Article

Investigation of Non-Thermal Plasma Assisted Charcoal Gasification for Production of Hydrogen-Rich Syngas

Yin Pang 1,*, Thomas Hammer 2, Dominik Müller 1, Jürgen Karl 1

- ¹ Chair of Energy Process Engineering, Friedrich-Alexander-Universität Erlangen-Nürnberg, Fürther Str. 244f, 90429 Nürnberg, Germany; dominik.mueller@fau.de; juergen.karl@fau.de
- ² Siemens AG, Corporate Technology, Research on Energy and Electronics, Günther-Scharowsky-Str. 1, 91058 Erlangen, Germany; thomas.hammer@siemens.com
- * Correspondence: yin.pang@fau.de; Tel.: +49-911-5302-9032

Abstract: The motivation of this work is to investigate experimentally the influence of non-thermal plasma (NTP) application on the reaction kinetics of atmospheric pressure steam gasification of charcoal using a thermostatically controlled drop tube reactor. A gliding-arc generator provides about 1 kW electrical power NTP. For comparison thermal gasification is investigated under comparable flow and specific energy input conditions providing additional heat to the steam. Optical temperature measurement 20 cm flow down of the NTP zone is utilized to characterize the specific enthalpy of the reactive flow. The composition of produced syngas is measured by a gas analyzer and used for the calculation of gas flow rates. The results show a NTP-enhancement on the production of individual syngas components (H₂, CO, CH₄), especially on hydrogen production by around 39%. The syngas-based carbon conversion and hydrogen release are calculated from the carbon and hydrogen balance between the correspondent content in syngas and in the feedstock. The NTP promoted the carbon conversion and hydrogen release by 25% and 31%, respectively. The first-order reaction kinetics are determined by data-fitting in an Arrhenius diagram. The plasma enhanced the reaction rate coefficients by 27%. Based on experimental results and other literature, possible plasma-induced reactions are proposed.

Novelty statement: Non-thermal plasma application has been claimed for decades to enhance the syngas yield of reforming and gasification as compared to conventional thermal processes because of its potential for efficient radical formation at comparatively low temperature. Here evidence is given for this hypothesis by characterizing the enhancement in syngas yield in steam gasification experiments where a fraction of the heat supplied to the process is substituted by electric energy supplied to a non-thermal plasma.

Highlights:

- NTP-Enhancement in syngas-based carbon conversion by 25%
- NTP-Enhancement in syngas-based hydrogen release by 31%
- NTP-Enhancement in reaction kinetics (reaction rate coefficient) by 27%

Keywords: non-thermal plasma; charcoal gasification; carbon conversion; hydrogen release; reaction kinetics; reaction mechanism

Nomenclature

A Pre-exponential factor BDE Bond dissociation energy

DC Direct current
E_A Activation energy

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k Reaction rate coefficient

 $\dot{m}_{C,CO}$ Carbon mass flow of carbon monoxide $\dot{m}_{C,CO2}$ Carbon mass flow of carbon dioxide

 $\dot{m}_{C,CH4}$ Carbon mass flow of methane $\dot{m}_{C,fuel}$ Carbon mass flow of feedstock $\dot{m}_{H,fuel}$ Hydrogen mass flow of hydrogen $\dot{m}_{H,CH4}$ Hydrogen mass flow of methane

 $\dot{m}_{H,plasma}$ Hydrogen mass flow from the plasma-assisted water dissociation

NTP Non-thermal plasma

R Gas constant S/C Steam to carbon T Temperature

Xc Syngas-based carbon conversion YH Syngas-based hydrogen release

1. Introduction

Charcoal or coke is one of the major products from pyrolysis reactions besides gases and tar [1]. Compared to biomass pyrolysis, the gasification of charcoal or coke is much slower due to slow reaction kinetics limited by mass transfer, e.g. pore diffusion and external mass transfer [2]. Increasing the reaction rate of charcoal gasification could have the benefit of a smaller reactor size, lower capital costs and higher output.

Several influence parameters can enhance the charcoal/coke gasification kinetics. Among them, pyrolysis conditions, origin of feedstock, working medium and mineral content are discussed here.

The test condition of pyrolysis determines the structure of charcoal and thus the reaction rate. This could be explained by the influence of macro-, mecro- and micropores generated during pyrolysis. Li et al. [3] prepared char samples at different pyrolysis temperature ranging from 1073 K to 1473 K for the CO₂ gasification reaction. The results showed a decrease in gasification reactivity by increasing pyrolysis temperature. With rising temperature, the pore structure melted, collapsed and jointed together, which led to decrease of specific areas/active sites. Bui et al. [4] studied the influence of pyrolysis pressure between 1 and 20 bar on charcoal gasification kinetics with CO₂ and concluded lower reaction rates at higher pyrolysis pressure.

The origin of feedstock influences the internal chemical and physical structure. Yuan et al. [5] gasified three different char samples from rice straw, chinar leaves and pine sawdust. During gasification, the char structure of pine straw was destructed due to melting and accumulation, which led to smaller specific surface area and lower gasification rate. The sample from chinar leaves, with high porosity and specific surface area, had the highest gasification rate. Wang et al. [6] investigated six char samples in a thermobalance at atmospheric pressure with CO₂. The results showed herbaceous residues (pine sawdust, bamboo sawdust) had a faster gasification reactivity than wooden residues (peanut shell, maize cob, wheat straw, rice lemma). Dahou et al. [7] studied gasification kinetics with water steam using six different feedstock in a thermo-gravimetric balance. Among them, residues from apple orchard, apricot orchard and vineyard showed highest conversion rates, while residues from rice husk and wheat straw reacted with the slowest kinetics.

Different working medium influences reaction kinetics during charcoal gasification. The gasification agent can be chosen by carbon dioxide, water steam, (oxygen-enriched) air and pure oxygen, with increasing reaction kinetics. Yoon and Lee [8] concluded from the experimental results, that the carbon conversion of pure steam gasification is lower than the conversion using steam-air-mixture. Guizani et al. [9] concluded experimentally that the reactivity of char using H₂O as agent was almost two times faster than using CO₂. Reschmeier and Karl [10] determined that the reaction kinetics using water steam were faster than using carbon dioxide.

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Many researchers have studied the catalytic influence of mineral content in the ash on charcoal gasification reactions. Bouraoui et al. [11] varied the ratio of K/Si-ratio between 0.2 and 3.8 during CO₂ gasification at 800°C and concluded an acceleration of reaction by a K/Si-ratio higher than 3 at high conversion degree. Hengel and Walker [12] found a strong catalytic influence caused by calcium-dosage during char gasification.

Recently, research activities are reported in the field of plasma-assisted charcoal/coke gasification.

Tamosiunas et al. [13] used a thermal plasma torch to gasify olive-derived charcoal with water steam. The DC arc plasma torch provided approximately 50 kW to the gasification reactor. The syngas consisted mainly of CO, H₂ and CO₂ with concentrations of 41%, 13% and 13%, respectively. Yoon and Lee [8] gasified charcoal powder using microwave plasma operated by steam-air-mixture. For plasma gasification operated by pure water steam, a maximum hydrogen content of ca. 60% and a cold gas efficiency of ca. 23% have been reported.

The major research focus of this work is to investigate the influence of NTP on charcoal gasification compared to conventional thermal gasification at similar test conditions, in particular with respect to specific energy input.

2. Experiments

2.1. Concept

The major motivation of this work is to investigate the influence of non-thermal plasma (NTP) on charcoal gasification based on a comparison between plasma and thermal cases at similar test conditions. The NTP, also known as non-equilibrium plasma, does not have a thermodynamic equilibrium. Mean electron energy, degree of ionization and concentrations of excited species and radicals correspond to temperatures substantially higher than the kinetic gas temperature. This can only be achieved by taking measures impeding the establishment of thermal equilibrium such as pulsed power supply or strong spatial gradients.

In case of NTP assisted gasification electric power generating the electrical gas discharge plasma substitutes a fraction of the thermal power fed to a conventional gasification process. The NTP generated in the water steam interacts with the charcoal particles by means of ionic reactions, radical reactions and heating. The contact time of the particles with the NTP depends on the type of gasification reactor and the type of NTP. In our experiments a drop tube reactor having a height of 4 m and an alternating current gliding arc generator operated at a frequency of about 90 kHz are utilized. Due to the strongly filamentary character of the gliding arc plasma fluctuating in space and time the contacting time of the particles with the NTP is expected to be very short as compared to the residence time in the drop tube reactor. Since ionization and radical formation in the NTP filaments takes place on the sub-microsecond time scale, thermally induced reactions are slow as compared to NTP induced ion and radical reactions as long as the gas temperature is not too high (i.e. below 3500 K).

In order to give evidence about NTP effects it is essential to perform thermal and NTP assisted gasification under comparable conditions (including feed flows and net specific energy input) and compare the syngas compositions and yields. The specific energy input is characterized by means of optical temperature measurement $15-20\,\mathrm{cm}$ downstream of the plasma zone [14]. Since thermalization due to radical reactions and ion recombination takes place on the time scale of microto milli-seconds this temperature is assumed to be representative for the specific energy input not only for the thermal gasification case but also for the NTP assisted gasification case.

2.2. Test-rig

The test-rig is explicitly described in a previous publication [15]. Figure 1 presents a general overview of the used test facility. A double-lead screw conveyor (Coperion K-TRON, K-CV-KT20) feeds charcoal powder with nitrogen purge into the reactor. The nitrogen is also used as a tracer for gas analysis. A steam generator provides the gasification reaction with two steam flows: the primary

water steam is fed into the plasma generator as working medium and the secondary steam flow is used to vary the S/C-ratio. In the reference thermal gasification, a steam superheater heats the primary water steam to achieve similar gas temperatures.

The non-thermal plasma generator has two gliding-arc electrodes with a length of ca. 1.1 m. The plasma arc is triggered by max. 10 kV between a gap distance of electrodes of around 2 mm. In the upper part of the window zone, in which two rectangular quartz glasses are installed, the generated fluctuating plasma arc can be observed and measured optically [14-15], in which the results showed no substantial deviations in gas temperature between NTP and thermal cases.

The drop tube reactor consists of preheating, window, reaction and quench zones. An electrical heater heats and maintains the preheating, window and reaction zones to a defined experimental temperature. The heating of the secondary water steam takes place in the preheating zone. In the beginning of the window zone, the plasma arc is triggered, and two water steam flows joint together. The fed charcoal particles pass through the fluctuating plasma area and react with the plasma mixture. A vertical moveable sampling probe withdraws the produced syngas at a defined position for gas analysis.

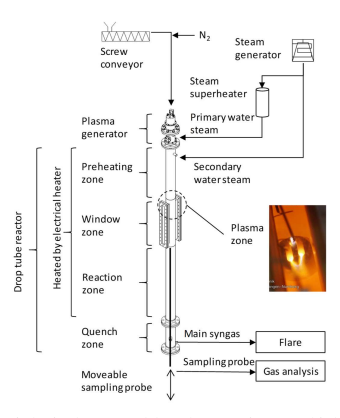


Figure 1. Test facility for plasma-assisted charcoal steam gasification (modified from [15])

2.3. Test Conditions

Table 1 describes the property of charcoal powder used in this work compared to two literature works. The determination of moisture, proximate analysis and ultimate analysis in this work were carried out according to German DIN standards [16–20]. Despite the moisture content, the feedstock used in this work and in literature, contains a similar elementary composition.

Table 1. Proximate and ultimate analysis of charcoal samples used in this work and comparable literature

Sources:	This work	Tamosiunas et al. [13]	Yoon and Lee [8]
Fuel Type	Charcoal (derived from wood)	Charcoal (derived from Olive pomace)	Charcoal (derived from oak)
Moisture (wt.%)	4.6	22	0.6
Proximate analysis			
(wt.%, dry)			
Volatile	21.3	17.4	27.6
Ash	2.3	5.6	1.4
Fixed carbon	76.4	77	70.4
Ultimate analysis			
(wt.%, dry)			
Carbon	84.5	80.4	83.3
Hydrogen	3.1	2.9	3.6
Nitrogen	0.5	0.4	0.4
Oxygen	9.2	16.3	11.2

Table 2 presents the test conditions in this work compared to the selected literature. In this work, the particle size of charcoal powder varies between 45 and 250 μ m with two categories. A mass flow of 0.16 kg/h of charcoal powder is fed with 4 l/min nitrogen purge into the reactor. The sum of primary and secondary water steam flows is 8 kg/h. The plasma generator supplies totally 0.9 kW power into the reaction. The measured operating current amplitude lying between 0.4 and 0.7 A shows clearly non-thermal characteristics, s. [15]. The drop tube reactor has around 0.1 m in diameter and 4 m in length. The wall temperature of the reactor is measured by thermocouples (type K) between 700 and 950 °C. The gasification takes place at atmospheric pressure.

Beside plasma and thermal gasification in this work, another test series has been conducted to determine the hydrogen production due to water dissociation. The only difference compared to plasma gasification is no fuel feeding.

The simplified charcoal structure has been adopted from Backreedy et al. [21], s. Figure 2. This is used for the discussion of NTP reaction mechanism in chapter 3.6.

Figure 2. Simplified biomass char matrix (adopted from [21])

Table 2. Test conditions of plasma-assisted charcoal gasification of this work and comparable literature

Sources:	This work	Tamosiunas et al. [13]	Yoon and Lee [8]
Feedstock			
Particle size (µm)	45 – 125; 125 – 250;	< 2000	75
Mass flow (kg/h)	0.16	4.7	1.3
Nitrogen purge	4	-	-
(l/min)			
Steam			
Flow rate (kg/h)	8	12.7	2.2
Temperature (°C)	270 (plasma);	n.a.	> 100
	650 (thermal)		
Plasma			
Electric power (kW)	0.9 (plasma);	50	5
	0 (thermal)		
Current amplitude	0.4 - 0.7 [15]	180	n.a.
(A)			
Electric heater			
Heating power (kW)	2.5	-	-
Reactor	cylindrical	cylindrical	cylindrical
Diameter (m)	0.1	0.4	0.058
Length (m)	4	1	0.1
Wall temperature (°C)	700 – 950	n.a.	n.a.
Pressure	atm.	atm.	atm.

n.a.: not available; atm.: atmospheric

3. Results

3.1. Syngas Composition

The syngas composition, including H₂, CO, CH₄ and CO₂, is measured by a gas analyzer (ABB, AO2020 system). The temperature, used as the vertical axis in this chapter, is the gas temperature which is optically measured inside the drop tube reactor [14-15].

Figure 3 shows the syngas composition in case of plasma and thermal gasification with systematic errors at dry-basis. At lower temperatures, large differences in hydrogen content can be explained by the plasma-induced reactions. The C-H bonds (s. Figure 2) are decomposed by possible electron impact forming hydrogen atoms and further hydrogen molecules. With the rising gas temperature, the hydrogen content increases rapidly in the thermal case due to enhanced steam gasification (1). Thus, the difference between two cases gets smaller.

$$CH_nO_m + (1-m)H_2O \rightarrow CO + (0.5n + 1-m)H_2$$
 (1)

The concentration of methane and carbon monoxide decreases with increasing gas temperature in the reactor. The content of carbon dioxide decreases slightly.

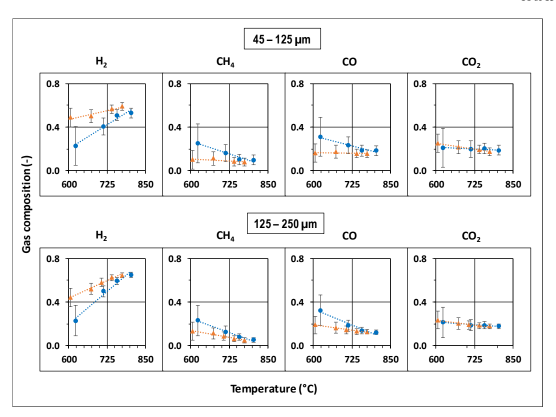


Figure 3. Syngas composition with dotted trend lines and systematic errors at dry-basis (orange: plasma; blue: thermal)

The measured syngas composition at 766 $^{\circ}$ C is summarized in Table 3 and compared to literature at similar test conditions. The comparison shows the results of this work are consistent to literature under consideration of systematic errors.

Table 3. Produced syngas composition in case of plasma and thermal charcoal gasification compared to literature

Sources:	H ₂ (vol.%)	CO (vol.%)	CH4 (vol.%)	CO ₂ (vol.%)
d = 45 – 125 mm, p	58.9 ± 4	15.8 ± 4	7.9 ± 4	17.5 ± 4
d = 45 - 125 mm, t	51.0 ± 5	18.4 ± 5	10.4 ± 5	20.3 ± 5
d = 125 - 250 mm. p	64.7 ± 2	12.8 ± 2	4.8 ± 2	17.7 ± 2
d = 125 - 250 mm, t	59.6 ± 3	13.7 ± 3	7.9 ± 3	18.8 ± 3
Tamosiunas et al. [13]	41.2	13	-	18.7
Yoon and Lee [8]	60	17	3	21

p: plasma case; t: thermal case

3.2. Syngas Production

The syngas production is calculated by its concentration and the known flow rate of nitrogen purge.

Figure 4 presents the syngas production from plasma-assisted and thermal charcoal gasification with systematic errors at dry-basis. With increasing gas temperature, individual syngas production was increased. Besides, NTP had a clear influence on the production of H₂, CO and CO₂.

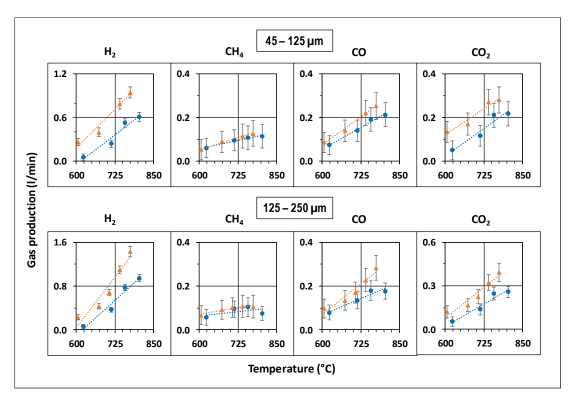


Figure 4. Syngas production with dotted trend lines and systematic errors at dry-basis (orange: plasma; blue: thermal)

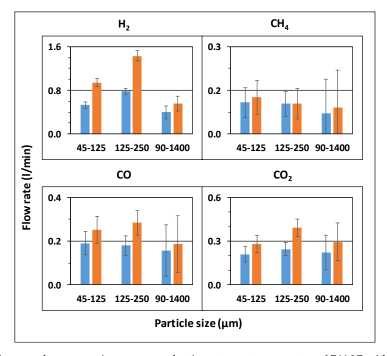


Figure 5. Plasma enhancement in syngas production at a gas temperature of 766 $^{\circ}$ C with systematic errors

The plasma enhancement in gas production at a gas temperature of 766 °C is shown in Figure 5. The results of particle size lying between 90 and 1400 μm are adopted from a previous own publication [15]. The NTP has promoted the production of hydrogen, methane, carbon monoxide and carbon dioxide by 39%, 8%, 26% and 29%, respectively.

3.3. Carbon Conversion

The syngas-based carbon conversion Xc is calculated by the carbon balance between feedstock and syngas products:

$$X_{C} = \frac{\dot{m}_{C,CO} + \dot{m}_{C,CO2} + \dot{m}_{C,CH4}}{\dot{m}_{C,fuel}}$$
(2)

with $\dot{m}_{C,fuel}$, $\dot{m}_{C,CO2}$ and $\dot{m}_{C,CH4}$ as the carbon mass flow of fuel, carbon monoxide, carbon dioxide and methane, respectively.

Figure 6 presents the calculated syngas-based carbon conversion in case of plasma and thermal charcoal gasification with systematic errors. In a), a temperature dependence between gas temperature in the reactor and carbon conversion is shown which can be explained by the undergoing chemical reactions. In b), the conversion degree is independent of particle size which indicates the mass transfer as the major influence factor. In average, the NTP has increased the carbon conversion by 25%.

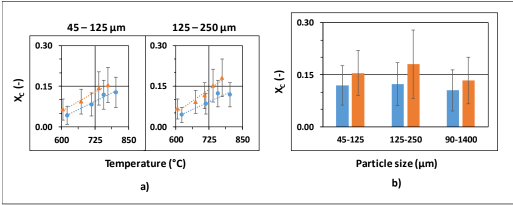


Figure 6. Calculated syngas-based carbon conversion in case of plasma (orange) and thermal (blue) charcoal gasification with systematic errors: a) variation of gas temperature; b) variation of particle size at $T=766\,^{\circ}C$

According to

Table 1, the fixed carbon content regarding the total mass is 72.9% (= 76.4% x (1 - 4.6%)). Therefore, the carbon content in the volatile matter can be calculated to 7.7% (= 84.5% x (1 - 4.6%) - 72.9%). With the assumption of equal composition of carbon content in tars and in gases, the expected Xc after a complete pyrolysis with regard to the total mass should be 3.9% (=7.7% * 0.5). In case of 45 – 125 µm charcoal gasification at 766 °C, the carbon conversion Xc is calculated to be ca. 15%, which corresponds to a conversion degree of 12.1% (= 15% x 84.5% x (1 - 4.6%)) with regards to the total mass. Considering the possible tar reforming, the released mass content of fixed carbon during gasification is between 4.4% (= 12.1% - 7.7%) and 8.2% (= 12.1% - 3.9%). The authors want to point out that the low carbon conversion degree is due to the limited particle residence time of less than 1 second.

3.4. Hydrogen Release

In this work a parameter Y_H , defined as syngas-based hydrogen release, is calculated from the hydrogen balance between the hydrogen mass flow in generated syngas ($\dot{m}_{H,CH4}$, $\dot{m}_{H,H2}$) minus the hydrogen production from the plasma-assisted water dissociation ($\dot{m}_{H,plasma}$) and the hydrogen mass flow of feedstock ($\dot{m}_{H,fuel}$) in feedstock. The authors want to point out that the hydrogen release is not equal to the hydrogen conversion from feedstock into syngas, because a certain amount of hydrogen comes from the water steam. Thus, the value of syngas-based hydrogen release can possibly exceed one.

$$Y_{H} = \frac{\dot{m}_{H,CH4} + \dot{m}_{H,H2} - \dot{m}_{H,plasma}}{\dot{m}_{H,fuel}}$$
(3)

Besides, the measured hydrogen production from the plasma-assisted water dissociation is potentially lower than the actual production during plasma gasification, because the H- and OH-radicals can recombine to form water. In case of charcoal gasification with plasma, the generated H- and OH-radicals can react with feedstock. In summary, the calculated syngas-based hydrogen release offers a possibility to quantitatively determine the influence of NTP on the hydrogen production from charcoal gasification.

Figure 7 presents the hydrogen release in case of plasma-assisted and thermal gasification with systematic errors. In a), similar temperature-dependent trend lines have been observed as those in Figure 6. In b), the hydrogen release of different particles has been compared at $T = 766 \,^{\circ}\text{C}$ in the reactor. An average NTP-enhancement of 31% has been calculated.

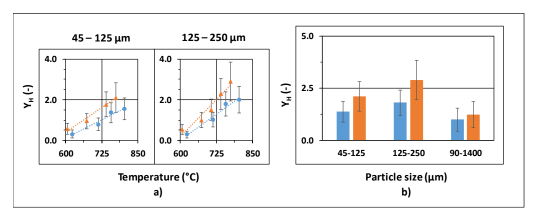


Figure 7. Calculated syngas-based hydrogen release in case of plasma (orange) and thermal (blue) charcoal gasification with systematic errors: a) variation of gas temperature; b) variation of particle size at $T=766\,^{\circ}C$

The relative high value of Y_H in case of $125-250~\mu m$ can be explained by the fluctuation of screw conveyor, which is already considered in the shown systematic errors.

3.5. Reaction Kinetics

The reaction kinetics are calculated by data-fitting in an Arrhenius diagram with first order, s. [15].

Figure 8 shows the calculated reaction kinetics in case of plasma and thermal charcoal gasification with systematic errors. In a), the reaction kinetics become faster with the rising gas temperature inside the reactor. In b), the particle size does not influence the reaction rate coefficient and the NTP promotes the reaction rate coefficient *k* by 27% in average.

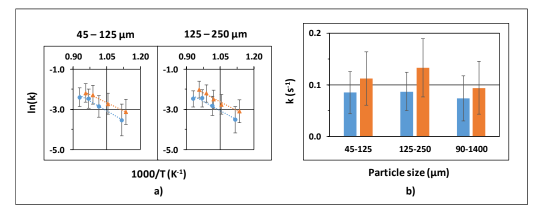


Figure 8. Calculated reaction kinetics in case of plasma (orange) and thermal (blue) charcoal gasification with systematic errors: a) variation of gas temperature; b) variation of particle size

The calculated reaction kinetics are summarized in Table 4. The calculated activation energies of plasma-assisted are lower than that of thermal gasification, which confirms the hypothesis of the NTP-induced kinetics promotion. However, the authors want to emphasis the complexity of the overall system including conventional heterogeneous gas-solid and plasma-induced reactions. The results in Table 4 are combined with considerable systematic errors, which is shown in Figure 8.

Table 4.	Calculated reaction	kinetics of own 1	esults (k=A*ex	p(-EA/RT)	, first reaction order)

Type of reaction	Particle size (µm)	Ea (kJ/mol)	A (s-1)
plasma-assisted	45 - 125	43.8	17.7
thermal	45 - 125	52.6	35.2
plasma-assisted	125 - 250	49.8	38.7
thermal	125 - 250	50.0	25.4
plasma-assisted	90 - 1400	34.0	4.6
thermal	90 – 1400	38.4	6.3

3.6. Proposed Reacton Mechanism based on Literature and own Results

Based on the own results and results in literature, the following plasma-induced reaction mechanism is proposed. The following reactions offer possible pathways of enhanced syngas production in case of plasma-assisted charcoal gasification. However, they represent only a small (possible) part of total plasma-assisted reaction mechanism.

The ionization and dissociation reaction takes place due to an electrical energy input, generating radicals, electrons and ions.

$$H_2O$$
 + electrical energy \rightarrow (4) radicals(e.g. ·H, ·OH) + charged particles(e.g. e-, H-, OH-) + excited particles + UV irradiation

Depending on the amount of electrical energy, the ionization and dissociation products can vary [22]. Thus, the reaction (4) gives only a general expression. For the purpose of simplicity, the negative charge of electron is neglected in the following reactions. In an electrical field, the ionized electron

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can be accelerated to achieve high kinetic energy (e.g. 1 - 10 eV [23]). The electrons with high kinetic energy are called as energetic electrons (e *).

$$e + electrical field \rightarrow e^*$$
 (5)

The enhancement of gas production, carbon conversion and hydrogen release can be explained by electron impact and radical-induced reactions. The symbol *R*- represents carbon-containing functional groups in charcoal structure.

$$e^* + R - H \rightarrow R + H + e \tag{6}$$

$$e^* + R-O-R \rightarrow R + O-R + e \tag{7}$$

The electron impact reactions (e.g. (6)-(7)) could cause the decomposition of chemical bonds, which indicates an energy transfer from kinetic energy into bond dissociation energy (BDE). According to BDEs listed in Table 5, the chemical bonds could be separated by the kinetic energy of energetic electrons. The probability of these reactions can be determined by the calculation of their reaction enthalpies, which is not done in this work.

Table 5. Selected bond dissociation energy of common bonds [24]

Type of bonds	BDE (kJ/mol)	BDE (eV)
H-O	459	4.8
С-Н	411	4.3
C-O	358	3.7
C=O	799	8.3
C-C	346	3.6
C=C	602	6.2
C≡C	835	8.7

The radicals such as ·H and ·OH, generated from water dissociation, have a high oxidation potential, which could also contribute to the destruction of carbon-carbon bonds in charcoal structures (e.g. (8)-(11)). These reactions have been investigated by Mulcahy and Young [25] with first reaction order. The test results showed the reaction rate of $200 - 710 \, \text{s}^{-1}$ and $100 - 360 \, \text{s}^{-1}$ at 298 K for (8)-(9) and for (10)-(11), respectively.

$$2\cdot OH + C \rightarrow CO + H_2O \tag{8}$$

$$4 \cdot OH + C \rightarrow CO_2 + 2H_2O \tag{9}$$

$$\cdot OH + C \rightarrow CO + 0.5H_2 \tag{10}$$

$$2 \cdot OH + C \rightarrow CO_2 + H_2 \tag{11}$$

The intermediate (radical) products could react with each other (e.g. (12)-(15)), producing possible final products such as water, hydrogen, and carbon dioxide. This could explain the enhanced gas production shown in Figure 5.

$$\cdot H + \cdot OH \rightarrow H_2O \tag{12}$$

$$\cdot \mathbf{H} + \cdot \mathbf{H} \to \mathbf{H}_2 \tag{13}$$

$$\cdot OH + \cdot OH \rightarrow H_2O_2 \rightarrow H_2O + 0.5O_2 \tag{14}$$

$$CO + H2O \rightarrow CO2 + H2$$
 (15)

The authors want to emphasize that the proposed reaction mechanism based on electron impact and radical oxidation is only a part of the total complex plasma-induced reaction mechanism. Other influencing factors, such as ions, excited particles and UV irradiation, could also contribute to fuel conversion. A review paper of plasma gasification/reforming with focus on reaction mechanism will be published in the near future.

4. Conclusions

In this work, charcoal gasification with water steam non-thermal plasma took place in a drop tube reactor. The particle size was varied between 45 and 250 µm with two categories.

The non-thermal plasma enhanced the production of H_2 , CH_4 , CO and CO_2 by 39%, 8%, 26% and 29%, respectively. The carbon conversion X_C and the hydrogen release Y_H were calculated from the individual carbon and hydrogen balance between the corresponding content in the syngas and the content in feedstock. The NTP-enhancement in X_C and Y_H were 25% and 31% respectively. The reaction kinetics were determined by data-fitting in an Arrhenius diagram and accelerated by 27% in plasma case. Based on the experimental results, plasma-induced reaction mechanism based on electron impact and radical reactions was proposed.

Author Contributions: Yin Pang conceived, designed and performed the experiments; All authors analyzed and evaluated the results; Yin Pang wrote the paper.

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References

- [1] J. Karl, "Dezentrale Energiesysteme, Neue Technologien im liberalisierten Energiemarkt." De Gruyter, Berlin, Boston, 2012.
- [2] J. Karl and T. Pröll, "Steam gasification of biomass in dual fluidized bed gasifiers: A review," *Renew. Sustain. Energy Rev.*, vol. 98, pp. 64–78, 2018.
- [3] R. Li, J. Zhang, G. Wang, X. Ning, H. Wang, and P. Wang, "Study on CO2 gasification reactivity of biomass char derived from high-temperature rapid pyrolysis," *Appl. Therm. Eng.*, vol. 121, pp. 1022–1031, 2017.
- [4] H.-H. Bui, L. Wang, K.-Q. Tran, and Ø. Skreiberg, "CO2 gasification of charcoals produced at various pressures," *Fuel Process. Technol.*, vol. 152, pp. 207–214, 2016.
- [5] S. Yuan, X. Chen, J. Li, and F. Wang, "CO2 Gasification Kinetics of Biomass Char Derived from High-Temperature Rapid Pyrolysis," *Energy & Fuels*, vol. 25, no. 5, pp. 2314–2321, May 2011.
- [6] G. Wang *et al.*, "Experimental and modeling studies on CO2 gasification of biomass chars," *Energy*, vol. 114, pp. 143–154, 2016.
- [7] T. Dahou *et al.*, "The Influence of Char Preparation and Biomass Type on Char Steam Gasification Kinetics," *Energies*, vol. 11, no. 8, 2018.
- [8] S. J. Yoon and J.-G. Lee, "Hydrogen-rich syngas production through coal and charcoal gasification using microwave steam and air plasma torch," *Int. J. Hydrogen Energy*, vol. 37, no. 22, pp. 17093–17100, 2012.
- [9] C. Guizani, M. Jeguirim, R. Gadiou, F. J. E. Sanz, and S. Salvador, "Biomass char gasification by H2O, CO2 and their mixture: Evolution of chemical, textural and structural properties of the chars," *Energy*, vol. 112, pp. 133–145, 2016.
- [10] R. Reschmeier and J. Karl, "Experimental study of wood char gasification kinetics in fluidized beds," *Biomass and Bioenergy*, vol. 85, 2016.
- [11] Z. Bouraoui, C. Dupont, M. Jeguirim, L. Limousy, and R. Gadiou, "CO2 gasification of woody biomass chars: The influence of K and Si on char reactivity," *Comptes Rendus Chim.*, vol. 19, no. 4, pp. 457–465, 2016.
- [12] T. late T. D. Hengel and P. L. Walker Jr., "Catalysis of lignite char gasification by exchangeable calcium and magnesium," *Fuel*, vol. 63, no. 9, pp. 1214–1220, 1984.
- [13] A. Tamošiūnas *et al.*, "The Potential of Thermal Plasma Gasification of Olive Pomace Charcoal," *Energies*, vol. 10, no. 5, 2017.
- [14] L. A. Bahr *et al.*, "Temperature determination of superheated water vapor by rotational-vibrational Raman spectroscopy," *Opt. Lett.*, vol. 43, no. 18, pp. 4477–4480, Sep. 2018.
- [15] Y. Pang et al., "Plasma-Assisted Biomass Gasification with Focus on Carbon Conversion and Reaction Kinetics Compared to Thermal Gasification," *Energies*, vol. 11, no. 5, 2018.
- [16] "DIN 51718, Prüfung fester Brennstoffe Bestimmung des Wassergehaltes und der Analysenfeuchtigkeit (DIN 51718:2002-06)."
- [17] "DIN 51719, Prüfung fester Brennstoffe Bestimmung des Aschegehaltes (DIN 51719:1997-07)."
- [18] "DIN 51720, Prüfung fester Brennstoffe Bestimmung des Gehaltes an Flüchtigen Bestandenteilen (DIN 51720:2001-03)."
- [19] "DIN 51734, Prüfung fester Brennstoffe Immediatanalyse und Berechnung des Fixen Kohlenstoffs (DIN 51734:2008-12)."
- [20] "DIN 51732, Prüfung fester Brennstoffe Bestimmung des Gesamtgehaltes an Kohlenstoff, Wasserstoff und Stickstoff Instrumentelle Methoden (DIN 51732:2014-07)."
- [21] R. I. Backreedy, J. M. Jones, M. Pourkashanian, and A. Williams, "Modeling the reaction of oxygen with coal and biomass chars," *Proc. Combust. Inst.*, vol. 29, no. 1, pp. 415–421, 2002.
- [22] N. S. J. Braithwaite, "Introduction to gas discharges," Plasma sources Sci. Technol., vol. 9, no. 4, p. 517, 2000.
- [23] A. Fridman, *Plasma chemistry*. Cambridge university press, 2008.

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[24] "Wired Chemist - Common Bond Energies and Bond Lengths," 2018. [Online]. Available: http://www.wiredchemist.com/chemistry/data/bond_energies_lengths.html. [Accessed: 26-Sep-2018].

[25] M. F. R. Mulcahy and B. C. Young, "The reaction of hydroxyl radicals with carbon at 298 K," *Carbon N. Y.*, vol. 13, no. 2, pp. 115–124, 1975.