Experimental Study of Steam and Carbon Dioxide Microwave Plasma for Advanced Thermal Treatment Application

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Abstract: Pollution reduction from waste management and energy generation is necessary to mitigate climate change and is one of the major challenges of the 21st century. This can be achieved through the development of innovative energy recovery technologies from biomass and wastes, such as microwave plasma gasification. An envelope of stable CO2 plasma operation is described, by varying working gas flow rate at applied microwave powers between 1 and 6 kW, whereas H2O plasma operation is possible with flow rate ranging from 20 to 50 g/min and microwave powers between 2.5 and 6 kW. The temperature generated in a large chamber connected to the plasma torch is recorded, reaching up to 850°C, showing a heterogeneous temperature distribution. In addition, optical emission spectroscopy measurements provide an insight into plasma chemistry and demonstrate the dissociation of CO2 and H2O molecules at extremely high temperatures of up to 6,300°C assuming local thermodynamic equilibrium. The experimental results demonstrate that the microwave plasma torch provides an ideal environment for gasification with high temperature and very chemically reactive species. This study provides valuable information for the design of microwave plasma gasification reactors with great potential for effective solid feedstock conversion into high quality syngas for energy production.

Keywords: Microwaves, Plasma, Carbon dioxide, Steam, Gasification

Abbreviation:
ATT: Advanced Thermal Treatment
DC: Direct Current
MPT: Microwave Plasma Torch
OES: Optical Emission Spectroscopy
SLPM: Standard Litre per Minute
SOFC: Solid Oxide Fuel Cell
1. Introduction

Industrialisation of society has enabled an enhanced standard of living, facilitated by easier access to electricity and goods. The consequence of this, however, is the emission of greenhouse gases, from fossil fuel combustion, which in turn contribute towards climate change [1]. In parallel the industrialisation of society is also driving the ever-increasing generation of waste which itself causes unintended health, safety and environmental issues, all of which are exacerbated in emerging economies [2]. As such, the provision of affordable, reliable and low-carbon energy and the development of sustainable waste management practices remain to be two of the largest challenges to mankind.

Today it is estimated that circa 70% of the solid waste generated globally is either sent to landfill or dumped [2], of which both practices are recognised as being the least sustainable option according to the Waste Hierarchy presented in European Directive 2008/98/EC [3]. In contrast, the directive stipulates that the optimum solutions for waste management are to reduce and reuse wastes at source, with recycling and energy recovery being the next best options should avoidance of waste generation not be possible [3]. As such, Governments across Europe have set ambitious targets, with respect to waste management, with an ultimate aim for less than 10% of all wastes generated to be disposed of to landfill by 2030 [4]. Energy recovery is recognised as the best treatment option for non-recyclable waste that would otherwise be disposed of to landfill. Waste can be used as a fuel for low-carbon energy generation and in doing so offset demand for fossil fuel derived energy. Incinerators are currently the predominant waste-to-energy technology deployed globally for waste destruction and subsequent energy recovery, whereby facilities are often large centralised units (increasing waste miles), that operate at low efficiencies, whilst also requiring significant exhaust gas treatment in order to comply with strict emissions regulations [5]. However, given the increasing transition of energy and waste management companies towards the development of distributed energy generation and waste management assets, an alternative opportunity exists whereby low-carbon energy can be generated from waste at source. Advanced thermal treatment (ATT), comprising gasification and pyrolysis process plant are one such emerging group of technologies that may present a significant opportunity for decentralised energy generation and waste management and in doing so reducing the polluting and costly requirement to transport waste to centralised facilities. Gasification can be defined as the thermal decomposition of a solid feedstock in an oxidizing environment, leading to the generation of a synthetic gas (syngas) composed of mainly CO and H₂. The resultant syngas from gasification process can be used for electricity or heat generation, using either a gas engine, gas turbine or fuel cell. One of the many virtues of gasification is its environmental performance [5]. This is especially true for thermal plasma gasification which can achieve almost complete waste conversion as well as the production of a clean syngas due to the exceptional properties of plasma [6, 7].

According to Fridman, plasma can be defined as the fourth state of the matter after solids, liquids and gases [8]. Plasma is formed through the phenomenon of ionization, which is the dissociation of electrons from atoms or molecules, leading to the formation of positively charged ions. The presence of ions, electrons, excited molecules and photons makes the plasma very reactive [8], enhancing chemical reactions similarly to the presence of a catalyst [9]. The unique properties of plasma have been employed in many applications such as surface treatments (etching, chemical vapour deposition or thermal spray coatings), metallurgy, sterilization of water and air, medicine (treatment of burns, ulcers or other skin diseases) and many others [8]. Whereas the study of the plasma torch presented herein could be beneficial for its application in various technologies, the author’s main interest is on the development of an ATT technology for waste and biomass energy conversion.

One of the advantages of plasma is its high energy density, enabling the design of compact gasification reactors whereby only a short feedstock residence time is required [6, 10]. Plasma gasification can be applied to a large range of feedstocks such as automobile residues, sludge, asbestos fibres, medical waste, municipal solid waste or refused-derived fuel [6, 10]. However, one of the challenges for the use of syngas from gasification is the presence of impurities that can damage the downstream electricity-generation devices. Typical impurities found in syngas, derived from standard gasifiers, are tars, sulphur and chloride compounds [11]. However, in plasma gasifiers, due to the high operational temperatures, tars within the syngas are broken down in situ resulting in a cleaner product gas [6, 9]. As such, plasma gasification offers a mechanism to reduce the cost and sophistication of syngas cleaning trains compared to that of conventional gasification systems. As a result plasma gasification systems are much more suited for integration with more efficient power generation technologies, such as solid oxide fuel cells (SOFC) achieving high electrical conversion efficiencies of between 55 and 65% [12].

At present, most plasma gasification systems are based on the use of direct current (DC) plasma torches. DC plasma torches operate by passing a gas, usually air, between two electrodes of high potential difference. This difference causes an arc to form between electrodes, and subsequently ionises the passing gas, creating a plasma. However, the major drawback of using DC torches relates to the lifetime of the electrodes, which will typically need replacing every 200 hours, especially when operated with oxidative gases [10]. A promising alternative is the use of microwaves for
Microwave plasma has been receiving a growing interest in recent years especially with respect to its applications in ATT [9, 13]. One of the advantages of microwave plasma, compared to conventional DC systems, is the possibility to operate with a wide range of working gases. ATT experiments using microwave plasma have been reported using N₂ [14-16], Ar [17-19], air [20], H₂O [21, 22] or CO₂ [23], as the ionising gas. Ar and N₂ can be used for pyrolysis treatment, however they are not preferred because they are inert gases that do not directly take part in chemical reactions, whilst the presence of non-combustible Ar and N₂ in the generated pyrolysis gas tends to lower its calorific value and quality [10]. Air has the advantages of being free and brings O₂ leading to exothermic oxidation reactions, supplying energy within the reactor while enhancing the carbon conversion rate [22, 24]. On the other hand, excess O₂ also causes combustion of combustible gases lowering the generated syngas calorific value [22, 24]. In addition, the N₂ content in air dilutes the syngas and lowers its heating value. Microwave plasma is a relatively expensive technology that is expected to generate a very high-quality syngas. H₂O and CO₂ are alternative gases that can have the oxidant role in gasification process. H₂O is a good candidate as it can generate syngas with a H₂ content that can exceed 50% by volume, and in doing so enhancing the heating value of the syngas [21, 22, 24]. H₂O is also readily available and could be generated using heat from hot syngas or the exhaust of an electricity generation device. H₂O has excellent reforming properties that will serve to enhance gasification reactions [5]. Another gas that has preferable chemical activity for gasification applications is CO₂, particularly through dry reforming and Boudouard reactions [5]. As such, reintroduction of CO₂ rich streams into the gasification reactor, such as the anode exhaust of a SOFC, would not only serve to enhance the gasification process, but also reduce process emissions of CO₂ to the atmosphere, mitigating against climate change.

Even though plasma gasification can produce a very clean syngas, the high energy consumption of the plasma torch tends to reduce the overall efficiency of the system and limit its deployment. That is why it is particularly important to have a thorough understanding of the plasma torch capabilities in order to optimise their integration within gasifiers. This study presents the range of operating conditions of which a stable microwave-induced plasma can be generated using H₂O and CO₂. As such this manuscript presents, for the first time, an empirical data set that describes the complete operating envelope of microwave induced plasma using H₂O and CO₂ as the ionisation gases. In addition, the operational temperature of the torches has been recorded within a large chamber yielding critical data to inform the sizing and configuration of a plasma reactor. An insight into the plasma chemistry through optical emission spectroscopy (OES) measurements is also presented, highlighting the presence of active species beneficial for efficient thermal conversion of solid feedstocks.

2. Materials and Methods

2.1. The microwave plasma torch

Figure 1 presents a schematic diagram of the experimental setup for this study. The main system of the experimental rig is the microwave plasma torch (MPT), comprising of a Downstream plasma source (Sairem SAS, Neyron, France). The microwave generator is a Sairem GMP G4 60K T400 that can generate microwaves at a frequency of 2.45 GHz up to a power of 6 kW. The microwaves propagate through standard WR340 waveguides equipped with a 3 stub-tuner and a sliding short circuit for microwave transmission optimisation. The microwave tuning devices enable the reflected microwave power to be minimized to less than 1% of the forwarded microwave power whilst maintaining a stable plasma.
The working gas is injected inside a quartz tube 450 mm long and with an outside and inside diameter of 30 and 25.6 mm respectively. The working gas flow rate is regulated using a mass flow controller (Alicat Scientific, Tucson AZ, USA) with a maximum capacity of 100 standard litre per minute (SLPM). The working gas can be pre-heated to temperatures exceeding 150°C using a high temperature oven. Process H₂O is generated using an E-3000 precision steam generator (Cellkraft, Stockholm, Sweden) operating at flow rates spanning 0.1 to 50 g/min at temperatures up to 200°C. The MPT is manually ignited by inserting a tungsten rod in the quartz tube from the top. For operation of a H₂O plasma, the working gas is pre-heated to >150°C, enabling H₂O to be bled into the working gas stream whilst avoiding condensation.

### 2.2. Temperature measurements

An insulated chamber is connected to the bottom chimney of the downstream plasma source as depicted in Figure 2. The chamber is fitted with three thermocouples. The aim of the insulated chamber is twofold, a) to safely capture the exhaust gases and b) to undertake temperature measurements to inform the design of a bespoke ATT reactor. The chamber is of cylindrical construction with a length of 50 cm and a diameter of 25 cm corresponding to an inside volume of 24.5 litres. The chamber is insulated with an inside layer of approximately 2 cm of fire cement on the side and bottom walls, and a 5 cm layer of fibre wool on the outside. In order to maximise operational temperatures within the chamber, the process exhaust gas outlet is located close to the downstream source on the side of the chamber to increase gas residence time. The thermocouples, named top, middle and bottom, are located across the operational profile of the plasma in order to capture operational temperatures across the plasma plume. The top, middle and bottom thermocouples are located at distances of 20, 50 and 70 cm from the centre of the waveguide, respectively. The top thermocouple is horizontal to the tip of the quartz tube and is not directly exposed to the plasma plume. In addition, the thermocouples protrude 3 cm into the chamber beyond the refractive lining and measure the temperatures close to the sidewalls. Once the plasma is ignited, the temperatures throughout the chamber are characterised by a rapid increase within the first few minutes, followed by a linear increase to an experimental maxima, which is typically achieved within 1 hour of operation. For example, the temperature at the middle thermocouple for a 20 SLPM CO₂ plasma at 3 kW of microwave power will reach 480°C within two minutes after plasma ignition, then taking approximately 45 minutes to reach a temperature maxima of 630°C. The experimental temperature maxima is defined when the average temperature increase rate is below 2°C per minute over a period of 5 minutes.

### 2.3. Optical emission spectroscopy

Plasma is generated in the Downstream MPT within the quartz tube. As such, a custom-made chimney, 14 cm long with a series of six holes as presented in Figure 2, was engineered and connected to the lower part of the MPT to enable optical emission spectroscopy (OES) measurements to be taken. The plasma spectra can therefore be measured in seven different locations along the plasma plume at intervals of 0, 6, 8, 9.5, 11, 12.5 and 14 cm. The measurement port located at 0 cm is horizontal to the middle of the waveguide and corresponds to where the plasma is formed, sometimes called the plasma heart. The port located at 6 cm corresponds to the top of the OES chimney, representing the closest point at which a chamber could be integrated with the MPT. OES measurements were captured using a HR2000+ES spectrometer (Ocean Optics Inc., Largo, FL, USA) measuring in the range 190-1100 nm, with an optical resolution of approximately 0.9 nm FWHM (Full Width at Half Maximum). A 74-UV collimating lens is used to receive the plasma emissions, which are then transmitted to the spectrometer via optical fibre. Analysis of plasma spectra and temperature estimations are undertaken by comparing the experimental data with computed data from the software Specair [25].

### 3. Results

#### 3.1. Operating conditions

This section presents the operating conditions, including the applied microwave power and the working gas flow rate, for which a stable plasma can be achieved within the experimental MPT unit. However, prior to presenting the results, it is first important to define the vocabulary employed in this section. Firstly, the plasma operation is defined as “steady” when it can be sustained for >10 mins without visible flickering of the plume, whereby when in this “steady” state, it is expected to operate for several
hours without issues. Relating to this, it was empirically
determined that the quartz tube starts melting within less
than 5 minutes of non-suitable operating conditions.
Secondarily, plasma is defined as “unsteady” when the
plasma cannot be sustained >10 mins. The third definition
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The experimentally determined range of operating
conditions for the MPT are presented in Figures 3 and 4 for
CO₂ and H₂O respectively. A “steady” CO₂ plasma can be
maintained for flow rates ranging from 20 to at least
80 SLPM at an applied microwave power ranging from 1
to 6 kW. Concerning H₂O operation, a “steady” plasma can
be sustained for flow rates from 20 to at least 50 g/min at
an applied microwave power ranging between 2.5 and
4 kW. When increasing the applied microwave power over
4 kW, it is necessary to gradually increase the H₂O flow
rate, a minimum H₂O flow rate of 30 and 40 g/min is
required at 5 and 6 kW microwave power respectively, in
order to keep a “steady” H₂O plasma operation.

There are three main reasons causing the plasma to be
“unsteady”. The first one relates to the gas flow rate. This
is particularly true for the CO₂ plasma at low flow rates
between 10 and 20 SLPM where the plasma is flickering
and tends to extinguish randomly. The second reason is that
a minimum applied microwave power is required to sustain
a stable plasma. It was experimentally determined that a
microwave power of 2.5 kW and 1 kW were necessary to
sustain a pure H₂O and CO₂ plasma, respectively. Below
these limits, the plasma plume tends to flicker and randomly extinguishes. No plasma could be formed at a
power below 1.75 kW and 0.75 kW for pure H₂O and CO₂
respectively. H₂O requires higher microwave power to
stabilise a plasma and this can be attributed to its higher
heat capacity compared to CO₂. In fact, more energy is
required for the H₂O to reach a certain temperature before
the ionization occurs. The third parameter limiting the
range of steady plasma operation is the life of the quartz
tube. This is especially an issue for the operation of H₂O
plasma. In fact, at low flow rates and high microwave
powers, the quartz tube tends to start melting. In Figures 3
and 4, “unsteady” plasma-operating conditions caused by
plasma flickering, either due to low gas flow rate or low
microwave power, are represented by black points,
whereas “unsteady” plasma operation conditions caused by
melting of the tube are represented by orange points. The
detection of when the quartz tube start melting is verified
using OES measurements. It was experimentally
determined that the first sign of quartz melting is the
emission of the Lithium peak at 671 nm in the plasma that
can also be visualized through glowing red points in the
quartz tube. Lithium is one of the common impurities in
quartz tube and exhibits a strong emission peak at 671 nm
when heated. This is the main issue encountered at low
H₂O flow rates. For microwave powers above 4 kW, it is
necessary to gradually increase the H₂O flow rate when
increasing the microwave power in order to protect the
quartz tube. On the other hand, high working gas flow rate
operation is possible and no higher limitations were found
in the working range of the experimental setup (maximum
flow of 100 SLPM for CO₂ and 50 g/min for H₂O). Operation of the MPT using mixtures of CO₂ and H₂O has
also been tested by the authors and is possible over a wide
range of operating conditions, but these results are not
presented here. Generally, steady plasma operation is
possible when adding CO₂ to stable pure H₂O plasma or
when adding H₂O to a steady pure CO₂ operating plasma.

Temperature measurements
Gasification is a thermal process and hence it is critical
to maintain a sufficient temperature within the gasification
chamber to enable reactions to proceed. In order to
investigate the suitability of the MPT for its integration in
an ATT system, the achievable temperature maxima
within the chamber were recorded for CO₂ and H₂O
plasmas, varying flow rates and magnetron powers across the steady operating conditions presented in section 3.1.

Figure 5: Temperature recorded in chamber with 20 SLPM CO₂ plasma varying the applied microwave power

Figure 5 presents the temperatures recorded when operating a CO₂ plasma across magnetron powers ranging from 1 to 6 kW at a working gas flow rate of 20 SLPM, whilst Figure 6 presents similar data whereby the working gas flow rate has been increased to 40 SLPM. For a CO₂ plasma, temperatures of between 50 and 800°C have been recorded at a working gas flow rate of 20 SLPM whereas temperatures of between 150 and 850°C were obtained at higher flow rate of 40 SLPM. The temperature is clearly increasing when increasing the applied microwave power, especially in the middle of the chamber. At microwave powers of >2 kW, the highest operating temperatures were achieved in the middle of the chamber, whilst at microwave powers of <2 kW the highest temperatures were recorded at the top of the chamber. It is important to remind the reader that the top thermocouple is located horizontally next to the tip of the quartz tube and is therefore not directly exposed to the plasma plume. As such, the top thermocouple provides a temperature estimation of the gas leaving the chamber. As presented in Figure 6, an increase CO₂ flow rate results in a slightly lower average temperature at the top and bottom of the chamber that can be explained by a heat dilution effect. On the other hand, an increase CO₂ flow rate also results in an increase temperature of the middle thermocouple, from 800°C at 20 SLPM to 850°C at 40 SLPM and both at 6 kW microwave power, suggesting an extension of the plasma plume closer to the middle thermocouple. Through comparison of Figure 5 and 6, it is evident that the microwave power has a greater effect on the temperature generated within the chamber than the working gas flow rate.

Figure 6: Temperature recorded in chamber with 40 SLPM CO₂ plasma varying the applied microwave power

Figure 7: Temperature recorded in chamber with 50 g/min H₂O plasma at applied microwave power of 3 and 6 kW

Figure 7 presents the temperatures recorded in the chamber with 50 g/min H₂O plasma for microwave powers of 3 and 6 kW. The temperature measurements with H₂O plasmas are only presented for working gas flow rates of 50 g/min as it was empirically determined that the effect of the flow rate was very low, with temperatures achieved at a flow rate of 25 g/min being similar to those achieved at 50 g/min. At microwave powers of 3 and 6 kW, the maximum chamber temperatures achieved were 600°C and 800°C respectively. As such, the temperature maxima achieved for H₂O plasmas are very similar to those achieved with CO₂. However, the temperature difference within the chamber is lower for H₂O than for CO₂. This can be attributed to the higher gas flow rates required to operate H₂O plasmas, making more homogeneous temperatures explained by greater turbulence within the reactor chamber.
3.3. Plasma chemistry

One of the advantages of plasma is its high chemical activity that will enhance gasification reactions rates avoiding the need for a catalyst [9]. In this study, an insight into the CO$_2$ and H$_2$O plasma chemistry is given using optical emission spectroscopy (OES). OES measurements of pure CO$_2$ and pure H$_2$O plasma are presented, providing information on the chemical species present in the plasma as well as information on the plasma temperatures. The aim of this study was to elucidate the active species present along the plasma plume.

Figure 8: CO$_2$ plasma spectra at 20 SLPM and 3 kW microwave power along the plasma

Figure 9: CO$_2$ plasma spectra varying the gas flow rate and microwave power 8 cm downstream of plasma heart

Figure 8 presents the emissions observed along a CO$_2$ plasma generated using a working gas flow rate of 20 SLPM at a microwave power of 3 kW. The C$_2$ Swan band is the dominant emissions from the CO$_2$ plasma, corresponding to the wavelength range of 400 to 600 nm. This wavelength range represents blue light in the visible light wavelength spectrum explaining the blue colour of the CO$_2$ plasma plume as presented in Figure 2. The O peaks are also present in the infrared wavelength range at 777 and 844 nm whereas the C peak is observed in the ultraviolet range at 248 nm. The presence of the C$_2$, C and O emissions illustrates the dissociation of CO$_2$ in the plasma. In fact, in the high temperature plasma, CO$_2$ tends to dissociate through the following reaction: CO$_2$ $\rightarrow$ CO + O but then recombines to CO$_2$ when the temperature decreases [23, 26-29]. The dissociation and recombination mechanisms are complex and involve many reactions with the dominant species being C, O, CO, C$_2$, O$_2$ and CO$_2$. Details of those mechanisms can be found in the literature [26-29].

According to Figure 8, it is evident that the intensity of the emissions are decreasing along the length of the plasma corresponding to lower temperature and species concentration. However, the C$_2$ Swan band and the O peaks can always be identified in CO$_2$ plasmas even 14 cm down from the centre of the waveguide in any operating conditions, whereas the C peak tends to disappear at lower plasma intensities. Figure 9 presents the influence of the microwave power and the CO$_2$ flow rate on the plasma emissions. It is clear that an increase of the microwave power and the CO$_2$ flow rate results in higher emissions intensities. However, a variation in the applied microwave power has more effect on the plasma spectra than varying the working gas flow rate. CO$_2$ plasma is very bright and the high emission intensities result in spectrometer saturation at approximately 15,000 counts/ms as observed in Figures 8 and 8 respectively. Spectrometer saturation generally occurs close to the CO$_2$ plasma heart and up to 8 cm down the plasma plume as can be seen in Figure 9.
Figure 10 presents the typical OES measurements of a pure H₂O plasma. An increase in microwave power results in higher H₂O emission intensities, reflecting the higher temperatures and concentrations of chemical species. However, this study served to identify that for H₂O plasmas the working gas flow rate has a negligible impact on the emission intensities. The H₂O plasma spectra is marked by the presence of mainly three species, which are •OH, •H and •O. The OH ion emits in the UV range between 275 and 340 nm with its main peak being at 309 nm. Emissions from hydrogen atoms are identified at 656 and 486 nm, respectively corresponding to the Hα and Hβ transitions. The OH ion excited to the level n=3 tends to come back to the less excited state n=2 with the emission of a photon at the wavelength 656 nm, corresponding to the Hα transition. Whereas the Hβ is the transition between the higher excited state of n=4 to n=2. The Hα and Hβ transitions are the only ones identified in experimental results from the Balmer series. The Hα is the dominant emission, also responsible for the orange-red colour of the H₂O plasma, whereas Hβ has a lower intensity and tends to disappear with decreasing the forward microwave power or as you transition along the plasma plume. The •O peaks at 777 and 844 nm are also present, similar to that of the CO₂ plasma. The presence of the •OH, •H and •O peaks is indicative of the dissociation of H₂O in plasma. In fact, similarly to CO₂, H₂O dissociates at high plasma temperatures predominately through the reaction: 

$$
\text{H}_2\text{O} \rightarrow \text{OH} + \text{H}
$$

prior to recombining as temperatures decrease [30]. The H₂O dissociation and recombination mechanisms are complex and mainly involve the dominant species •H, •OH, H₂, O₂ and H₂O [30].

The comparison of the •OH emissions with simulated spectra using the software Specair was carried out in order to estimate the plasma temperature, along the plasma plume transect, at varying operating conditions. The temperature estimations calculated by Specair are based upon rotational temperatures of the identified molecules. Importantly, for this study, it has previously been determined that CO₂ and H₂O microwave plasmas are in local thermal equilibrium, meaning that the gas temperature is estimated to be equal to that of species rotational temperatures [23, 26-28, 30]. Figure 12 presents an example of an experimental spectrum fit using the Specair software. This was repeated for H₂O and CO₂ (1 g/min H₂O was added in CO₂ to study the •OH emissions) plasmas along the quartz tube, at vary applied microwave powers and working gas flow rates. This data indicated that the temperature in pure H₂O plasma varies from 5,800°C in the heart of the plasma to 2,300°C 14 cm down the quartz tube. In contrast, higher temperatures were indicated for CO₂ with maximum observed temperature estimated to be 6,300°C. This study also served to identify that the temperature of the plasma plume could be varied by up to 500°C by varying the forward microwave power and the working gas flow rate. Furthermore, by increasing the microwave power or by decreasing the working gas flow rate, the diameter of the plasma plume is increased, reflecting an increase in energy density, favouring additional molecular ionization, rather than an increase of the plasma temperature. These results demonstrate that very active ions (•OH, •O, •H) are still present at relatively high temperatures in the quartz tube up to 14 cm down the centre of the waveguide.
4. Discussion

This section provides a discussion based on the experimental results presented in section 3, with emphasis on the application of the MPT into a gasification system for waste or biomass conversion.

The operating temperatures in gasifiers generally vary between 550°C and 1,600°C according to the reactor design, the working gas and the feedstock characteristics [5]. Temperatures >600°C are typically required for methane and hydrocarbon decomposition into CO and H2, the main components of syngas [5]. However, in this study, the maximum temperature recorded at the bottom of the connected chamber was 600°C at a maximum operational microwave power of 6 kW. Given that extremely high operating temperatures of >2,000°C have been observed in the MPT plume, the significantly lower temperatures measured in the bottom of the chamber illustrates the requirement for compact plasma gasification reactor integration to avoid steep thermal gradients across the system. The injection of air or oxygen into a gasification vessel could also be used to increase the reaction temperature through exothermic oxidation reactions. However, the use of air or O2 for this process would result in a reduced syngas calorific value due to fuel oxidation and N2 dilution. The study serves to demonstrate how the manipulation of MPT operating parameters could be used to rapidly regulate the temperature within a gasification reactor, which in turn can be used to maximise the efficiency of the gasification process. Whereas the applied microwave power is directly proportional to the temperature in the chamber, the flow of working gas can be tuned to optimise the quantity of reactant required for feedstock conversion.

One of the main advantages of using plasma in ATT processes is the quality of the syngas generated, reducing the requirement for clean-up prior to its use in power generation [6]. In fact, this study highlights plasma temperatures ranging from 2,000°C to 6,300°C within the first 14 cm of the plasma plume, which is appropriate for tars dissociation which occurs at temperatures of >1,000°C [5]. In addition, the presence of ions, electrons, excited molecules and photons within plasma should also serve to enhance the chemical reactions, including gasification reactions as well as tars conversion [8]. The very active •O and •OH identified in plasma will take part in chemical reactions thus improving solid conversion and syngas reforming. Furthermore, the consumption of •O and •OH in gasification reactions will prevent CO2 and H2O recombination, resulting in the generation of additional CO and H2, which in turn enhances the quality of the resultant syngas. The benefit of H2O and CO2 plasma chemistry is therefore twofold as it a) provides very chemically active species enhancing gasification reactions and b) generates additional CO and H2 from CO2 and H2O dissociation respectively. As such, these processes are in theory able to enhance the energy in the syngas to exceed that of the initial energy content of the solid feedstock, assuming complete conversion and additional CO or H2 generated from the plasma working gas dissociation.

Nevertheless, the exceptional properties of plasma can only be efficiently exploited if the solid and the gas resulting from its devolatilization have sufficient time in contact with plasma. In fact, studies integrating MPT in gasification reactors at laboratory scale generally operate through the injection of powdered feedstock within the plasma plume [21, 22, 24, 31] or from the top of the quartz tube [20]. These plasma entrained-flow bed configurations result in low feedstock conversion efficiency, because of the short feedstock residence time, despite the contact with the plasma at high temperatures. The design of a gasification reactor integrated with the MPT is one of the main challenges to improve the efficiency of the process. The experimental study of the MPT, described here, provides useful parametric data to inform the design of a microwave plasma ATT reactor. The plasma moving-bed reactor such as described in Tang et al. [10] is an alternative that could improve feedstock conversion efficiencies as well as the syngas quality. The reactor could be connected, effectively replacing the OES chimney 6 cm down the plasma heart, where the plasma temperature is still >4,000°C. This would enable the solid feedstock to be thermally dissociated at the bottom of the chamber whereas the resultant syngas would go through the high temperature plasma to exit at the top of the reactor, enhancing gas reforming and tars dissociation. Further gas reforming could be achieved by recycling part of the generated syngas in the plasma working gas, which has been proved very efficient for tars reforming [32].

Whereas plasma properties are ideal for gasification applications, the main drawback is the energy consumption of the plasma torch that tends to lower the overall efficiency of the process. The gasification efficiency is generally estimated by the cold gas efficiency, which is defined by the energy in the syngas generated divided by the energy content of the feedstock. In case of allothermal gasification using a plasma torch, the energy consumption of the plasma torch is added to the feedstock energy in the denominator. Conventional authothermal gasification efficiencies are generally circa 80% [33]. Considering a 90% feedstock conversion when using plasma (the energy in syngas is then assumed equal to 90% of the energy content of the feedstock), the energy consumed by the plasma torch should then be less than 12.5% of the energy content of the feedstock in order to achieve similar cold gas efficiencies as conventional gasification. For example, a 5 kW plasma torch should be able to treat at least 40 kW of feedstock, corresponding to a feed rate of 9.6 kg/hr of woody biomass or 4.2 kg/hr of plastic (assuming woody biomass and plastic heating values of 15 and 34 MJ/kg, respectively [33]). With respect to power generation through an integrated gasification power plant, the clean, low tar syngas generated when using plasmas can enable
higher net electrical efficiencies to be achieved, compared to that of autothermal gasification [6]. This observation is further reinforced by the emergence of SOFC systems, that enable greater efficiencies to be achieved by converting syngas to electricity through electrochemical reactions [12].

5. Conclusion

Waste management and energy generation are two major challenges of the 21st century for the mitigation of climate change. This study investigates the feasibility of incorporating a MPT in a gasification system to create an innovative energy from waste or biomass solution. The MPT can be operated in a wide range of operating conditions, including flow rates ranging 20 to 80 SLPM and forward microwave powers of between 1 to 6 kW for pure CO₂, and flow rates of 25 to 50 g/min and forward microwave powers of 2.5 to 6 kW for pure H₂O. Experimental studies have served to prove the flexibility of the device, with respect to operating parameters and temperature control, which is a great advantage for its integration in a gasification system. Temperatures up to 850°C have been measured in a large chamber connected to the plasma torch providing valuable information for the design of an integrated gasification system and showing the benefit of a compact reactor. An insight into the plasma chemistry has been obtained through OES measurements showing the dissociation of H₂O and CO₂ molecules at high temperature up to 6,300°C. Very reactive species such as •O and •OH are formed and are present in the plasma plume even 14 cm downstream of the plasma heart. Those species will enhance gasification reactions whereas additional CO and H₂ will enrich the syngas from CO₂ and H₂O dissociation respectively. The MPT provides an ideal environment for gasification with high temperature and very reactive chemical species. The integration of microwave plasma in gasification has promising potential for the generation of high-quality syngas that lowers the requirement for gas cleaning and eases the coupling of high efficiency energy conversion devices such as SOFC.

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7. References


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