- (i) In the gravimetric analysis the isopropanol is vaporized out of the sample to gain a solvent free tar residue from whose mass the gravimetric tar concentration is derived. According to [40] this sample covers all "heavy" tar species with a molar mass over approximately 180 g mol<sup>-1</sup> (phenanthrene). Also tars with lower molar mass can be detected with this method, but with decreasing molar mass the detectability of tar species by gravimetric analysis decreases. Very "light" species, such as benzene, toluene, and xylene (BTX), are not detected or only to a small extend by gravimetric analysis. The gravimetric tar concentration is particularly important to assess the amount of very heavy tars with high boiling temperatures, that may condense easily on cold facility parts (e.g., gas coolers, valves) and can cause blockage and operation failure. The gravimetric analysis is performed three times for each tar sample and a mean value was calculated. As a novel method, elemental analysis of C, H, N, S, and Cl, as well as the heating value of the gravimetric tar was conducted for some samples to assess its composition.
- (ii) With gas chromatography (GC) analysis of the isopropanol-tar sample, actual tar components can be individually intentified and quantified. Especially "light" species can be detected very well. These species are partially still present after gas cooling, condensation and possibly after gas washing and thus have to be considered for downstream equipment. On the contrary, heavy tar species can only be detected up to a certain molar mass with a GC, in this work this was pyrene with 202 g mol<sup>-1</sup>. Quantification of all evaluated tar species was conducted with GC-FID by the institute of energy and process engineering of

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FAU University of Erlangen-Nuremberg with the following equipment and specifications: Agilent GC 7890A, column CP Sil 8 CB (L = 25 m, d = 0.25 mm), temperature programme: 3 min at 40 °C, heating rate 4.7 K min<sup>-1</sup> to 300 °C [41]. For simplification, benzene is also considered a tar component despite the fact that it is not named as tar in most of the literature.

Elemental and heating value analysis of fuel and gravimetric tar residues were conducted by a CHN-analyzer and combustion in a bomb calorimeter with wet sampling and ion-chromatography of S and Cl. Inorganic elemental analysis of sewage sludge ash and limestone was done by using acid digestion with subsequent ICP analysis.

Particle size distributions were measured by sieve analysis and with a laser diffraction particle size analyser.

## 2.4. Experimental Parameters

The main operation parameters for the steam-oxygen gasification process are gasification temperature, steam to carbon ratio, oxygen ratio, weight hourly space velocity and the bed material type.

For the choice of the gasification temperature, the following trade-off needs to be considered: Higher temperatures lead to faster gasification reactions and thus improved char and tar conversion but, on the other hand, can result in bed agglomeration when fuels with low temperature ash melting behaviour like straw are used. Also, higher gasification temperatures require more oxygen and thus reduce the overall conversion efficiency. The used bed material, sewage sludge ash, has a rather high ash melting temperature of 1200 °C but it already begins sintering at about 900 °C [42]. Therefore, the maximum bed temperature should be kept below that. In this work gasification temperatures from 650 °C to 900 °C were investigated to generate insight in the temperature influence on the process. Because of the advantages of higher gasification rates and lower tar yields, it is expected that an industrial sewage sludge gasifier will be operated at a temperature above 800 °C but below 900 °C, so as to prevent sintering and agglomeration of the bed.

The steam to carbon ratio,  $n_{SC}$ , describes the ratio of the total mole flow of water, including the fuel's moisture, to the mole flow of biomass carbon introduced into the gasifier:

$$n_{SC} = \frac{\dot{N}_{\text{H2O,steam}} + \dot{N}_{\text{H2O, moist}}}{\dot{N}_{\text{C,fuel}}}$$
(1)

Generally, higher  $n_{SC}$  results in a reduced energetic efficiency as additional steam needs to be introduced to the gasifier. On the other hand, low S/C ratios can reduce char conversion and cause higher tar yields. The water-gas-shift reaction (CO +  $H_2O \leftrightarrow H_2 + CO_2$ ) is severely influenced by the steam to carbon ratio, since  $H_2O$  pushes the equilibrium to the right side. Therefore, the  $H_2$  yield as well as the  $H_2/CO$ -ratio, that is important for downstream synthesis processes, can be increased with increasing S/C ratio.

The oxygen ratio  $n_{O2}$  (analogue to air ratio  $\lambda$  in combustion processes), often also referred to as equivalence ratio ER, expresses the ratio of the amount of oxygen actually used in the process to the amount of oxygen which would be required for the stoichiometric oxidation of the fuel:

$$n_{O_2} = \frac{\dot{N}_{O_2}}{\dot{N}_{O_2,\text{stoic}}} \tag{2}$$

In a technical process,  $n_{\rm O2}$  cannot be chosen independently of the gasifier's temperature, but is a function thereof. However, due to the electrical heating of the experimental facility used in this work,  $n_{\rm O2}$  is not coupled with the gasifier temperature and hence needs to be set by the operator. Nevertheless, in order to simulate industrially relevant conditions within the experiments,  $n_{\rm O2}$  was chosen according to adiabatic conditions. These calculations resulted in ranges of n from 0.2 to 0.3. To generate comparable conditions, the base case of  $n_{\rm O2} = 0.25$  was chosen.

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T weight hourly space velocity,  $n_{WHSV}$ , is the ratio of the fuel feeding rate to bed inventory mass:

$$n_{\text{WHSV}} = \frac{\dot{M}_{fuel,daf}}{M_{hed}} \tag{3}$$

Therefore  $n_{\rm WHSV}$  is a design value that brings the fuel load and the size of the fluidized bed reactor vessel in correlation. The  $n_{\rm WHSV}$  is correlated reciprocally to the fuel residence or space time in the bed. That means, for higher  $n_{\rm WHSV}$  the fuel has a shorter residence time in the bed.

The limestone additive ratio  $n_{\text{CaCO3}}$  expresses the mass ration of limestone additive to sewage sludge feed.

$$n_{CaCO_3} = \frac{\dot{M}_{CaCO_3}}{\dot{M}_{fuel,ar}} \tag{4}$$

The superficial velocity  $u_0$  in the fluidized bed was calculated by dividing the volume flow of steam and oxygen at operation conditions by the cross sectional area  $A_{\rm gasifier}$  of the gasifier (d<sub>gasifier</sub> = 0.15 m):

$$n_{CaCO_3} = \frac{\dot{M}_{CaCO_3}}{\dot{M}_{fuel,ar}} \tag{5}$$

The minimum fluidizing velocity  $u_{\rm mf}$  was derived from the particle Reynolds number at minimum fluidizing conditions  $Re_{\rm p,mf}$  which was calculated according to Kunii and Levenspiel and Wen and Yu.

$$u_0 = \frac{\dot{V}_{\text{H}_2\text{O,steam}} + \dot{V}_{\text{O}_2}}{A_{\text{gasifier}}} \tag{6}$$

The experimental parameters are summarized in Table 4.

## 2.5. Experimental Procedure

Prior to the experiments, the gasifier was heated to the desired bed temperature with around 7 kg of bed material, consisting of sewage sludge ash. The flows of sewage sludge, steam and oxygen were adjusted according to the desired experimental parameters. When the temperature and gas concentrations were steady, the experimental point commenced. Each experimental point was held in steady state conditions for at least 30 min and mean values for the continuously recorded data were calculated over this period. The tar and  $NH_3$  measurements, with a duration of 20 min each, were conducted during steady state operation conditions.

For the experiments with limestone as bed additive, a respective batch of limestone was added prior to the experiment to set up the desired fraction of CaO in the bed. After that, a continuous dosing of limestone with the additive ratio of  $0.25 \text{ kg kg}^{-1}$  (mass limestone per sewage sludge as used) was started.

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Variation	Run	ð	$n_{SC}$	$n_{\mathrm{O2}}$	$n_{ m WHSV}$	$\dot{m}_{ m fuel}$	$m_{ m bed}$	$n_{\text{CaCO3}}$	$u_0$	$u_0 u_{\rm mf}^{-1}$
		°C	$ m mol\ mol^{-1}$	$ m mol\ mol^{-1}$	$\mathrm{h}^{-1}$	${ m kg}{ m h}^{-1}$	kg	${ m kg}~{ m kg}^{-1}$	${ m m}~{ m s}^{-1}$	-
	O1	848	1.03	0.20	0.49	7.2	6.1	0	0.34	4.5
n <sub>O2</sub> _	O2/ref	842	1.05	0.25	0.44	7.1	7.8	0	0.36	4.7
	O3	842	1.01	0.27	0.41	7.2	7.6	0	0.36	4.8
-	O4	852	1.04	0.28	0.49	7.2	7.2	0	0.37	5.0
	S1	852	0.59	0.24	0.52	11.3	7.4	0	0.37	5.0
-	S2/ref	842	1.05	0.25	0.44	7.1	7.8	0	0.36	4.7
$n_{\rm SC}$ -	S3	839	1.48	0.25	0.43	7.2	7.3	0	0.47	6.3
-	S4	849	1.98	0.26	0.54	7.0	6.8	0	0.60	8.0
	θ1	659	1.03	0.26	0.54	7.0	6.8	0	0.29	3.9
θ -	θ2	778	0.97	0.26	0.53	7.1	7.1	0	0.32	4.2
0 -	ϑ3/ref	842	1.05	0.25	0.47	7.1	7.8	0	0.36	4.7
-	θ4	894	1.03	0.25	0.50	7.0	7.3	0	0.37	4.9
	θCa1	655	1.01	0.25	0.48	6.9	5.8	0.25	0.28	3.8
-	θCa2	708	1.04	0.26	0.26	6.9	7.6	0.25	0.31	4.1
$\theta$ with $\overline{}$ CaCO <sub>3</sub>	∂Са3	770	1.06	0.26	0.29	6.8	7.4	0.25	0.33	4.3
additive	θCa4 θCa5	798 844	1.12 1.08	0.28 0.27	0.40 0.38	6.4 6.6	7.9 8.4	0.25 0.25	0.33 0.35	4.5 4.6
-	θСа6	897	1.26	0.26	0.48	7.0	7.1	0.25	0.43	5.7
	W1	848	1.21	0.25	0.27	7.2	12.9	0	0.41	5.4
-	W2/ref	842	1.05	0.25	0.44	7.1	7.8	0	0.36	4.7
$n_{ m WHSV}$	W3	829	1.02	0.26	0.71	13.6	9.4	0	0.67	9.0
-	W4	849	0.99	0.26	1.19	7.0	2.9	0	0.34	4.5
-	W5	846	0.95	0.26	1.29	13.9	5.3	0	0.65	8.7

Table 4. List of all experimental of this work with respective operation conditions.

# 2.6. Thermochemical Equilibrium Calculations for Sulfur Capture with Calcium

The thermochemical equilibrium can be used to calculate the theoretical optimum of the  $H_2S$  and COS capture by reaction with CaO during gasification [34,43]. CaO is present in sewage sludge ash but can also be added in form of limestone (CaCO<sub>3</sub>) bed additive. The  $H_2S$  and COS capture follows the equilibrium reaction equations [31,34,44,45]:

$$CaCO_3 \leftrightarrow CaO + CO_2$$
 (7)

$$H_2S + CaO \leftrightarrow CaS + H_2O$$
 (8)

$$COS + CaO \leftrightarrow CaS + CO_2 \tag{9}$$

Since, in the gasifier, a reducing atmosphere is present, no  $CaSO_4$  can be formed, but instead, CaS.

With the data from Tables 1 and 2, the molar ratio of calcium and sulfur fed into the gasifier by the sewage sludge can be calculated:

$$n_{\text{Ca,S}} = \frac{\dot{N}_{\text{Ca, fuel}}}{\dot{N}_{\text{S, fuel}}} \tag{10}$$

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It can be seen that the sewage sludge has enough CaO present in its ash to capture all of the sludge's sulfur ( $n_{Ca,S} = 3.5 \text{ mol mol}^{-1}$ ). However, sulfur capture can only occur until the equilibrium concentration of  $H_2S$  and COS is reached in the syngas. Since also  $H_2O$  and  $CO_2$  are part of the relevant chemical equations (Equations (4)–(6)), the total syngas atmosphere influences the equilibrium concentrations of  $H_2S$  and COS. Hence, equilibrium calculations of the whole gasification process needed to be performed.

For this, the software FactSage 7.3, described in [46], with the database FactPS (pure substances) including all species relevant for this gasification process was used. FactSage's numerical solver then calculated the product composition where the Gibbs energy is at its minimum—the thermodynamic equilibrium:

$$\Delta G_{\rm R} = 0 \tag{11}$$

The input masses necessary for the gasification of 1 kg sewage sludge as used together with the gasification agents according to Tables 5 and 6 were entered into the FactSage solver. The gasification temperature was set. All calculations were done for ambient pressure of 1 bar. The database FactPS was used. Then, the equilibrium product composition, including the permanent gases like  $H_2$ , CO,  $CO_2$ , as well as  $H_2S$  and COS, was calculated. From this result, the equilibrium concentrations of  $H_2S$  and COS were calculated as volume fraction based on the dry syngas.

**Table 5.** Fuel input masses for FactSage equilibrium calculation for gasification of 1 kg as used sewage sludge, ash components other than CaO were neglected.

m <sub>H2O,moist</sub>	$m_{C}$	$m_{ m H}$	$m_{\rm O}$	$m_{ m N}$	$m_{\mathrm{S}}$	$m_{\text{Cl}}$	$m_{\text{CaO}}$
			in	kg			
0.065	0.250	0.034	0.157	0.037	0.012	0.001	0.098

**Table 6.** Gasification agent input masses and operation parameters for FactSage<sup>®</sup> equilibrium calculation for gasification of 1 kg as used sewage sludge.

Variation		Operation	Input Gasification Agent			
	p	θ	$n_{\mathrm{O2}}$	$n_{\rm SC}$	$m_{\mathrm{O2}}$	$m_{ m H2O}$
	in bar	in °C	in mol mol <sup>-1</sup>	in mol mol <sup>–1</sup>	in kg	in kg
$ref/n_{WHSV}$	1	850	0.25	1	0.212	0.309
θ	1	550 1000	0.25	1	0.212	0.309
$n_{O2}$	1	850	01	1	01	0.309
$n_{SC}$	1	850	0.25	$0 \dots 4$	0.212	0 1.5

## 3. Results and Discussion

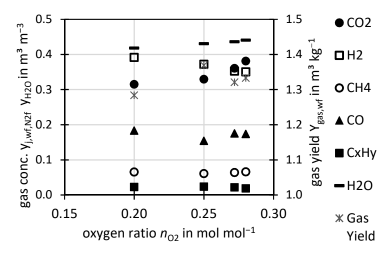
In this paper experimental runs investigating the influence of key operation parameters on sewage sludge gasification are presented. The actual operation values for these runs are summarized Table 4.

#### 3.1. Permanent Gases

In this chapter the influence of the operation parameters on the permanent gas composition and yields are shown. The concentration of the permanent gases  $H_2$ , CO,  $CO_2$ ,  $CH_4$  and  $C_xH_y$  ( $C_2$  to  $C_4$  hydrocarbons) are given as volume fractions on a dry  $N_2$ -free basis and the permanent gas yield as dry  $N_2$ -free volume in STP conditions (273.17 K, 101,300 Pa) per mass of dry ash free sewage sludge. In addition, the water concentration is shown as volume fraction on wet  $N_2$ -free basis.

## 3.1.1. Variation of Oxygen Ratio $n_{O2}$

Figure 2 shows the influence of the oxygen ratio  $n_{O2}$  on the permanent gases at constant temperature  $\vartheta$ . With increasing oxygen ratio, the  $H_2$  concentration decreased and the  $CO_2$  and  $H_2O$  concentration increased. Also, a slight decrease of CO was observed. This reflects the increased combustion of  $H_2$  and CO to  $CO_2$  due to the higher oxygen supply. The  $CCO_4$  and  $CCO_4$  concentrations were not influenced by the oxygen ratio in the investigated range. A rather constant value was observed for the gas yield.

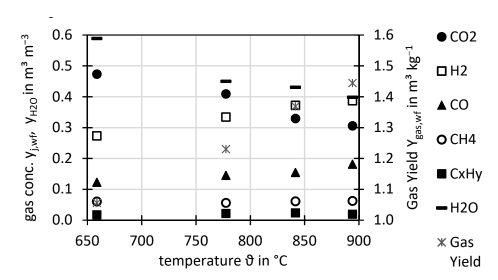


**Figure 2.** Permanent gas concentrations in dry N<sub>2</sub>-free basis, for H<sub>2</sub>O wet N<sub>2</sub>-free basis and dry N<sub>2</sub>-free gas yield per dry ash free fuel at different oxygen ratios  $n_{O2}$  and constant  $\vartheta$ .

In most experiments found in the literature, other parameters such as temperature also varied when different  $n_{\rm O2}$  were investigated. Therefore, it is difficult to compare the trends [22,47–49]. For coal gasification in a steam-oxygen blown slightly pressurized spouting bed at 940 °C [50],  $n_{\rm O2}$  variation also showed increasing CO<sub>2</sub> concentration whereas H<sub>2</sub> and CO were increasing until  $n_{\rm O2}$  = 0.35 but then decreased with increasing  $n_{\rm O2}$ . The reference connects this to increased gas yield and carbon conversion up to  $n_{\rm O2}$  = 0.35 followed by constant values for higher  $n_{\rm O2}$ . The trend of higher  $n_{\rm O2}$  for higher  $n_{\rm O2}$  in that reference is fitting well to this work. The variation of gas yield and carbon conversion with  $n_{\rm O2}$  is significantly dependent on the fuel reactivity and temperature, which could be the explanation for the differences between both studies.

# 3.1.2. Variation of Temperature $\vartheta$

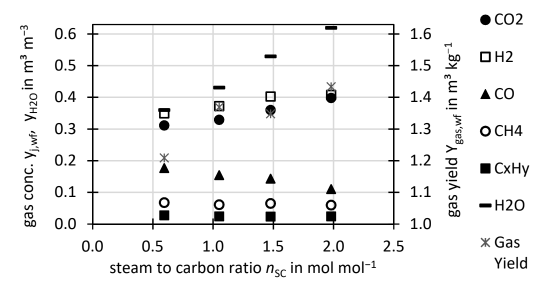
Figure 3 shows the influence of the gasification temperature  $\vartheta$  on the permanent gases for a constant  $n_{O2}$ . With increasing temperature, the gas yield increased due to higher carbon conversion to permanent gases. Also the  $H_2$  and CO concentration increased and the  $H_2O$  concentration decreased with increasing temperature, which is also related to an increased fuel conversion by steam gasification. No influence was seen on CH<sub>4</sub> and  $C_xH_y$  concentrations. Similar trends were found for steam-air gasification of corn straw [51] and sewage sludge [32] also, with regard to the  $N_2$ -free basis, and similar gas concentrations of all permanent gases were reported. Also, the temperature influence in steam-oxygen gasification of wood is reported in literature [22] and shows similar trends compared to this study concerning  $H_2$ ,  $CO_2$ , and gas yield. However, differences were reported in the trend of the CO, CH<sub>4</sub>, and  $C_xH_y$  concentrations, which decreased slightly with temperature in the reference whereas in this study they slightly increased (CO) or stayed constant (CH<sub>4</sub>,  $C_xH_y$ ). Similar trends to this study were also obtained by steam gasification of sewage sludge in the same gasifier operated in DFB mode [52].



**Figure 3.** Permanent gas concentrations in dry  $N_2$ -free basis, for  $H_2O$  wet  $N_2$ -free basis and dry  $N_2$ -free gas yield per dry ash free fuel for different temperatures  $\vartheta$  at constant  $n_{O2}$ .

## 3.1.3. Variation of Steam to Carbon Ratio $n_{\rm sc}$

Figure 4 shows the influence of the steam to carbon ratio  $n_{SC}$  on the permanent gases. The  $n_{SC}$  had a strong influence on the permanent gas composition, since  $H_2$  and  $CO_2$  strongly increased and CO decreased with  $n_{SC}$ . This was due to the water gas shift reaction ( $CO + H_2O \rightarrow H_2 + CO_2$ ) that was driven to the right side by adding steam. Therefore also the dry gas yield rose, since water was converted to  $H_2$ . The  $CH_4$  and  $C_xH_y$  concentrations were mostly unaffected. Naturally, as more steam was introduced, the  $H_2O$  concentration increased with  $n_{SC}$ . Parameter variation in steam-oxygen gasification of wood in a comparable facility was conducted by [22] and in air-steam gasification of corn straw by [51], where similar trends for  $H_2$ , CO,  $CO_2$  and  $H_2O$  were found, but other than in this work, a slight decrease of  $CH_4$  and  $C_xH_y$  was reported with increasing  $n_{SC}$  in the reference. High CO and lower CO concentrations at low CO as in this work were also reported by [21]. For steam-air gasification of refuse-derived fuel in a rotary kiln also a rise in gas yield with CO was reported [53].

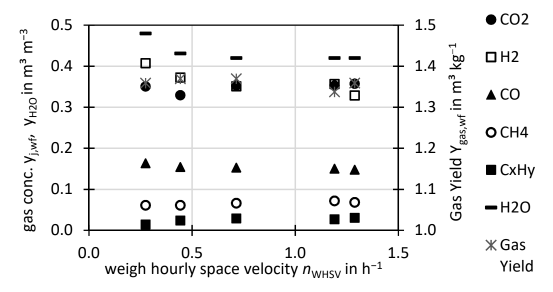


**Figure 4.** Permanent gas concentrations for different  $n_{SC}$  in dry  $N_2$ -free basis, for  $H_2O$  wet  $N_2$ -free basis and dry  $N_2$ -free gas yield per dry ash free fuel.

These results indicate, that through a higher  $n_{SC}$  a conversion of CO to  $H_2$  and  $CO_2$  can be achieved to improve the  $H_2$ -yield or to adjust the  $H_2$ /CO-ratio. However, higher  $n_{SC}$  requires more energy-intensive steam production, so that in practice it always needs to be considered if a higher  $n_{SC}$  or a downstream catalytic WGS unit, which are currently developed for uncleaned syngas [54], is preferred.

# 3.1.4. Variation of Weight Hourly Space Velocity $n_{WHSV}$

Figure 5 shows the influence of the weight hourly space velocity  $n_{\rm WHSV}$ , given in mass flow of dry ash-free sewage sludge per bed inventory mass, on the permanent gases. It has to be mentioned, that  $n_{\rm WHSV}$  was changed, both, by variation of fuel throughput and bed inventory as can be seen in Table 4. For the gases CO, CO<sub>2</sub>, CH<sub>4</sub>, and C<sub>x</sub>H<sub>y</sub>, no clear trend was observed for different  $n_{\rm WHSV}$ . A slight decrease for the H<sub>2</sub> concentration with increased  $n_{\rm WHSV}$  was present, which could be related to slightly decreased water-gas-shift reaction due to the decreasing residence time of the gas. Similar behavior was reported for sewage sludge steam-air gasification by [15], where for a higher turnover rate, respectively higher  $n_{\rm WHSV}$ , lower H<sub>2</sub> concentrations, and no or only little change in the other permanent gas concentrations were found.



**Figure 5.** Permanent gas concentrations for different  $n_{WHSV}$  in dry N<sub>2</sub>-free basis, for H<sub>2</sub>O wet N<sub>2</sub>-free basis and dry N<sub>2</sub>-free gas yield per dry ash free fuel.

## 3.2. Tar Concentration

As described previously, tar sampling was conducted wet chemically and two sample analysis methods were used: GC analysis and gravimetric analysis. By GC analysis 18 tar species (named in [14]) were quantified. In the following, the GC tars are shown as ECN tar classes [40]. The majority of the GC-detectable tars were found to be benzene, toluene, xylene (BTX) included in the here used classes "benzene" and "ECN3". In previous work [36] it was found, that also unneglectable amounts of heterocyclic species, mainly pyridine, are present in tar from sewage sludge gasification. However, pyridine was not quantified in the GC analysis presented in this work since the used GC was not calibrated for that.

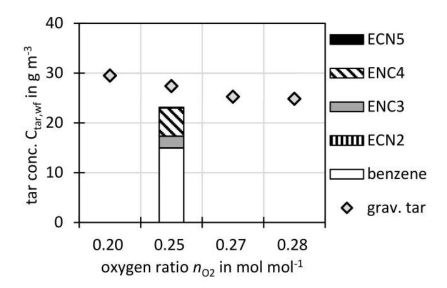
The gravimetric analysis detects mostly heavy tars, but also inorganic components e.g., salts from ammonia, sulfur and chlorine can be present (see previous work [14,36]). The carbon mass fraction in the gravimetric tars was found to be around  $0.5 \ kg \ kg^{-1}$ , which has to be considered when using the presented tar concentrations for calculating a carbon mass balance.

All concentrations are shown as tar mass per volume of dry and  $N_2$ -free syngas in STP conditions. The tar yields can be calculated by multiplying the tar concentration with the gas yields from Section 3.1.

In the following, the influence of different operation conditions on the tar concentrations are shown. In this study the influence of the operation parameters  $n_{O2}$ ,  $\vartheta$ ,  $n_{SC}$ ,  $n_{WHSV}$  is shown. The tar concentration is also significantly dependent on the catalytic activity of the bed material e.g. in case of limestone or dolomite addition which was not subject of this work but was investigated in literature [51].

# 3.2.1. Variation of Oxygen Ratio $n_{O2}$

Figure 6 shows the influence of the oxygen ratio  $n_{\rm O2}$  on the tar concentration. The gravimetric tar concentration is dropping slightly with increasing oxygen ratio. This could be related to enhanced tar oxidation. Since due to technical reasons only one GC analysis was performed for this parameter variation, no trend can be observed.



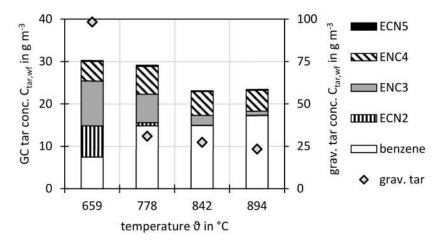
**Figure 6.** Gravimetric and GC tar concentrations dry, N<sub>2</sub>-free and STP basis at different oxygen ratios  $n_{O2}$  and constant  $\vartheta$ .

Again, although  $n_{\rm O2}$  was varied by several researchers [22,47–49], in all found references, other significant parameters such as temperature were altered as well. Therefore, no exclusive variation of  $n_{\rm O2}$  was found in the literature. This study can thus at least shed some light on the behaviour of gravimetric tar concentration at different  $n_{\rm O2}$ .

## 3.2.2. Variation of Temperature $\vartheta$

Figure 7 shows the tar concentration at different gasification temperatures  $\vartheta$ . At the 659 °C a very high gravimetric tar concentration of 98 g m<sup>-3</sup> was measured. When the temperature was increased to 778 °C the concentration drastically dropped to 31 g m<sup>-3</sup>. With further temperature increase, the gravimetric tar concentration steadily dropped to 27 g m<sup>-3</sup> and 21 g m<sup>-3</sup> for 842 °C and 894 °C, respectively. As known [55], at low gasification temperatures, considerable fractions of the fuel are not converted into small molecules, but form large tar molecules which are detected well by gravimetric analysis. At higher temperatures those gravimetric tars are cracked or reformed into smaller tar species or into permanent gases.

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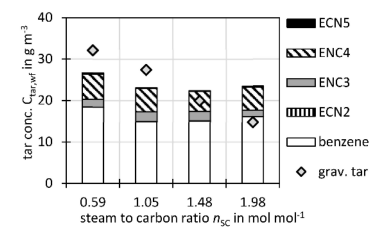
**Figure 7.** Gravimetric and GC tar concentrations on dry,  $N_2$ -free and STP basis at different gasifier temperatures.

The total GC tar concentration was 31 g m $^{-3}$  at 659 °C. When the temperature was increased, the GC tars were found to be decreasing slightly but steadily to 23 g m $^{-3}$  at 894 °C. It has to be noted that the concentration of tar species from ECN2 and ECN3 classes decreased severely with temperature, while benzene is increasing.

In the literature, overall similar behaviour is reported [22,47,52,56–59].

## 3.2.3. Variation of Steam to Carbon Ratio $n_{SC}$

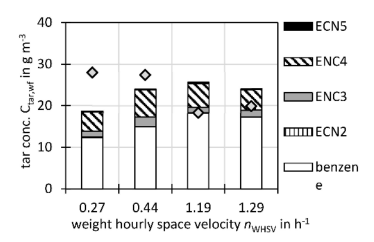
Figure 8 shows the tar concentrations against steam to carbon ratio  $n_{SC}$ , where a decrease of the gravimetric tar concentration with increasing  $n_{SC}$  was observed. This could be due to the known [55,60] effect of higher steam concentrations on reforming of gravimetric tars. The GC tars stayed rather constant over  $n_{SC}$ . This corresponds well to other gasification experiments where higher steam concentrations also reduced tar concentrations [22,60,61].



**Figure 8.** Gravimetric and GC tar concentrations on dry,  $N_2$ -free and STP basis at different steam to carbon ratios  $n_{SC}$ .

## 3.2.4. Variation of Weight Hourly Space Velocity $n_{WHSV}$

Figure 9 shows the tar concentration against the weight hourly space velocity  $n_{\rm WHSV}$ . For the gravimetric tars no clear trend was observed. It also has to be noted that the point at  $n_{\rm WHSV} = 1.3 \, \rm h^{-1}$  was conducted with a higher fuel mass flow of around 14 kg h<sup>-1</sup> compared to around 7 kg h<sup>-1</sup> (see Table 4) for the other points. Despite that, the tar concentration for the points at  $n_{\rm WHSV} = 1.2 \, \rm h^{-1}$  and  $1.3 \, \rm h^{-1}$  were found to be similar.



**Figure 9.** Gravimetric and GC-MS tar concentrations on dry,  $N_2$ -free and STP basis at different weight hourly space velocities.

When comparing the two gravimetric tar concentrations lower  $n_{\rm WHSV}$  with the two ones at higher  $n_{\rm WHSV}$ , a decrease can be seen. For the GC tars no trend is visible. However, a more comprehensive study is needed to fully assess the influence of  $n_{\rm WHSV}$  on tar concentration.

In literature [15] for steam-air gasification it was reported that tar concentration is slightly increased with higher "turnover rate" respective higher  $n_{\text{WHSV}}$ . For steam gasification, [56] found slightly increased gravimetric tar yields but constant GC tar yields for increasing  $n_{\text{WHSV}}$  which is consistent with this work.

## 3.2.5. Gravimetric Tar Composition and Heating Value

For some runs, the gravimetric tar sample was retrieved after the completed gravimetric analysis and analysed. Figure 10 shows the elemental composition of gravimetric tars. It can be seen that the carbon mass fraction is only around 0.5 kg kg $^{-1}$  compared to up to 0.87 kg kg $^{-1}$  for tar from wood [14]. Also, significant amounts of S, N, and Cl were found in sewage sludge gravimetric tar in this study. It is therefore assumed that, when applied for sewage sludge derived syngas, the gravimetric analysis does not only sample typical tar species (e.g., PAH), but also salts like NH<sub>4</sub>Cl, (NH<sub>4</sub>)<sub>2</sub>CO<sub>3</sub>, and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> that are formed from NH<sub>3</sub>, HCl, and H<sub>2</sub>S which are present in high concentrations in the syngas. This hypothesis is supported by the measured higher heating value of the gravimetric tars, which were found to be around 25 MJ/kg and are thus around 40% below the heating value of a typical gravimetric tar species (e.g., naphthalene 40 MJ kg $^{-1}$ ).

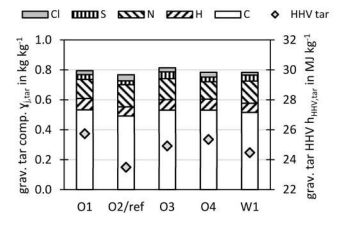


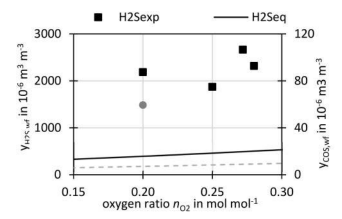
Figure 10. Elemental analysis and heating values of gravimetric tars.

## 3.3. H<sub>2</sub>S and COS Concentration

In this chapter the concentrations of the gaseous sulfur species H<sub>2</sub>S and COS are discussed.

## 3.3.1. Variation of Oxygen Ratio $n_{O2}$

Figure 11 shows the measured  $H_2S$  and COS concentrations as well as the equilibrium concentrations, calculated with FactSage<sup>®</sup> (see Section 2.6), over the oxygen ratio  $n_{O2}$ . All concentrations are presented in water free (wf) and  $N_2$ -free basis. For oxygen ratios from 0.20 to 0.28,  $H_2S$  concentrations between  $2000 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup> and  $3000 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup> were measured at the respective conditions of  $847 \pm 5$  °C. From the measurements of this work, it is hard to deduct a specific trend for  $H_2S$ , but it can be noted that the two points at higher  $n_{O2}$  show slightly higher concentrations. For COS only one measurement was performed of  $60 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup> at  $n_{O2} = 0.2$  mol mol<sup>-1</sup>.



**Figure 11.** Measured and equilibrium  $H_2S$  and COS concentrations on dry and  $N_2$ -free basis at different oxygen ratios  $n_{O2}$  and constant  $\vartheta$ .

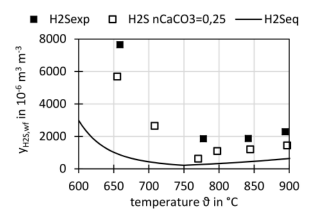
The equilibrium predicts a slight increase of  $H_2S$  and COS concentration with  $n_{O2}$ . This is related to higher feed of oxygen to the gasifier at higher  $n_{O2}$ , leading to the formation of  $H_2O$  and  $CO_2$ . This drives Equations (5) and (6) away from sulfur capture following Le Chatelier's principle.

## 3.3.2. Variation of Temperature $\vartheta$

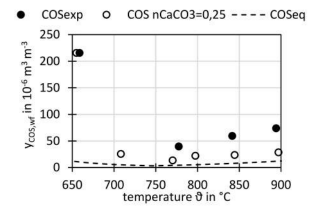
Figure 12 shows the  $H_2S$  concentration and Figure 13 the COS concentration at different temperatures for cases with the reference bed material, sewage sludge ash, but also for runs with limestone additive. Additionally, the  $H_2S$  and COS concentrations in the thermochemical equilibrium, calculated by the software FactSage<sup>®</sup> as described Section 2.6 are shown.

A strong dependence of the concentration of the two sulfur species on the temperature could be observed in the experiment as well as in the equilibrium calculations. At low temperatures high concentrations of up to  $7653 \times 10^{-6}$  m³ m⁻³ for H₂S and up to  $215 \times 10^{-6}$  m³ m⁻³ for COS were measured with sewage sludge ash as bed material. With increasing temperatures the H₂S and COS concentrations were decreasing until a minimum was reached before the concentration increased again at higher temperatures. This is related to the facts, that sewage sludge ash contains CaO which is can capture sulfur through reactions Equations (5) and (6) which are exothermic and therefore favored at lower temperatures. However the calcination reaction (Equation (4)) is favored at higher temperatures. The overlay of calcination and sulfur capture leads to the here seen temperature dependency.

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**Figure 12.** Measured and equilibrium  $H_2S$  concentration on dry and  $N_2$ -free basis at different temperatures  $\vartheta$ .



**Figure 13.** Measured and equilibrium COS concentration on dry and  $N_2$ -free basis at different temperatures  $\theta$ .

For the case with limestone additive, a similar trend was observed but consequently lower concentrations were measured. The equilibrium has a similar trend as the experiments, but shows for all cases lower concentrations, which is in agreement with the fact that the equilibrium predicts optimal sulfur capture and thus the lowest possible concentration. The lowest measured concentration without limestone additive was  $1855 \times 10^{-6} \text{ m}^3 \text{ m}^{-3}$  at  $778 \,^{\circ}\text{C}$  and with limestone additive the lowest concentration was  $624 \times 10^{-6} \text{ m}^3 \text{ m}^{-3}$  at  $770 \,^{\circ}\text{C}$ . With the limestone additive the H<sub>2</sub>S concentration could thus be reduced by around factor 3 at these temperatures. The absolute equilibrium minimum is was calculated for  $740 \,^{\circ}\text{C}$  with  $243 \times 10^{-6} \, \text{m}^3 \, \text{m}^{-3}$ .

For COS similar behavior was observed, at 770 °C–778 °C the concentration for sewage sludge ash bed is  $40 \times 10^{-6}$  m³ m<sup>-3</sup>, with limestone additive it is reduced to  $13 \times 10^{-6}$  m³ m<sup>-3</sup>, also by factor 3. The COS minimum equilibrium concentration was calculated for 740 °C with  $6 \times 10^{-6}$  m³ m<sup>-3</sup>.

As said, with further increasing temperatures the measured and equilibrium  $H_2S$  and COS concentrations are rising again. At the reference temperature of 840 °C, which is also a very common temperature to operate sewage sludge incinerators and gasifiers [62], slightly higher concentrations of  $H_2S$  with  $1873 \times 10^{-6}$  m³ m<sup>-3</sup> and COS of  $59 \times 10^{-6}$  m³ m<sup>-3</sup> were observed without additive. Again, the limestone additive brings a reduction. It was noticed that, with limestone additive, the COS concentrations can be brought very close to the equilibrium for all investigated temperatures above 750 °C, while for  $H_2S$ , the concentrations with additive still have some distance to the equilibrium.

In literature [34,43], similar trends of  $H_2S$  concentrations over temperature in the presence of CaO were also reported in experiments and equilibrium calculations.

## 3.3.3. Variation of Steam to Carbon Ratio $n_{SC}$

Figure 14 shows the  $H_2S$  and COS concentrations for different steam to carbon ratios  $n_{SC}$ . The measured  $H_2S$  concentrations increased with  $n_{SC}$ . The lowest S/C ratio tested was 0.6 mol mol<sup>-1</sup> which lowered the  $H_2S$  concentration to  $1340 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup>. A similar trend was found by [34]. This follows well the trend of the equilibrium where the  $H_2S$  concentrations are increasing with  $n_{SC}$  following Le Chatelier's principle: With higher  $n_{SC}$ , more steam is introduced into the gasifier, driving Equation (5) to the left towards less  $H_2S$  capture.

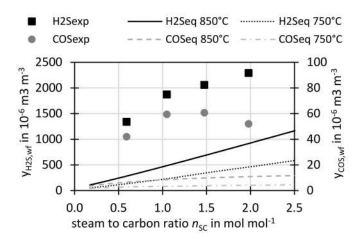


Figure 14. Measured and equilibrium  $H_2S$  and COS concentration on dry and  $N_2$ -free basis at different steam to carbon ratios, additionally equilibrium calculations for  $\vartheta = 750$  °C are showing the best possible sulfur capture.

The measured COS concentration stayed on the same level throughout the  $n_{\rm SC}$  variation, whereas the equilibrium predicts a very small increase. Since steam is not present in Equation (6),  $n_{\rm SC}$  has no direct effect on COS capture. However through the water gas shift reaction, steam addition also produces more CO<sub>2</sub> influencing the COS capture.

Additionally, the equilibrium for 750 °C is shown since sulfur capture is maximized at this temperature. It can be seen that, at operation conditions of 750 °C and low  $n_{SC}$ , very low concentrations are achievable according to the equilibrium.

In literature [34,43], similar trends of  $H_2S$  concentrations over  $n_{SC}$  in the presence of CaO were also reported in experiments and equilibrium calculations.

## 3.3.4. Variation of Weight Hourly Space Velocity $n_{WHSV}$

Figure 15 shows the  $H_2S$  and COS concentration at different fuel space velocities  $n_{WHSV}$ . For the measured values, no specific trend could be observed. Since the equilibrium already assumes infinite residence time of the fuel, it cannot produce trends of the space velocity and thus a constant equilibrium concentration is shown.

#### 3.4. NH<sub>3</sub> Concentration

Figure 16 shows the measured  $NH_3$  concentrations in the syngas with respect to different operation conditions. As indicated above,  $NH_3$  was measured wet chemically. Therefor these measurements were not performed for every experimental point. Due to the limited points, it is not always possible to deduct a clear trend, therefore further studies with more comprehensive  $NH_3$  analysis are needed.

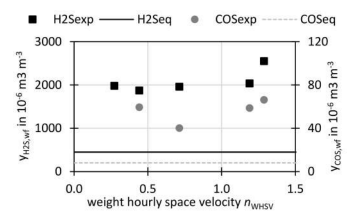


Figure 15. Measured and equilibrium H<sub>2</sub>S and COS concentration at different fuel space velocities.

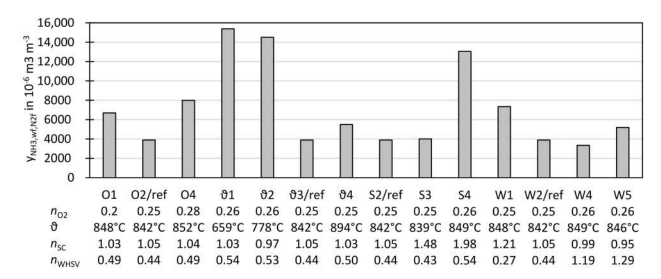


Figure 16. Compilation of all conducted NH<sub>3</sub> measurements.

For most runs, NH<sub>3</sub> concentrations of between  $4000 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup> and  $8000 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup> were measured. However, for runs with lower  $\vartheta$  or with higher  $n_{SC}$  higher NH<sub>3</sub> concentrations of up to  $15,000 \times 10^{-6}$  m<sup>3</sup> m<sup>-3</sup> were measured.

It can be concluded that NH<sub>3</sub> is present in considerable concentrations in sewage sludge derived syngas, which is clearly related to the high nitrogen content in the fuel. Other research also shows high NH<sub>3</sub> concentration for high N-containing fuels such as sewage sludge [14,52], manure [52], peat [26], or coal [63]. It has to be noted that steam-oxygen gasification seams to show considerably lower NH<sub>3</sub> concentration than steam gasification [52] in the same facility at IFK University of Stuttgart. Literature also reports decreasing NH<sub>3</sub> concentrations with increasing temperature, which is in line with this study [64].

## 3.5. Ash Yields and Carbon Contents

Depending on the process design, different kinds of ash streams are produced and are usually divided in bed ash and fly ash. Carbon contained in ashes leaves the reactor ungasified and therefore reduces the conversion efficiency of the process.

#### 3.5.1. Bed Ash

In the used research facility as well as in some commercial sewage sludge gasifiers [65], the bed ash is the main ash stream since granular dried sludge is used as fuel. The bed ash is therefore also serving as bed material. Since constantly new bed ash is produced, it is

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discharged continuously over an overflow. In this work, the bed ash had a particle size of  $d_{\rm p,50}$  = 520  $\mu$ m and was only slightly smaller than the used fuel (see [14] for detailed particle distribution).

The bed ash yield per dry fuel is around 0.41 kg/kg as calculated from the difference of the fuels ash content (proximate analysis Table 1) and the measured fly ash yields (Table 6). The carbon content of the bed material after discharge from the reactor was around 0.017 kg/kg at a reference gasification temperature of 850  $^{\circ}$ C however for lower temperature the carbon content was found to be higher up to 0.04 kg/kg at 668  $^{\circ}$ C (Table 7).

run	ð	$n_{\rm SC}$	$n_{O2}$	$n_{ m WHSV}$	$Y_{ash,fly}$	$Y_{ash,bed,bal}$	YC,ash,fly	YC,ash,bed
θ1	659	1.03	0.259	0.54	0.072	0.404	0.110	0.045
ϑ2	778	0.97	0.256	0.53	0.063	0.413	0.092	0.031
ref	842	1.05	0.250	0.47	0.062	0.414	0.085	0.017
$\vartheta 4$	894	1.03	0.254	0.51	0.059	0.417	0.047	0.006
<b>W1</b>	848	1.21	0.252	0.27	0.065	0.411	0.066	0.016
SC4	849	1.98	0.256	0.54	0.070	0.406	0.055	0.014

**Table 7.** Fly ash and bed ash yields and carbon content.

#### 3.5.2. Fly Ash

Smaller sized fuel particles, dust produced by attrition in the bed or soot formed by fuel conversion are entrained from the fluidized bed and are considered fly ash. As described in Section 2.2, this fly ash is in the used research facility captured by a primary cyclone, a secondary cyclone and a ceramic candle filter. The yields of the respective fly ash fractions for the reference run ( $n_{O2} = 0.25$ ,  $\vartheta = 842$ ,  $n_{SC} = 1.05$  mol mol<sup>-1</sup>) are  $Y_{ash,prim.cyclone} = 0.051$  kg kg<sup>-1</sup>,  $Y_{ash,sec.cyclone} = 0.008$  kg kg<sup>-1</sup> and  $Y_{ash,filter} = 0.003$  kg kg<sup>-1</sup>. It can be seen that the majority of the fly ash is separated by the primary cyclone.

In Table 7, the fly ash yield and the carbon mass fraction are shown for different runs, wherein all respective fly ash fractions (prim. cyclone, sec. cyclone, filter) have been cumulated. It can be seen that the fly ash yield per fuel mass is for all presented runs between 0.059 kg kg $^{-1}$  and 0.072 kg kg $^{-1}$ . The carbon mass fraction of the fly ash varied between 0.047 kg kg $^{-1}$  and 0.110 kg kg $^{-1}$  and was found to be dependent on gasification temperature.

## 3.6. Ash Main Elements and Heavy Metal Content

Table 8 shows the main elemental analysis of the ash fractions of the reference run. It can be seen that the elemental composition for all ash fractions was similar. The major components of the ash were calcium, silica, phosphorous, aluminium and iron. These elements originated from the mineral content of waste water and additives for phosphorous precipitation. Due to the high phosphorous content, the ash can be used as fertilizer if it fulfils the respective regulations or as raw material for industrial fertilizer production.

Table 9 shows the heavy metal concentration in dry sewage sludge and the ash fractions from the gasification experiments. Also the German legal limits for fertilizers [66] in general and specifically for the deployment of sewage sludge as fertilizer [67] are given. It can be seen that the used dry sewage sludge already fulfils the legal limits for all components except Thallium. Depending on the species, the concentration of the heavy metals in the ashes are different to those of the sewage sludge. For more volatile elements such as Cd, Pb and Tl, the concentration in the bed ash is reduced compared to the sewage sludge, but these elements are found again in the fly ashes with increased concentrations. It is gratifying, that the bed ash is therefore, compared to sewage sludge, depleted of Cd, Hg, Tl and has reduced Pb concentrations. The bed ash fulfils all legal limits except of Nickel, however the high Ni concentrations of the bed ash are assumed to come from the high temperature steel pipe of the reactor and are not attributed to the Ni present in the sewage sludge. Since in industrial plants are refractory lined, the industrial ash is expected to meet the Ni limits. The bed ash is only slightly below the Cu limit, though.

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Table 8. Ash main elemental analysis.

	Bed Ash		Fly Ash	
	Overflow	Prim. Cyclone	Sec. Cyclone	Filter
		mass fraction	n in kg kg <sup>-1</sup>	
$Al_2O_3$	0.105	0.105	0.107	0.108
BaO	0.001	0.001	0.001	0.001
CaO	0.235	0.252	0.302	0.335
$Fe_2O_3$	0.098	0.095	0.115	0.108
K <sub>2</sub> O	0.015	0.017	0.014	0.011
MgO	0.025	0.029	0.031	0.031
$MnO_2$	0.003	0.002	0.003	0.002
Na <sub>2</sub> O	0.004	0.004	0.004	0.003
$P_2O_5$	0.141	0.128	0.137	0.135
$SO_3$	0.040	0.033	0.033	0.028
$SiO_2$	0.307	0.267	0.228	0.228
SrO	0.001	0.001	0.001	0.001
TiO <sub>2</sub>	0.005	0.006	0.006	0.006

**Table 9.** Heavy metal concentration in sewage sludge and ash fractions, for comparison: German legal limit. <sup>a</sup> German fertilizer law (DüMV), <sup>b</sup> German sewage sludge treatment law (AbklärV2017), <sup>c</sup> for dry sewage sludge with  $8\% P_2O_5$ , <sup>d</sup> for ash with  $12\% P_2O_5$ , <sup>e</sup> stricter limits for Cr<sup>VI</sup>:  $2 \text{ mg kg}^{-1}$ .

	Sewage Sludge	Bed Ash	Prim. Cyclone	Sec. Cyclone	Filter	German Legal Limit
		mas	s fraction of t	race elements	in mg/kg	
As	10	38	40	86	38	40 a
Be	1	1	1	1	1	-
Cd	1.7	0.2	3.3	29	0	$4^{a,c}/6^{a,d}$
Co	3	29	29	41	29	-
Cr	377	828	1018	1939	828	900 <sup>a</sup> /2 <sup>e</sup>
Cu	515	865	946	1552	865	900 <sup>b</sup>
Hg	0.4	0.0	0.1	0.5	0.0	1 <sup>a</sup>
Mo	7	25	36	81	25	-
Ni	25	592	4639	6338	592	80 a
Pb	60	134	219	710	134	150 <sup>a</sup>
Sb	2	15	12	23	15	-
Se	13	43	103	217	43	-
Sn	134	75	122	195	75	-
Tl	3	< 0.3	< 0.3	< 0.3	< 0.3	1 <sup>a</sup>
V	31	52	53	64	52	-
Zn	1010	1607	1589	2291	1607	4000 <sup>b</sup>

It has to be noted, that in this study only the total value of chrome was measured, but not  $Cr^{VI}$  for which strict limits are imposed [66]. However, from a commercial air-blown sewage sludge fluidized bed gasifier the heavy metal concentrations of the bed ash were reported, and the  $Cr^{VI}$  concentration was below 0.5 mg kg $^{-1}$  and with that below the legal limit [62]. From this commercial facility also very low concentrations of Cd, Hg and Tl were reported, each below 0.2 mg kg $^{-1}$ , 0.05 mg kg $^{-1}$  and 0.5 mg kg $^{-1}$ , respectively, and Pb was reported with 20 mg kg $^{-1}$ . These values correspond well to the findings of this study. The Ni concentration was reported to be much lower in the commercial gasifier, underlining the hypothesis that, in this study, the Ni loading of the ash came from the high temperature steel of the reactor pipe. Arsenic, at 3 mg kg $^{-1}$ , was reported to be much lower in the commercial gasifier compared to this study, however there could also be differences in the sewage sludge composition, which is not considered in [62].

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## 3.7. Hydrodynamic Behavior of the Sewage Sludge Ash Bed Material

Overall, the fluidization behaviour of the bed material was good during the experiments. The facility was in operation with sewage for around 25 trial days with each a gasification operation of 4 h to 8 h. In most cases the experimental conditions were varied several times per day to investigate different experimental points. During the majority of the experiments, stable hydrodynamical conditions were reached quickly, visible by a constant pressure drop over the fluidized bed.

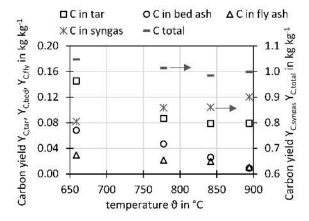
However, bed melting and agglomeration have been observed on few occasions for operation points with low superficial fluidization velocity of  $u_0 < 0.2 \text{ m s}^{-1}$  corresponding to  $u_0 u_{mf}^{-1} < 3$  and temperatures over 850 °C. In these cases at the beginning stable condition could be reached, but then a rapid temperature rise accompanied with partial bed material melting and agglomeration was triggered by a short interruption of the fuel supply. After that, fluidization was not possible anymore and the agglomerated bed material needed to be removed from the reactor. Such short fuel supply interruptions also occurred at other operation points with higher fluidization velocity, but no negative effect was present there. Leading to the explanation, that after the fuel supply interruption the endothermic effect of the fuel conversion ceased and the further supplied oxygen could exothermally oxidize char and reduced metals (e.g., FeO) in the bed. Due to the low fluidization velocity, the bed was poorly mixed and therefore the temperature could rise above the ash melting temperature locally at the gasifier bottom. The authors therefore advise to always ensure a good bed material mixing by maintaining sufficient fluidization velocity of  $u_0 u_{mf}^{-1} >> 3$ . However, it is likely that this problem was more prominent in the small electrically heated research reactor than in a bigger industrial facility.

#### 3.8. Carbon Balance

Figure 17 shows the carbon yields for different gasifier temperatures. On the left y-axis, the carbon yield in tars, in bed ash and in fly ash is shown. On the right y-axis, the carbon yield in the syngas main components (CO,  $CO_2$ ,  $CH_4$ ,  $C_xH_y$ ) as well as the total sum of all measured carbon yields is depicted.

The carbon yield in tars was calculated as sum of carbon in gravimetric tars as measured from gravimetric tar concentration (Figure 7) and its elemental analysis (Figure 10) as well as the carbon in benzene, toluene and xylene (ECN3 in Figure 7) taken from the GC analysis. This approach adds the light tars to the gravimetric tars to represent a total tar amount. The carbon yield in tars decreased with temperature, since the tars are thermally cracked at higher temperatures.

The carbon yield in bed ash and fly ash decreased with increasing temperature since more char was gasified, e.g., by the water-gas reaction at higher temperatures due to faster reaction rates.



**Figure 17.** Carbon balance: Yield of carbon in syngas, tar, bed ash, fly ash and total sum for different gasifier temperatures.

The carbon yield in the syngas main components increased with temperature, since at higher temperature more char and tar is converted to gases such as CO, CO<sub>2</sub>, and CH<sub>4</sub>.

The total carbon yield sum for all shown carbon points are close to 1 kg kg $^{-1}$ , which means that almost the same amount of carbon was found in the products than introduced with the sewage sludge.

## 4. Conclusions and Outlook

This work offers a comprehensive experimental study of the steam-oxygen gasification of sewage sludge for synthesis gas production in a 20 kW fluidized bed plant. The experimental results can be utilized for process design and to set-up and validate a gasifier model, since product gas, char, tar and impurity yields as well as their dependencies on the operation conditions are presented.

Since the gas yield was enhanced and the tar yield decreased with increasing gasification temperature it can be concluded, that the gasification temperature should be chosen as high as practically feasible (i.e.,  $850~^{\circ}$ C to  $900~^{\circ}$ C) taking into account the ash melting behavior.

The  $H_2/CO$ -ratio can be controlled by altering the steam to carbon ratio  $n_{SC}$  as steam promotes the water gas shift reaction. It could be shown that the  $H_2/CO$ -ratio could be tailored to 2 (e.g., for methanol or DME-production) or 3 (for methanation) with a reasonable range of  $n_{SC}$ .

In the investigated process, the majority of the ash is received as bed ash. Hazardous cadmium, mercury and thallium, contained in the sewage sludge, is evaporated in the gasification process, therefore the bed ash was found to be free of or very lean in those elements, which is beneficial for the ash application for fertilizer production. Since the sewage sludge ash contains calcium, the majority of the sludge's sulfur is bound in the bed ash. By adding limestone to the gasifier this sulfur capture is enhanced and the  $H_2S$  and COS concentrations in the syngas are lowered.

Overall, the steam-oxygen gasification proved to be an efficient and technically feasible process for sewage sludge treatment. The process can play a role in closing the loop for carbon by converting the organic fraction of sewage sludge to carbon-containing bio-fuels or chemicals as well as in closing the loop for phosphorous that can be recovered from the heavy metal lean ash. Therefore, in future, this process can be considered as an alternative to fluidized bed incineration.

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## Abbreviations

as used au daf dry ash free combined heat and power plant CHP DME dimethyl ether FR Equivalence ratio, synonymous to oxygen ratio  $n_{O2}$ fcfixed carbon **FID** flame ionizing detector GC gas chromatography **SNG** Substitute Natural Gas volatile matter vm wf water free

## **Symbols**

A	m²	cross sectional area
$Ar_{p}$	-	Archimedes number of particle
$C_{\text{tar}}$	${ m g}~{ m m}^{-3}$	tar mass per volume of dry and N2-free gas in STP conditions
$d_{p}$	μm	particle size
Υj	${ m kg~kg^{-1}}$	mass fraction of element j in solid sample
m	kg	mass
m	$ m kgh^{-1}$	mass flow
N	mol	substance amount
$\dot{N}$	$ m mol~h^{-1}$	mole flow
$n_{\text{Ca,S}}$	$\mathrm{mol}\mathrm{mol}^{-1}$	molar ratio of calcium feed to sulfur feed
$n_{\text{CaCO3}}$	${ m kg~kg^{-1}}$	limestone additive ratio
$n_{O2}$	$\mathrm{mol}\mathrm{mol}^{-1}$	oxygen ratio
$n_{SC}$	$\mathrm{mol}\mathrm{mol}^{-1}$	steam to carbon ratio
$n_{\mathrm{WHSV}}$	$h^{-1}$	weight hourly space velocity of dry ash free fuel in bed
$Re_{p,mf}$	-	Reynolds number for particle at minimum fluidizing condition
$u_0$	${ m m\ s^{-1}}$	superficial velocity
$u_{\rm mf}$	${ m m~s^{-1}}$	minimal fluidization velocity
$Y_{gas}$	$\rm m^3~kg^{-1}$	gas yield (volume of dry and $N_2$ -free gas in STP conditions per mass of dry ash free fuel)
$y_{i}$	$\mathrm{m}^3~\mathrm{m}^{-3}$	gas concentration (volume fraction)
$y_{j}$ , $\Delta G_{R}$	$kJ \text{ mol}^{-1}$	Gibbs reaction enthalpy

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