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Influence of steam on gasification of millimetric wood particles in a drop tube reactor: Experiments and modelling

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Keywords:
Biomass
Steam
Gasification
Drop tube reactor
Modelling

HIGHLIGHTS
- Wood particles were gasified in a drop tube reactor at 1000–1400 °C.
- The influence of the presence of steam in the atmosphere was investigated.
- The particle size – 0.35–0.80 mm – has no influence on experimental products yields.
- After 4.4 s at 1400 °C with steam, the char is totally gasified but soot still remains.
- A comprehensive model was developed, which satisfactorily simulates the experiments.

ABSTRACT
Wood particle conversion in a drop tube reactor was studied between 1000 °C and 1400 °C, both with experiments and simulations using a comprehensive model. Particular attention was paid to the influence of steam on the gasification process, and to the wood particle size. The model included a description of the different phenomena involved in biomass conversion: pyrolysis, gas phase reactions, soot formation and carbonaceous solid gasification. Satisfactory results were obtained in comparison with experiments.

The addition of steam in the atmosphere influenced char, tar, soot and gas yields, especially at 1200 °C and 1400 °C. These changes were linked to char and soot steam gasification, and to gas phase reactions, among which hydrocarbons steam reforming and water–gas shift.

At 1400 °C, with 25 mol% H2O and after a residence time of 4.4 s, the char seems to be completely gasified, while the soot yield still represents about 5 wt% of the initial dry biomass.

Wood particle size in the range 0.35–0.80 mm was experimentally shown to have no influence on products yields for a residence time of a few seconds.

1. Introduction

Entrained flow reactor is one of the most promising biomass gasification technologies for biofuel production. Even if this technology is well advanced on an industrial scale for coal gasification, the development of reactors fed with biomass has not really been achieved yet. The main advantage of this technology is the high conversion of biomass into a syngas with very low tar and hydrogen contents. The entrained flow reactor (EFR) is characterised by high operating temperatures, above 1300 °C. The particles are introduced under a pulsed form and, submitted to a high heating flux, are converted into syngas in a short residence time of a few seconds. On the basis of the experience feedback on coal injection and gasification in EFR, particles size is recommended to be under 200 μm. This implies high additional costs in the process due to wood grinding and/or pre-treatment such as torrefaction in order to transform biomass into a material which can be more easily ground [1]. So the use of larger particles, of approximately 1 mm, would be of interest, but it has to be proved that such large particles can be completely converted in the EFR [2].

The phenomenology linked to biomass conversion in conditions similar to those of an EFR has rarely been studied until now. Under a high heating flux, biomass particles, as they enter the reactor, are submitted to fast pyrolysis which mainly produces H2, H2O, CO, CO2, light and heavy hydrocarbons (tar) and char. Hydrocarbons then undergo complex reactions of cracking and polymerization, which can lead to soot particles formation. These carbon-rich particles can induce some operational problems in the process, such as blocking or fouling of equipments, and must be removed from the gas. In the presence of an oxidising gas (steam, oxygen, carbon dioxide), hydrocarbons can be reformed, and the carbonaceous solids can be gasified.
The drop tube reactor (DTR) is an experimental apparatus well adapted to study biomass conversion in the conditions of an EFR at the laboratory scale. Even if the phenomena listed above cannot be completely decoupled, a variation in the composition of the atmosphere can allow isolation of some of them: pyrolysis and hydrocarbon cracking/polymerization reactions are the main phenomena occurring in an inert atmosphere, while hydrocarbon reforming and solid gasification also play an important role in a steam containing atmosphere.

Few works in the literature deal with biomass gasification studies in a DTR at temperatures above 1000 °C, either in an inert atmosphere [3–5], or an atmosphere containing steam and/or oxygen [5–8]. These works show that the gas yield from fast pyrolysis is higher than 80 wt% of dry biomass for particles smaller than 0.5 mm above 800 °C [4,9]. The main gaseous products are CO, CO2, H2, H2O, CH4, and also in smaller amounts C2H2, C2H4, C2H6 and C3H8; gas composition varies with temperature. Tar, char and soot are the other pyrolysis products. Soot is already detected at 900 °C, but its yield is the highest around 1100–1200 °C, where it can reach about 15 wt% of dry biomass in an inert atmosphere [4,5]. The char yield was shown to be lower at 800 °C or above with the addition of steam in the atmosphere [5,7]. The same influence was shown for soot yield [5,8].

In order to better understand the different phenomena which control biomass conversion, a comprehensive modelling tool can be useful. This model should include a representation of biomass pyrolysis, of carbonaceous solid gasification, of gas phase reactions and if possible of soot formation. Few researchers have developed models with such characteristics for biomass gasification in an EPR or DTR. Vilas et al. focused on the biomass reburning phenomenon [10]; the CMIC department at Politecnico di Milano also developed a modelling tool including a detailed description of biomass particles pyrolysis [11–13].

The present study aims to investigate wood particle conversion in a drop tube reactor between 1000 °C and 1400 °C, both with experiments and simulations. Particular attention is paid to the influence of steam on the gasification process, and to wood particle size, which has very rarely been studied in these conditions [3]. The experiments were conducted in a DTR with beech wood particles of 0.35 mm and 0.80 mm as feedstock. First results obtained in an inert atmosphere were already presented in another publication [5,6]. The micro-gas chromatograph (µ-GC) concentrations were considered for quantification of the major gases (CO2, CO, CH4, N2, H2), as well as benzene. For quantification of C2H2, C2H4, C2H6, the FTIR (Fourier Transformed InfraRed) spectrometer value was considered. A Flame Ionization Detector (FID) also gives the total hydrocarbons by difference.

### 2. Experimental

#### 2.1. Biomass samples

The feedstock used in this study is beech sawdust. Two particles sizes ranges were selected after sieving: 0.313–0.400 mm and 0.730–0.900 mm. These samples are referred to as 0.35 mm and 0.80 mm particles samples respectively.

The proximate analysis (ash, moisture, volatile matter and fixed carbon) and the ultimate analysis (C, H, O, N, S) of the two samples are shown in Table 1. No significant difference between the compositions of the two samples is confirmed.

#### 2.2. Description of the drop tube reactor

The drop tube reactor (DTR) is represented in Fig. 1. It consists of an alumina tube inserted in a vertical electrical heater with three independent heating zones. The dimensions of the tube are 2.3 m in length and 0.075 m in internal diameter. The heated zone is 1.2 m long. The DTR works at atmospheric pressure and can reach a maximum temperature of 1600 °C.

The wood particles are continuously fed into the reactor using a gravimetric feeding system, which consists of a feedstock hopper on a conveyor belt. The whole system is placed on a balance and is connected to a computer. The conveyor belt velocity is controlled so as to follow the set point of the feeding flow rate. The wood particles initially introduced in the hopper are entrained by the conveyor belt to a vibrating channel and then to a pneumatic ejector from which they are injected into the reactor with a 2 NL min−1 transport nitrogen stream through a water-cooled (30 °C) feeding probe. A dispersion dome is placed at the outlet of the feeding probe to distribute the solid particles over the reactor cross section.

The main gas stream, which can be N2 or a mixture of N2 and H2O, is electrically pre-heated before entering the reactor. For the introduction of H2O into the DTR, a steam generator working at 180 °C is linked to the reactor inlet.

An oil-cooled (110 °C) sampling probe can be inserted at different heights in the bottom half of the reactor to collect gas and the remaining solid. A fraction of the exhaust gas is sucked in the sampling probe and passes through a settling box and a filter. This part of the experimental facility is heated (150 °C) to avoid steam condensation. After the filter, the sampled gas passes through a condenser if steam is introduced into the reactor, and is finally analyzed.

Different analyzers are available for gas analyses, and have been described in details in a previous article [14]. The micro-gas chromatograph (µ-GC) concentrations were considered for quantification of the major gases (CO2, CO, CH4, N2, H2), as well as benzene. For quantification of C2H2, C2H4, C2H6, the FTIR (Fourier Transformed InfraRed) spectrometer value was considered. A Flame Ionization Detector (FID) also gives the total hydrocarbons content of the gas. H2O was measured by a psychrometer in the case of experiments under nitrogen. For experiments in a wet atmosphere, the water yield was not determined. Indeed, water condensation at the outlet of the reactor (in the condenser) appeared to be not efficient enough for a precise quantification by weighing.

#### 2.3. Conditions of the experiments

Experiments were performed in a nitrogen inert atmosphere and a steam containing atmosphere (25 mol% H2O–75 mol% N2), at several temperatures, 1000 °C, 1200 °C and 1400 °C, and with each of the two particle size samples (Table 1). Gas sampling was performed at two different levels, after 0.6 m – middle of the heated zone – or 1.2 m of reaction length. Only at 1400 °C no

### Table 1: Proximate and ultimate analyses of beech sawdust samples.

<table>
<thead>
<tr>
<th>Proximate analysis</th>
<th>0.35 mm particles</th>
<th>0.80 mm particles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (wt%)</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>Volatile matter (wt% dry basis)</td>
<td>85.3</td>
<td>85.3</td>
</tr>
<tr>
<td>Fixed carbon (wt% dry basis)</td>
<td>14.3</td>
<td>14.3</td>
</tr>
<tr>
<td>Ash (wt% dry basis)</td>
<td>0.4</td>
<td>0.4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ultimate analysis (wt% dry, ash free basis)</th>
<th>0.35 mm particles</th>
<th>0.80 mm particles</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>50.8</td>
<td>50.4</td>
</tr>
<tr>
<td>H</td>
<td>5.9</td>
<td>5.9</td>
</tr>
<tr>
<td>N</td>
<td>0.3</td>
<td>0.3</td>
</tr>
<tr>
<td>S</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>O</td>
<td>42.9</td>
<td>43.3</td>
</tr>
</tbody>
</table>

*By difference.*
experiment could be performed with sampling at the middle of the heated zone because of cooling oil overheating.

The atmosphere gas flow rate at the inlet was varied in order to keep a constant gas velocity and thus a constant gas residence time for experiments at different temperatures (Table 2). Due to slip velocity, the residence time of the 0.80 mm particles was lower than the residence time of the 0.35 mm particles, which was shown to be roughly equal to the gas residence time [14]. Note that the experiments were performed under diluted condition, as the total gas flow rate was hardly modified by gas release from biomass thermal decomposition.

Biomass feeding rate was varied between 0.3 g min⁻¹ and 1.3 g min⁻¹. In this range, no influence of feeding rate on products yields was observed.

<table>
<thead>
<tr>
<th>Temperature of the experiment (°C)</th>
<th>Inlet gas flow rate (NL min⁻¹)</th>
<th>Reaction length (m)</th>
<th>Gas residence time (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1000</td>
<td>15.9</td>
<td>0.6</td>
<td>2.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.2</td>
<td>4.4</td>
</tr>
<tr>
<td>1200</td>
<td>13.8</td>
<td>0.6</td>
<td>2.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.2</td>
<td>4.4</td>
</tr>
<tr>
<td>1400</td>
<td>12.1</td>
<td>1.2</td>
<td>4.4</td>
</tr>
</tbody>
</table>

Our DTR is designed for small biomass feeding rates and for working in diluted conditions. So it was not possible, for tests in a steam containing atmosphere, to find operating conditions which would give representative values of steam/biomass inlet rates ratio and atmospheric steam partial pressure of an industrial EFR. Steam was in a large stoichiometric excess, about 30–35 times higher than the quantity needed for the gasification of all the carbon introduced into the reactor. However, steam partial pressure was fixed at 25 mol%, a value close to the one encountered in an EFR.

In the experiments, different kinds of solids – char and soot – were recovered in the sampling line. Char was mainly found in the settling box, whereas soot particles were mainly observed in the filter.

2.4. Gas and solids yields determination

The gas yields were determined from the gas composition measurements and the nitrogen inlet flow rate, using nitrogen as a tracer. An estimation of the relative error based on repeatability experiments gives ±5% for H₂ and CO yields (g/g of dry biomass), ±10% for CO₂, H₂O and CH₄ yields, and ±20% for C₂H₂, C₂H₄, C₂H₆ and C₆H₆ yields.

Char was recovered into the settling box, and its ash content was measured. As already detailed previously [14], its yield was calculated using the ash tracer method. The relative error for the char yield is estimated at ±25%.
Tar and soot yields could not be directly quantified. In particular, it appeared that soot particles could not be all collected on the filter, as a part of them remained in the sampling probe and could not be quantified. So the lacking mass in the calculated global mass balance for experiments in nitrogen, and in the calculated carbon mass balance for experiments in a wet atmosphere, was attributed to soot and tar. This means that for the latter type of experiments, the hydrogen and oxygen contents of tar and soot were neglected. The soot recovered at the bottom of the DTR was found to be composed of more than 99.2 wt% of carbon [15], which validates this assumption for soot. As for tar, the study of Zhang et al. [5] performed in conditions similar to ours show that for experiments in a steam containing atmosphere, at 1000 °C and above, carbon represents more than 93.5 wt% in tars. Moreover, the same authors show that the tar yield at 1200 °C and above is negligible, which was confirmed in our study, as detailed in Section 4.1. So, our assumption implies that at 1000 °C only in a wet atmosphere, the experimental tar + soot yield could be a bit underestimated, of 6.5% at maximum.

Moreover, the uncertainty derived from repeatability experiments for the tar and soot yield is estimated to be of ±25%. These two sources of errors were taken into account for representation of tar + soot yields (Figs. 2, 6 and 11).

3. Modelling

The experiments were simulated using a numerical model named GASPAR, which allows modelling a drop tube reactor. The GASPAR software, originally developed for thermochemical applications such as combustion and pollutants formation [16–18], was modified in order to adapt it to biomass gasification [19], and specifically in this study for biomass gasification above 1000 °C.

3.1. The GASPAR model

For the experiments described above, the gas flow in the tubular reactor is laminar (Reynolds number between 600 and 1000). The drop tube reactor is modelled by a plug flow reactor. A Lagrangian approach is used to represent the biomass particles, which are here considered to be spherical. Particle isothermicity assumption is made and no slip between the gas and the particles is considered. The entire flow in the drop tube reactor is simulated as the sum of elementary parts, each one representing a single particle and the surrounding gas volume associated to it. The model simulates the time or space evolution of these samples along the reactor, which are equivalent here. The model can be therefore qualified as a 0D model. As a consequence, it does not allow to completely characterize the reactor temperature and hydraulics, contrary to CFD simulations used by other authors in complement to biomass pyrolysis experiments in drop tube reactors [20,21].

Gas phase reactions are computed using subroutines from CHEMKIN II software using a detailed chemical mechanism [22]. The model includes thermal and chemical phenomena:

- heat transfer of the carrier gas (convection and radiation) and of wood particles (external conduction and radiation), described in more details in [16];
- particle drying using an Arrhenius law;
- chemical reactions: particle pyrolysis, gas phase reactions including soot formation, soot and char gasification.

The gas temperature profile calculated along the drop tube reactor was validated against measurements performed at several heights in the reactor.

All the differential equations are simultaneously time integrated with the help of the Gear algorithm that can solve stiff ordinary differential equation systems [23]. The whole model is included in a FORTRAN program. The model is able to predict with a very low running time (under 20 s) the evolution of several variables versus the gas residence time: char, gas, soot and tar yields, particle temperature, and also gas composition with all the species considered in the detailed kinetic scheme. Note that as slip particle velocity is neglected in these simulations, solid and gas residence times are equal. Therefore, the model results are compared to the experimental results obtained with 0.35 mm wood particles only, for which the slip velocity can be neglected.

Each chemical phenomenon is described below in more details.

3.2. Pyrolysis

The term “pyrolysis” here refers to biomass devolatilization and vapour thermal cracking. Pyrolysis is modelled by a one-step reaction, where biomass is decomposed into volatile species and a carbonaceous residue, the char.

The time evolution of the dry biomass weight \( m_{\text{dry biomass}} \) is expressed by the following relation:

\[
\frac{dm_{\text{dry biomass}}}{dt} = -m_{\text{dry biomass}} \cdot k_{\text{pyro}} (s^{-1})
\]

(1)

\( k_{\text{pyro}} \) (s\(^{-1}\)) is the pyrolysis kinetic parameter, and is expressed by means of an Arrhenius law:

\[
k_{\text{pyro}} = k_{0,\text{pyro}} \exp \left( \frac{-E_{\text{a,pyro}}}{RT} \right)
\]

(2)

with \( k_{0,\text{pyro}} = 7.4 \times 10^{5} \) s\(^{-1}\) and \( E_{\text{a,pyro}} = 70 \times 10^{3} \) J mol\(^{-1}\). These values were adjusted on experimental results obtained for biomass pyrolysis at 800 °C in the same reactor [9,19].

The products are gaseous species (\( \text{H}_2, \text{CO}, \text{H}_2\text{O}, \text{CO}_2, \text{CH}_4, \text{C}_2\text{H}_6, \text{C}_3\text{H}_8, \text{C}_6\text{H}_{12} \)), char, and tar compounds, for which model species were searched. For experiments in an inert atmosphere at 800 °C in the same DTR, tar was shown to have a mean formula close to \( \text{C}_{1.4}\text{H}_{2.2} \text{O}_{0.3} \) [15]; this suggests that tar is a mixture of oxygen containing and aromatic hydrocarbons. This is in agreement with the experimental results of Zhang et al. [4] obtained under similar conditions, who found out that methanol, phenol, toluene and naphthalene were the main tar species at 800 °C. Among these species, phenol was not selected as its decomposition was not correctly represented in the kinetic scheme selected for gas phase reactions modelling (Section 3.3). It appeared then that a combination between methanol – \( \text{CH}_3\text{OH} \) – and naphthalene – \( \text{C}_{10}\text{H}_{8} \) – only was the best representation for our tar composition. These two model compounds were introduced in the gas phase and their reactions were modelled with the same detailed kinetic scheme as the one used for all other gaseous compounds. Char is supposed to be composed of carbon only.

So, biomass pyrolysis is altogether represented by the reaction:

\[
\begin{align*}
\text{C}_w\text{H}_4\text{O}_y + z\text{H}_2\text{O} & \rightarrow a\text{H}_2 + b\text{CO} + c\text{CO}_2 + d\text{H}_2\text{O} + e\text{CH}_4 \\
+ f\text{C}_2\text{H}_2 + g\text{C}_2\text{H}_4 + h\text{C}_2\text{H}_6 + i\text{C}_6\text{H}_6 + j\text{C}_6\text{H}_5 \\
+ k\text{C} + m\text{CH}_3\text{OH} + n\text{C}_{10}\text{H}_8 + o\text{CH}_4\text{O}_x
\end{align*}
\]

(3)

where the dry biomass \( \text{C}_w\text{H}_4\text{O}_y \) has been individualized from its moisture. The composition of the biomass is indicated in Table 3. A “rest” with the mean formula \( \text{C}_{1.4}\text{H}_{2.2} \text{O}_{0.3} \) was introduced into the reaction so that the stoichiometry is respected. Physically, the mass fraction of this rest can be attributed to tar. In the model, as this rest does not correspond to any identified molecule, the choice was made that it would not participate in any other reaction.

The coefficients \( a, b, c, d, e, f, g, h, i, j, k \) and \( o \) were determined from previous experiments under an inert atmosphere at 800 °C.
The products in these conditions are assumed to mainly result from pyrolysis and tar cracking, which are supposed in the model to give the same products at any temperature higher than 800 °C. This means in particular that the char yield from the pyrolysis step is assumed to be independent on temperature between 800 and 1400 °C. This hypothesis lies on several experimental observations: in our previous experiments in an inert atmosphere, we observed that the char yield was steady between 800 and 1200 °C [14]; this is in agreement with the experimental findings of Zhang et al. [4] who observed a plateau in char yield between 800 and 1100 °C for pyrolysis of Hinoki cypress sawdust.

Having made the choice that methanol and naphthalene would represent all tar, the coefficients \( m \) and \( n \) were adjusted in order to minimize the coefficients \( s \), \( r \) and \( u \). The mass of the rest then represents less than 3 wt% of initial dry biomass. A single set of parameters was used for all simulations (Table 4).

### 3.3. Gas phase reactions and soot formation

The Skjæth-Rasmussen and co-workers kinetic scheme [159 species, 773 reactions] [24] was selected for gas phase modelling. This kinetic scheme can predict the Polycyclic Aromatic Hydrocarbons (PAH) formation, which play an important role in soot formation. It was validated with a series of methane oxidation experiments in a plug-flow reactor, some of them in the presence of steam [24]. This mechanism was compared with a good accuracy to gas phase experiments between 1000 °C and 1400 °C on gas mixtures with a composition representative of biomass pyrolysis gas [25].

Soot formation implies several steps [26]:

- Nucleation: the primary soot particles, whose size is under 20 nm, are formed from the inception of two PAH molecules,
- Surface growth, by the addition of \( C_2H_2 \) or of PAH molecules on the primary particles surface,
- Coagulation of primary soot particles.

Surface growth and coagulation lead to the formation of spherical particles of several tens of nm, called spherules. Finally, these spherules agglomerate to form cluster-like or chain-like micrometric structures.

Some models of soot formation represent the elementary soot formation steps in a simplified [27] or detailed [28] way. The existing models can predict soot mass yield, particles number density and even soot composition for the most complex ones. The present study focuses on the modelling of soot mass yield only, which can be compared to experimental results.

In the selected kinetic scheme [24], soot formation can be modelled by the collision of two pyrene molecules, which corresponds to the particle inception step. However, this approach did not give satisfactory results in our conditions as soot formation modelling through particle inception only is not complete. In fact, its use would require the introduction of complex parameters for the representation of mass growth in order to obtain more accurate results, while a simple approach is suitable in the present study. So, another soot formation model was developed in this work and introduced in the kinetic scheme. In this model, soot is formed from \( C_2H_2 \) and from all PAH, following one-step independent reactions where soot particles inception and surface growth are implicitly included. Soot is assumed to be composed of carbon only. In our calculations, the main PAH compounds formed by gas phase reactions and considered as soot precursors are: naphthalene \((C_{10}H_8)\), acenaphthalene \((C_{12}H_{10})\), phenanthrene \((C_{14}H_{10})\), fluoranthene \((C_{16}H_{10})\), and pyrene \((C_{16}H_{10})\).

The reactions leading to soot formation are written below, where PAHs are represented by the general formula \( C_{n}H_{m} \):

\[ C_2H_2 \rightarrow 2C_{soot} + H_2 \]  
\[ C_{n}H_{m} \rightarrow nC_{soot} + \frac{m}{2}H_2 \]

Each reaction kinetic is supposed to follow a first order Arrhenius law with respect to \( C_2H_2 \) or PAH concentration. The same activation energy as that obtained by Ziegler [27] for soot formation from \( C_2H_2 \) in propane pyrolysis experiments \((E_a = 167 \cdot 10^{-3} \text{ J mol}^{-1})\) was used for reactions (4) and (5). As shown before, \( C_2H_2 \) and PAH participate in soot formation following different mechanisms, thereby a different pre-exponential factor was used for reactions (4) and (5). These factors were adjusted in order to minimize the difference between experimental and calculated \( C_2H_2 \) and tar + soot yields in an inert atmosphere at 1000 °C and 1200 °C.

The values of the pre-exponential factors obtained by this fitting are \(1.0 \times 10^6 \text{ s}^{-1} \) and \( 5 \times 10^5 \text{ s}^{-1} \) for reactions (4) and (5) respectively.

### 3.4. Char and soot particles gasification

According to a previous analysis based on characteristic times comparison, char particles gasification is controlled by chemical reaction only and not by steam mass transfer in the conditions of our experiments in a wet atmosphere with the 0.35 mm wood particles [15]. The same result was obtained for soot particles steam gasification.

Kinetic models previously developed on the basis of thermogravimetric analysis experiments with soot and char samples recovered in the drop tube reactor facility were introduced into GASPAR. The kinetic laws taken from [15] are given in Table 5 as a function of the conversion \( X \), which is calculated as:

### Table 4

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Values of the parameters fixed in reaction (3).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coefficients of the pyrolysis reaction (3)</td>
<td></td>
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<tr>
<td>( a )</td>
<td>0.6999</td>
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<tr>
<td>( b )</td>
<td>2.0156</td>
</tr>
<tr>
<td>( c )</td>
<td>0.3527</td>
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<td>( d )</td>
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<td>( e )</td>
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<tr>
<td>( n )</td>
<td>0.1445</td>
</tr>
</tbody>
</table>

Formula for “rest”

\[ s \]
\[ t \]
\[ u \]

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Dry biomass formula and moisture coefficient.</th>
</tr>
</thead>
<tbody>
<tr>
<td>( w )</td>
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<tr>
<td>( x )</td>
<td>3.7654</td>
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<tr>
<td>( y )</td>
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<table>
<thead>
<tr>
<th>Table 5</th>
<th>Coefficients of the pyrolysis reaction (3).</th>
</tr>
</thead>
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<td>( a )</td>
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<td>( b )</td>
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<tr>
<td>( c )</td>
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</tbody>
</table>
The total dry gas (H\textsubscript{2}, CO, CO\textsubscript{2}, CH\textsubscript{4}, C\textsubscript{2}H\textsubscript{6}, C\textsubscript{3}H\textsubscript{8}, C\textsubscript{4}H\textsubscript{10}) and tar + soot mass yields obtained for 0.35 mm particles are firstly presented in Section 4.1, with a special focus on the influence of steam in the atmosphere. The simulation results are then compared to the experimental ones in Section 4.2, at all temperatures and for both inert and wet atmospheres. The experimental results for 0.80 mm particles are finally presented in Section 4.3 where they are compared to the 0.35 mm particles experimental results.

### 4.1. Influence of steam on the experimental results

The experimental char, tar + soot and gas yields obtained for 0.35 mm wood particles are firstly presented in Section 4.1, with a special focus on the influence of steam in the atmosphere. The simulation results are then compared to the experimental ones in Section 4.2, at all temperatures and for both inert and wet atmospheres. The experimental results for 0.80 mm particles are finally presented in Section 4.3 where they are compared to the 0.35 mm particles experimental results.

#### Fig. 2

![Graph showing mass yields for different conditions](image)

**Fig. 2.** Experimental mass yields of char, tar + soot, and total dry gas under inert and wet atmospheres at 1000°C, 1200°C and 1400°C.

\[ X = 1 - \frac{m(t)}{m(t = 0)} \]

where \( m \) is the mass of char or soot (kg) and \( t \) is the time (s).

Gasification by steam only was considered in the model.

### 4. Results and discussion

The experimental char, tar + soot and gas yields obtained for 0.35 mm wood particles are firstly presented in Section 4.1, with a special focus on the influence of steam in the atmosphere. The simulation results are then compared to the experimental ones in Section 4.2, at all temperatures and for both inert and wet atmospheres. The experimental results for 0.80 mm particles are finally presented in Section 4.3 where they are compared to the 0.35 mm particles experimental results.

#### 4.1. Influence of steam on the experimental results

The total dry gas (H\textsubscript{2}, CO, CO\textsubscript{2}, CH\textsubscript{4}, C\textsubscript{2}H\textsubscript{6}, C\textsubscript{3}H\textsubscript{8}, C\textsubscript{4}H\textsubscript{10}) and tar + soot mass yields obtained for 0.35 mm particles under an inert atmosphere at the bottom of the reactor, already presented in [14], are compared to the yields obtained under a wet atmosphere (25% H\textsubscript{2}O, 75% N\textsubscript{2}) in Fig. 2. Note that the total dry gas mass yield was divided by 10 to be represented on the same scale as char and tar + soot.

As explained before, the sum of tar and soot yields was determined by difference from the mass balance. At 1200°C and 1400°C, the experimental observations showed that almost no tar was present in the DTR. Besides, the sum of CH\textsubscript{4}, C\textsubscript{2}H\textsubscript{6}, C\textsubscript{3}H\textsubscript{8} and C\textsubscript{4}H\textsubscript{10} molar fractions, measured by \( \mu \)-GC and FTIR at the outlet, was equal to the total hydrocarbons content measured by FID. This means that the experimental tar yield at 1200°C and 1400°C is negligible, so the experimental “tar + soot” mass yield at 1200°C and 1400°C is considered to be that of soot.

Fig. 2 shows that the (tar+) soot yield is higher under an inert atmosphere at all temperatures, with a significant difference at 1200°C and 1400°C. Note that the soot yield still represents about 5 wt% of the initial dry biomass in a wet atmosphere at 1400°C. This result shows that even at high temperature with significant steam content in the atmosphere, soot particles cannot be completely avoided. This was also already stated by Qin et al. [8] even with some oxygen added to the inert gas mixture for biomass gasification tests in a DTR.

In both atmospheres, char yield tends to decrease as temperature increases (Fig. 2). At 1000°C, char yield is not significantly influenced by the atmosphere, whereas at higher temperature, char yield is much lower under a wet atmosphere. Considering that char formation by pyrolysis gives the same yield in inert and wet atmospheres, the lower Char yield in a wet atmosphere at 1200°C and 1400°C can be explained by its steam gasification. Note that at 1400°C in a wet atmosphere, no carbonaceous char could be collected at the output of the drop tube reactor. The mass yield represented in Fig. 2 is then that of the ashes.

The total dry gas yield is higher with the presence of steam in the atmosphere, especially at 1200°C and 1400°C (Fig. 2). In these conditions, the total dry gas yield is higher than 100% of initial dry biomass mass, which shows that steam takes part in reactions leading to dry gas formation.

The individual gas mass yields are presented in Figs. 3 and 4 at 1000°C, 1200°C and 1400°C in a wet atmosphere, and compared to the previous results obtained in an inert atmosphere [14]. Note that the H\textsubscript{2} mass yield was multiplied by 10 in Fig. 3 to be represented on the same scale as CO and CO\textsubscript{2}.

In the presence of steam in the atmosphere, the H\textsubscript{2} and CO\textsubscript{2} yields are much higher, whereas the CO yield is steady or lower than in an inert atmosphere (Fig. 3). At 1200°C and 1400°C, C\textsubscript{2}H\textsubscript{6}, C\textsubscript{3}H\textsubscript{8} and C\textsubscript{4}H\textsubscript{10} yields are significantly lower in a wet atmosphere (Fig. 4). The CH\textsubscript{4} yield is surprisingly higher in a wet atmosphere at 1000°C and 1200°C. At 1400°C, no hydrocarbon at all, including CH\textsubscript{4}, can be detected in a wet atmosphere.

As shown in Fig. 2, at 1200°C and 1400°C, the presence of 25% of steam in the atmosphere induces a significantly higher gas yield. A contribution to this higher gas yield may come from char or soot steam gasification, expressed by reaction (7) assuming that char and soot are composed of carbon only:

\[ C + H\textsubscript{2}O \rightarrow CO + H\textsubscript{2} \]  

(7)

However, even by making the gross and probably wrong assumption that the lower soot yield in a wet atmosphere is only due to soot gasification, carbonaceous solids gasification cannot quantitatively explain the difference in dry gas yields between inert and wet atmospheres at 1200°C and 1400°C (Fig. 3). So, some gas phase reactions implying steam also contribute to this higher gas yield.

Hydrocarbon steam reforming may be one explanation to this observation, and also to the lower C\textsubscript{2}H\textsubscript{6}, C\textsubscript{3}H\textsubscript{8} and C\textsubscript{4}H\textsubscript{10} yields in a wet atmosphere (Fig. 3). Hydrocarbon steam reforming, together with steam gasification of soot, were also mentioned in [8] to explain the increase of gas yield with the addition of steam. Concerning the higher CH\textsubscript{4} yield in a wet atmosphere at 1000°C and 1200°C, the same result was also obtained in other works focusing on methane reforming [25,29]. This result could be due to an inhibiting effect of H\textsubscript{2}O on methane reforming at temperatures below 1300°C [29].
ments. In the model, char formation yield is the same whatever spheres. Simulations are globally in good agreement with experi-

DTR, as a function of temperature and for both inert and wet atmos-

tivation, with its equilibrium shifted towards CO

Fig. 3. Experimental mass yields of H\textsubscript{2}, CO, and CO\textsubscript{2} under inert and wet atmospheres at 1000 °C, 1200 °C and 1400 °C.

Fig. 4. Experimental mass yields of hydrocarbons under inert and wet atmospheres at 1000 °C, 1200 °C and 1400 °C.

Fig. 5. Experimental (symbols) and calculated (lines) char mass yields for 0.35 mm particles.}

Finally, we think that the higher CO\textsubscript{2} and H\textsubscript{2} yields correlated with a lower CO yield are largely due to the water–gas shift reac-
tion, with its equilibrium shifted towards CO\textsubscript{2} and H\textsubscript{2} production in presence of a large excess of steam:

\[
\text{CO} + \text{H}_2\text{O} \leftrightarrow \text{CO}_2 + \text{H}_2 \tag{8}
\]

The higher gas yield in a wet atmosphere above mentioned would then be largely due to this reaction.

This discussion shows that the influence of steam on the products yields cannot be both quantitatively and simply explained. In the GASPAR model, the reactions mentioned above are all taken into consideration, even in a more detailed way for gas phase reactions. The comparison between simulation and experiment results in next section aims at verifying if our representation of phenomena in the model is well adapted to the experiments, and at better understanding the results.

4.2. Comparison of experiment and simulation results

Char yields obtained by experiments and by simulations are represented in Fig. 5 for 0.35 mm particles, at the bottom of the DTR, as a function of temperature and for both inert and wet atmospheres. Simulations are globally in good agreement with experiments. In the model, char formation yield is the same whatever the temperature – about 0.05 g/g of dry biomass, calculated from coefficients of Table 4. Lower char yields values are due to char steam gasification, which is described by the kinetic law presented in Table 5. The gasification rate is dependant on the steam partial pressure in the atmosphere. This is why at constant temperature, gasification is enhanced in a wet atmosphere, leading to a lower char yield. In an inert atmosphere, only the steam released by biomass pyrolysis or subsequent gas phase reactions can take part in the char gasification process. Gasification by the steam released during biomass conversion is observed at 1400 °C in an inert atmosphere (Fig. 5), with a char yield at about 0.03 g/g of dry biomass. However, the higher temperature – 1400 °C – and the addition of steam in the atmosphere are both necessary to almost completely gasify the char, as only ashes then remain.

As said before, gasification by CO\textsubscript{2} was not considered in the model. Several experimental studies show that biomass char gasi-

Fig. 3. Char mass yields obtained by experiments and by simulations are

Table 4

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>CH\textsubscript{4}</th>
<th>C\textsubscript{2}H\textsubscript{2}</th>
<th>C\textsubscript{2}H\textsubscript{4}</th>
<th>C\textsubscript{6}H\textsubscript{6}</th>
</tr>
</thead>
<tbody>
<tr>
<td>N\textsubscript{2}, 1000 °C</td>
<td>0.10 ± 0.01</td>
<td>0.01 ± 0.005</td>
<td>0.02 ± 0.005</td>
<td>0.00 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2} + 25% H\textsubscript{2}O, 1000 °C</td>
<td>0.12 ± 0.01</td>
<td>0.015 ± 0.005</td>
<td>0.03 ± 0.005</td>
<td>0.002 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2}, 1200 °C</td>
<td>0.15 ± 0.02</td>
<td>0.02 ± 0.01</td>
<td>0.04 ± 0.01</td>
<td>0.003 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2} + 25% H\textsubscript{2}O, 1200 °C</td>
<td>0.18 ± 0.02</td>
<td>0.025 ± 0.01</td>
<td>0.05 ± 0.01</td>
<td>0.004 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2}, 1400 °C</td>
<td>0.20 ± 0.03</td>
<td>0.03 ± 0.015</td>
<td>0.06 ± 0.015</td>
<td>0.006 ± 0.0015</td>
</tr>
<tr>
<td>N\textsubscript{2} + 25% H\textsubscript{2}O, 1400 °C</td>
<td>0.23 ± 0.035</td>
<td>0.035 ± 0.015</td>
<td>0.07 ± 0.015</td>
<td>0.007 ± 0.0015</td>
</tr>
</tbody>
</table>

Table 5

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>CH\textsubscript{4}</th>
<th>C\textsubscript{2}H\textsubscript{2}</th>
<th>C\textsubscript{2}H\textsubscript{4}</th>
<th>C\textsubscript{6}H\textsubscript{6}</th>
</tr>
</thead>
<tbody>
<tr>
<td>N\textsubscript{2}, 1000 °C</td>
<td>0.10 ± 0.01</td>
<td>0.01 ± 0.005</td>
<td>0.02 ± 0.005</td>
<td>0.00 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2} + 25% H\textsubscript{2}O, 1000 °C</td>
<td>0.12 ± 0.01</td>
<td>0.015 ± 0.005</td>
<td>0.03 ± 0.005</td>
<td>0.002 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2}, 1200 °C</td>
<td>0.15 ± 0.02</td>
<td>0.02 ± 0.01</td>
<td>0.04 ± 0.01</td>
<td>0.003 ± 0.001</td>
</tr>
<tr>
<td>N\textsubscript{2} + 25% H\textsubscript{2}O, 1200 °C</td>
<td>0.18 ± 0.02</td>
<td>0.025 ± 0.01</td>
<td>0.05 ± 0.01</td>
<td>0.004 ± 0.0015</td>
</tr>
<tr>
<td>N\textsubscript{2}, 1400 °C</td>
<td>0.20 ± 0.03</td>
<td>0.03 ± 0.015</td>
<td>0.06 ± 0.015</td>
<td>0.006 ± 0.0015</td>
</tr>
<tr>
<td>N\textsubscript{2} + 25% H\textsubscript{2}O, 1400 °C</td>
<td>0.23 ± 0.035</td>
<td>0.035 ± 0.015</td>
<td>0.07 ± 0.015</td>
<td>0.007 ± 0.0015</td>
</tr>
</tbody>
</table>

Fig. 5. Char mass yields obtained by experiments and by simulations are

For experiments in nitrogen at 1000 and 1200 °C for char, and at all temperatures for soot, the steam gasification is very low so that taking into account gasification by CO\textsubscript{2} would hardly change the results.

For experiments in nitrogen at 1400 °C for char, and at all temperatures for soot, the steam gasification is very low so that taking into account gasification by CO\textsubscript{2} would hardly change the results.
A comparison between experimental and calculated tar + soot mass yields is now performed in Fig. 6a, at different temperatures and for both inert and wet atmospheres. Note that according to the simulations, the tar species at 1000 °C and above are only PAHs. Moreover, the “rest” yield is not included in the calculated tar yield. Tar + soot calculated mass yields at 1000 °C and 1200 °C in an inert atmosphere agree with experimental results, as gasification kinetic law parameters were adjusted for this purpose, as explained in Section 3.3. In an inert atmosphere at 1400 °C, the simulation is still in good agreement with the experiment. In a wet atmosphere, the simulation results are rather satisfactory at 1400 °C, whereas they are too high compared to experimental results at 1000 °C and 1200 °C. So, in these conditions, the influence of steam on tar and soot yields is not quantitatively well represented. This may be explained by an underestimation of tar reforming in the model, which would then lead to CO and H₂ formation. The model seems to wrongly favour reaction paths leading to PAHs polymerization and soot formation.

The individual tar and soot mass yields obtained by simulations are shown in Fig. 6b. The calculated tar yields are close to 0 at 1200 °C and 1400 °C, which is in agreement with the experimental observations mentioned in previous section.

Supplementary simulations were performed at 1400 °C in order to better understand why soot yield is much lower in a wet atmosphere. The difference can a priori be explained by two phenomena both represented in the model: steam reforming of soot precursors, which limit soot formation, and/or steam gasification of soot particles in a H₂O rich atmosphere. Fig. 7 shows the calculated soot yield at 1400 °C:

- in an inert atmosphere,
- in a wet atmosphere without taking soot particles steam gasification into account: the only influence of steam on soot yield then comes from its reaction with soot precursors,
- in a wet atmosphere considering soot particles steam gasification.

Comparing the two first results in Fig. 7 shows that, according to the model, part of soot yield reduction in a wet atmosphere comes from reactions of steam with soot precursors, thus leading to a lower soot formation yield. However, comparing the last two results, it can be seen that soot particles gasification also contributes to the lower soot yield in a wet atmosphere, even if this phenomenon is less important.

The gas yields experimental results are now compared to the simulation results for 0.35 mm particles at 1200 °C in Figs. 8 and 9 for an inert and a wet atmosphere respectively. Results are shown here as a function of the reaction zone length. Note that no model result is shown at 1000 °C, as the same conclusions can be drawn at 1000 °C and 1200 °C. Let us remind that individual gas yields from biomass pyrolysis were adjusted to fit experimental results at 800 °C (Section 3.2); C₂H₂ yield was also adjusted to fit the experimental results at 1200 °C (Section 3.3). So the simulation results shown here at higher temperature depend on gas reaction and solid gasification modelling.

Fig. 8 shows that the simulation results are globally in good agreement with experimental results in an inert atmosphere. The H₂ yield is slightly underestimated, of about 15%, while the CH₄ yield is overestimated by the model.

Fig. 9 shows that at 1200 °C the CO, CH₄ and C₂H₂ yields given by the simulation are in rather good agreement with experimental results in a wet atmosphere. The CO₂ and H₂ yields are underestimated by the model, but their increase along the reaction zone length is qualitatively well represented. When comparing the experimental and calculated tar + soot yields above (Fig. 6a), we made the hypothesis that tar reforming might be underestimated by the model at 1000 °C and 1200 °C in a wet atmosphere. This would be in agreement with an underestimation of gas yield in these conditions, in particular H₂ yield and CO or CO₂ yield, the latter being dependant on the CO yield via the water–gas shift reaction.

Another comparison of the experimental and calculated gas mass yields is shown in Fig. 10 for inert and wet atmospheres at 1400 °C. The agreement between model and experiments is then good, which was also shown before for char and tar + soot yields at the same temperature (Figs. 5 and 6a respectively).

**4.3. Influence of particle size on experimental products yields**

The influence of biomass particle size (0.35 mm and 0.80 mm) on char, tar + soot and total dry gas yields is shown in Fig. 11 for
5. Conclusion

This study allowed the investigation of wood particle conversion in a drop tube reactor between 1000 °C and 1400 °C, both through experiments and simulations. It focused in particular on the influence of steam and of wood particle size on the gasification process.

The addition of steam to the atmosphere was experimentally shown to have an influence on tar, soot, char, and gas yields, especially at 1200 °C and 1400 °C. Due to char steam gasification enhancement in a steam-rich atmosphere, char yield is then lower. Concerning soot yield, from 15 to 20 wt% in an inert atmosphere, it decreases to about 5 wt% in a wet atmosphere. The total dry gas yield is much higher under a wet atmosphere at 1200 °C and 1400 °C, and its composition is modified. This can be partly explained by steam gasification of solids – char and soot – which produce gases. However, gas phase reactions, such as hydrocarbons steam reforming and water gas shift, probably also have an important influence on gas yields.

In the experimental conditions which are the closest to the operation conditions of an EFR − 1400 °C, atmosphere with 25 mol% H2O, residence time of 4.4 s – the char seems to be completely gasified. However, the soot yield still represents about 5 wt% of the initial dry biomass.

Wood particle size in the range 0.35–0.80 mm was experimentally shown to have no influence on products yields for a residence time of a few seconds. This lets us think that particles of about 1 mm could be well gasified in an EFR. The model appeared to give globally satisfactory results in comparison with experimental ones, concerning char, tar + soot and...
individual gas yields. Only the influence of steam at 1000 °C and 1200 °C could still be better represented, maybe by improving the modelling of reactions involved in tar reforming.

The model was used to reach a better comprehension of the involved phenomena. The simulation results showed that the lower soot yield in a wet atmosphere could mainly be explained by reforming of soot precursors, but also by steam gasification of soot particles. In the future, we intend to use the different individual models of GASPAR representing biomass conversion in other numerical models developed for EFR design.

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References