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Influence of H₂O, CO₂ and O₂ addition on biomass gasification in entrained flow reactor conditions: Experiments and modelling

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HIGHLIGHTS

- Wood particles were gasified in CO₂, H₂O and O₂ atmospheres.
- H₂O and CO₂ addition decrease char and soot production above 1200 °C.
- O₂ addition lowers char, hydrocarbons, soot and tar formation.
- Experiments were modelled with a 1D-model using detailed chemical scheme.
- Gas, char, tar + soot were satisfactorily simulated in the whole range of conditions.

Keywords:
Biomass
Gasification
Entrained flow reactor
Drop tube reactor
Modelling

ABSTRACT

Biomass gasification in Entrained Flow Reactor (EFR) is both studied with experiments in a drop tube reactor and modelling with a 1-D model (GASPAR). Operating conditions are chosen thanks to results of a preliminary modelling of an industrial EFR. Influence of addition of steam (0.55 g/g db), carbon dioxide (0.87 g/g db) and oxygen (Equivalent Ratio: 0–0.61) is investigated between 800 and 1400 °C with beech wood particles sieved between 315 and 450 μm as feedstock. The model takes into account pyrolysis reaction, gas phase reaction with a detailed chemical scheme (176 species, 5988 reactions), char gasification by steam and CO₂ and soot formation. H₂O or CO₂ addition has no influence on gasification product yields at 800 and 1000 °C, while at 1200 and 1400 °C the char gasification is significantly enhanced and soot formation is certainly inhibited by OH radical which reacts with soot precursors. The modification of output gas phase composition is mostly due to WGS reaction which reaches thermodynamic equilibrium from about 1200 °C. As expected, O₂ has a significant influence on gas and tar yields through combustion reactions. Char and soot yields decreases as ER increases. The GASPAR model allows a good prediction of gas and char and gives relevant evolution of soot and tar yields on the large majority of conditions studied.

1. Introduction

The world is facing a major energy crisis in the recent years. Fossil fuel energy is the first source of primary energy and we are facing two major issues: the depletion of fossil resources while the world energy demand grows [1], and the climate change, generally attributed to the greenhouse gases emissions directly linked to the fossil fuel usage [2]. To cope up with this challenge, numerous alternative sources of energy are explored. Currently, biomass is the first renewable resource in the world and is well spread across the world. Different biomass conversion processes have been developed and biomass gasification appears to be one of the most promising processes to produce syngas for power generation or biofuel synthesis.

The Entrained Flow Reactor (EFR) is a well-known technology for coal gasification but it still needs development to be operated with biomass as feedstock. The main advantage of this technology is the high conversion of biomass into syngas with a very low tar content which is particularly appropriate for biofuel synthesis processes. The EFR is characterized by a high operating temperature (~1500 °C), a high pressure (>30 bar), a short particle residence time (~5 s) and a high heat flux at particle surface (>1 GW m⁻²). It is generally operated as an autothermal reactor which means that a part of biomass is burnt to supply enough energy for endothermic gasification reactions. Biomass is injected as small particles (smaller than 300 μm).
In order to improve the knowledge on this technology, experimental and modelling studies have been conducted. Experimental ones are generally performed with lab-scale reactors which allow reproducing some important EFR characteristics as temperature, heat flux, residence time and particle size: the Drop Tube Reactor (DTR). Only few studies have dealt with woody biomass pyrolysis in a DTR above 1000 °C [3–6]. Woody biomass gasification has been studied at high temperature (T > 1000 °C) in presence of oxygen [4,7,8] and/or in presence of steam [4,7,9–11] or in presence of carbon dioxide [11]. These studies show a high influence of gasification atmosphere both on carbon conversion and on the product gas composition. Recently, several biomass EFR pilot facilities have been built to study biomass gasification in conditions close to those of an industrial plant [12,13].

In parallel, gasification models using equilibrium model [12] or CFD calculations [14] have been developed and tested. These models, which include simple reaction schemes, generally allow predicting the syngas composition with a good accuracy for the main gases (CO, H₂, CO₂, CH₄, H₂O) but are unable to predict tar and soot production. On the other hand, detailed chemical schemes based on elementary reactions are able to predict the production of minor products [15,16] and are very useful to better understand the mechanisms leading to undesirable products like soot or tars. These detailed chemical schemes have rarely been integrated into biomass gasification reactor models and validated with experimental results [17,18].

The present study aims to investigate wood particle conversion in a DTR between 800 °C and 1400 °C, both with experiments and simulations. Experiments were conducted in inert and in oxidizing atmospheres with beech wood particles as feedstock. The oxidizing atmospheres included: oxygen which is necessary to heat an autothermal reactor, steam which could be added to improve biomass gasification and carbon dioxide which could be recycled from the reactor output to be used as carrier gas. The amounts of oxygen, steam or carbon dioxide injected were chosen to be representative of an EFR. An existing model [10] was improved to represent biomass particle conversion in presence of these oxidants and was validated by comparison with the experimental results.

2. Materials and methods

2.1. Feedstock

The feedstock used for the experiments is beech sawdust sieved in a size range of 0.315–0.450 mm. The proximate analysis and the ultimate analysis are given in Table 1.

2.2. The drop tube reactor

Experiments were conducted in a Drop Tube Reactor (DTR) presented in Fig. 1. It consists of an alumina tube inserted in a vertical electrical heater with three independent heating zones. The internal diameter of the tube is 0.075 m and the heated zone length is 1.2 m. The DTR works at atmospheric pressure and can reach a maximum temperature of 1400 °C.

The wood particles are continuously fed into the reactor using a gravimetric feeding system, controlled by a computer enabling a ±1% accuracy for the flowrate. The wood particles are entrained to a pneumatic ejector from which they are injected into the reactor. For the introduction of O₂ into the DTR, a steam generator is set at 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 or in a tar protocol if tars are sampled. 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.
2.3. Gas analysis

Main gaseous compounds are analyzed online with a micro-gas chromatograph (H₂, CO, CO₂, CH₄, C₂H₄, C₂H₆, C₃H₈, C₆H₆, O₂ and N₂), a psychrometer (H₂O), a paramagnetic detector (O₂), a Thermal Conductivity Detector (H₂), and a Non-Dispersive Infrared Spectrometer (CO₂). Three different columns are settled in the micro-gas chromatograph: H₂, O₂, N₂, CO and CH₄ are analyzed using a molsieve 5A; CO₂, C₂H₄, C₂H₆, C₂H₂ and C₃H₈ are analyzed using a Propyl-pyridyl-urea column (PPU) connected to a back flush inlet. At last, a non-polar dimethylsilicone capillary column (OV1) is used to measure C₆H₆ concentration.

Gas yields are calculated using the tracer method with N₂ as tracer. Repeatability of these experiments was checked at several days and months of interval; the relative difference between experiments is inferior to 15% in most cases but for some species in some conditions repeatability is not as good. In Section 3, the experimental results are not given as mean values but all repeatability experiments results are plotted. The experimental relative uncertainties on gas yields are estimated at about 10% for CO, CO₂, H₂, CH₄ and 15% for C₂H₂, C₆H₆, C₂H₄, C₂H₆, C₁₃H₈ and steam.

2.4. Tar sampling and analysis

A part of the product gas stream containing tar is continuously sucked and passed through a sampling train composed of a filter, five Erlenmeyer flasks filled with isopropanol – three in a “hot” bath at 40 °C and two in a “cold” bath at –70 °C made with a mixture of isopropanol and carbon ice, a pump and a volumetric gas counter. Each tar sampling lasts about 30 min at a flow rate of 2 NL min⁻¹.

Isopropanol samples are analyzed using a gas chromatographer connected to a Flame Ionization Detector (GC–FID). Knowing the volume of gas passing through the tar protocol, the time of sampling and the concentration of tars in each Erlenmeyer flasks, the tar yields can be deduced. Tar analyses were only performed for experiments at 800 °C in pyrolysis conditions defined in Table 3.

2.5. Solid residue analysis

Char and soot are both collected in the settling box and the filter after each experiment. Char is mainly retained in the settling box whereas soot is mainly in the filter. This segregation phenomenon has been previously observed [6].

Char yield is calculated thanks to the ash tracer method. The ash contents of initial biomass and of chars are measured at 815 °C. Considering that ash totally remains in the char, the char yield can be calculated according to the following equation:

\[ \eta_{\text{char}} = \frac{\theta_{\text{biomass}}}{\theta_{\text{char}}} \]

with \( \eta_{\text{char}} \), the char yield and \( \theta_{\text{char}} \) and \( \theta_{\text{biomass}} \) the ash contents in char and biomass respectively.

Note that this ash tracer method was validated with other furnaces in pyrolysis conditions at 800 °C and 1100 °C. The char yield was determined by weighing on the one hand, and by the ash tracer method on the other hand. The relative error between the two measurements was about 3%.

Two different protocols were used to measure the ash content in char:

- If the available mass of char was superior to 300 mg, a laboratory furnace was used following the ISO 1171: 2010 standard. The minimal char quantity for the ash content measurement was 300 mg.
- If the available mass was under 300 mg, the ash content was measured with a Thermo Gravimetric Analyser (TGA), following the same temperature history as in the laboratory furnace. The required mass of char was about 10 mg. This protocol was validated by comparison with the first one and the maximal relative difference was 5%.

C and H contents in chars were also measured with a microanalyzer when the remaining quantities of chars were sufficient (\( \theta_{\text{char}} > 50 \) mg). The relative uncertainty was calculated for each char yield measurement and ranges between 5% and 15%.

Fig. 1. Scheme of the drop tube reactor.

Fig. 2. Input data and assumptions for the EFR modelling.
Flowrate, $C_p$ the molar heat capacity and between the inlet $O_2$ and outlet temperatures and density were considered to be 20 vol% and 650 kg/m$^3$ feeding line at 25 $^\circ$C and a pressure of 40 bar. The gas residence time was assumed to be about 4 s which allowed sizing the EFR: length = 4.1 m and diameter = 1.6 m. In the pressurized biomass pyrolysis experiments at 25 $^\circ$C, the wood particle volume concentration and density were considered to be 20 vol% and 650 kg/m$^3$ respectively. Heat loss was estimated at 100 kW m$^{-2}$ of internal surface. The biomass specifications are those of the beech sawdust used for the drop tube reactor experiments (Table 1).

The energy balance of the reactor can be written as:

$$\sum_j Q_j \int_{T_i}^{T_{out}} C_p \, dT - \sum_j Q_j \int_{T_i}^{T_{in}} C_p \, dT + \sum_j Q_j \Delta H^f_j(j) - \sum_i Q_i \Delta H^b_i(i) + P_{\text{loss}} = 0$$  \hspace{1cm} (2)

where $i$ concerns the reactants and $j$ the products, $Q$ is the molar flowrate, $C_p$ the molar heat capacity and $\Delta H^b_i$ the formation enthalpy of products and reactants. $P_{\text{loss}}$ is the heat loss through the wall of the reactor. $T_{\text{in}}$ and $T_{\text{out}}$ are respectively the inlet and outlet temperatures and $T_0$ is the reference temperature.

The oxygen inlet flowrate is adjusted to close the energy balance of the system and is the main result of the calculation. From this flowrate, the Equivalent Ratio (ER) which is defined as the ratio between the inlet $O_2$ flowrate and the $O_2$ flowrate needed for complete combustion of biomass can be calculated. Results are given in Table 2 for three levels of steam addition and with $N_2$ or $CO_2$ as carrier gas in the biomass feeding line, whose flowrates were calculated with CHEMKIN. Then the differential system is solved in the DTR. It takes into account different phenomena as particle heating, particle drying, pyrolysis reaction, gas phase reactions and char gasification. The differential equations related to gas, tar and soot production and reaction in the gas phase are calculated with CHEMKIN. Then the differential system is solved with LSODE solver which is appropriate for stiff systems.

### Table 2
Calculated ER as a function of S/B with $N_2$ or $CO_2$ as carrier gas.

<table>
<thead>
<tr>
<th>Carrier gas</th>
<th>$N_2 = 0.28$ g/g db</th>
<th>$CO_2 = 0.44$ g/g db</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steam/biomass (g/g db)</td>
<td>0</td>
<td>0.27</td>
</tr>
<tr>
<td>Steam content (vol%)</td>
<td>0%</td>
<td>33.6%</td>
</tr>
<tr>
<td>ER</td>
<td>0.43</td>
<td>0.46</td>
</tr>
<tr>
<td>$O_2$ content (vol%)</td>
<td>63.7%</td>
<td>43.5%</td>
</tr>
</tbody>
</table>

### 2.6. Experimental conditions

#### 2.6.1. Experimental conditions determination

In order to choose relevant conditions for our experimental study, an autothermal full scale EFR was modelled (Fig. 2) with simple assumptions. Thermodynamic equilibrium was assumed at the output of the reactor (Gibbs Energy minimization) at a temperature of 1500 $^\circ$C and a pressure of 40 bar. The gas residence time was assumed to be about 4 s which allowed sizing the EFR: length = 4.1 m and diameter = 1.6 m. In the pressurized biomass feeding line at 25 $^\circ$C, the wood particle volume concentration and density were considered to be 20 vol% and 650 kg/m$^3$ respectively. Heat loss was estimated at 100 kW m$^{-2}$ of internal surface. The biomass specifications are those of the beech sawdust used for the drop tube reactor experiments (Table 1).

The energy balance of the reactor can be written as:

$$\sum_j Q_j \int_{T_i}^{T_{out}} C_p \, dT - \sum_j Q_j \int_{T_i}^{T_{in}} C_p \, dT + \sum_j Q_j \Delta H^f_j(j) - \sum_i Q_i \Delta H^b_i(i) + P_{\text{loss}} = 0$$  \hspace{1cm} (2)

where $i$ concerns the reactants and $j$ the products, $Q$ is the molar flowrate, $C_p$ the molar heat capacity and $\Delta H^b_i$ the formation enthalpy of products and reactants. $P_{\text{loss}}$ is the heat loss through the wall of the reactor. $T_{\text{in}}$ and $T_{\text{out}}$ are respectively the inlet and outlet temperatures and $T_0$ is the reference temperature.

The oxygen inlet flowrate is adjusted to close the energy balance of the system and is the main result of the calculation. From this flowrate, the Equivalent Ratio (ER) which is defined as the ratio between the inlet $O_2$ flowrate and the $O_2$ flowrate needed for complete combustion of biomass can be calculated. Results are given in Table 2 for three levels of steam addition and with $N_2$ or $CO_2$ as carrier gas in the biomass feeding line, whose flowrates were calculated as described above.

As expected, the required $O_2$ to reach 1500 $^\circ$C increases with the addition of steam, by about 18% when 0.64 g of steam per gram of dry biomass is added. The increase does not depend on carrier gas ($CO_2$ or $N_2$).

With these calculations, the operating conditions of an EFR fed with biomass are specified. They were used to choose the test conditions in the DTR, which are presented in the following section.

### 2.6.2. Experimental conditions in the DTR

In this study, four different temperatures were selected: 800 $^\circ$C, 1000 $^\circ$C, 1200 $^\circ$C and 1400 $^\circ$C. Total inlet gas flowrates were respectively 18.8 NL/min, 15.3 NL/min, 13.7 NL/min and 12.1 NL/min, in order to keep a constant gas mean residence time of 4.3 s. In all tests, the wet biomass feeding rate was 1 g/min.

Different atmospheres were studied: inert ($N_2$) and oxidative with $O_2$, $H_2O$ and $CO_2$ as added reactants. The experimental conditions are listed in Table 3 and are related to the preliminary calculation results (Table 2). Our drop tube reactor is designed for small biomass feeding rates and for working in diluted conditions (the dilution gas was nitrogen). So it was impossible to find operating conditions which allow reproducing both reactants/biomass mass ratio and composition of inlet gas of an EFR. Here, the reactant/biomass mass ratio was kept as representative as possible of an EFR.

Several experimental conditions are out of the ranges calculated previously because of technical limitations of the experimental facility. Experiments were conducted at the four temperatures for each reactive atmosphere.

### 3. Modelling

The experiments were simulated using a numerical model named GASPAR. It was previously adapted and validated to model pyrolysis and steam gasification of biomass in a DTR at high temperature [10]. The GASPAR model having been previously described in [10], only a global description is given. A focus is then made on the chemical reaction modelling, which has been modified.

#### 3.1. The GASPAR model

The GASPAR model is a 1-D model describing biomass gasification in the DTR. It takes into account different phenomena as particle heating, particle drying, pyrolysis reaction, gas phase reactions and char gasification. The differential equations related to gas, tar and soot production and reaction in the gas phase are calculated with CHEMKIN. Then the differential system is solved with LSODE solver which is appropriate for stiff systems.

The major hypotheses of GASPAR are:

- The drop tube reactor is modelled as a plug flow reactor.
- The particles are supposed to be spherical.
- Temperature and concentrations are supposed to be uniform inside the particles.

The slip velocity between particles and gas is taken into account. The slip velocity is calculated by Eq. (3) [19] with the drag

### Table 3
Test conditions in the drop tube reactor.

<table>
<thead>
<tr>
<th>Oxidizing reactant injected</th>
<th>Air flow rate (NL/min)</th>
<th>ER</th>
<th>$H_2O$ flow rate (NL/min)</th>
<th>$H_2O$/biomass (g/g db)</th>
<th>$CO_2$ flow rate (NL/min)</th>
<th>$CO_2$/biomass (g/g db)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrolysis experiments</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Oxygen</td>
<td>1</td>
<td>0.24</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>1.8</td>
<td>0.44</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>2.5</td>
<td>0.61</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Steam ($H_2O$-experiments)</td>
<td>–</td>
<td>0.62</td>
<td>0.55</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Carbon dioxide ($CO_2$-experiments)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.4</td>
<td>0.87</td>
<td>–</td>
</tr>
</tbody>
</table>
coefficient correlation from [20], given in Eq. (4). The model takes into account the modifications of the particle characteristics during the pyrolysis reaction, namely the particle bulk density \( \rho_p \) and the particle equivalent spherical diameter \( d_{eq} \).

\[
\frac{d\rho_p}{dt} = \frac{(\rho_p - \rho_p^d)}{\rho_p} - \frac{3}{4} \frac{\rho_p}{d_{eq}} \frac{1}{\rho_p} \frac{d^2 \rho_p}{dt} \quad \text{(3)}
\]

\[
C_d = 1.5 \left[ \frac{24}{Re_p} \left( 1 + 0.173 \rho_p^{657} \right) + \frac{0.413}{1 + 20 \rho_p^{1.89}} \right] \quad \text{(4)}
\]

This slip velocity model has been validated with experimental particle velocity measurement [19].

3.2. Pyrolysis

Particle pyrolysis is modelled by a one-step reaction where biomass is decomposed into gas, tar and char. The kinetic of pyrolysis reaction follows an Arrhenius law given in Eq. (5).

\[
\frac{dm_{dry \ biomass}}{dt} = -m_{dry \ biomass} \rho_{pyro} \exp \left( -\frac{E_{pyro}}{RT} \right) \quad \text{(5)}
\]

Kinetic parameters were adjusted so that simulation results fit with experimental results obtained for biomass pyrolysis at 800 °C in a DTR similar to the one used for this study [19].

The pyrolysis reaction for dry ash free biomass is represented by:

\[
C_{3}H_{5}O_{2} \rightarrow x_1 H_2 + x_2 CO + x_3 CO_2 + x_4 CH_4 + x_5 C_2H_2 + x_6 C_2H_4 + x_7 C_3H_6 + x_8 CH_6 + x_9 H_2O + x_{10} C_3H_8 + \frac{x_{11}}{x_{12}} C_6H_{12} + \frac{x_{13}}{x_{14}} C_2H_8 + \frac{x_{15}}{x_{16}} C_7H_{16} + \frac{x_{17}}{x_{18}} C_8H_{18} + \frac{x_{19}}{x_{20}} C_9H_{20} + \frac{x_{21}}{x_{22}} C_{10}H_{22} + \frac{x_{23}}{x_{24}} \ldots + \gamma C + C_5H_2O_2 \quad \text{(6)}
\]

Table 4

<table>
<thead>
<tr>
<th>Biomass composition</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Residue composition</td>
<td>s</td>
<td>u</td>
<td>v</td>
</tr>
<tr>
<td>1.2 E^{-3}</td>
<td>1.45 E^{-3}</td>
<td>0.2405</td>
<td></td>
</tr>
</tbody>
</table>

In order to improve the prediction capability of the model, the gas phase reactions were modelled using a detailed kinetic scheme. The Ranzi and co-workers kinetic scheme [176 species, 5988 reactions] [16] was selected for gas phase modelling. It is a detailed lumped mechanism of the pyrolysis, partial oxidation and combustion of Primary Reference Fuels. It takes into account all the tars used here in the global pyrolysis reaction. It was validated in relevant experimental conditions and it predicts PAH formation up to C_{20} which is convenient for the soot formation modelling [21]. Main studies dealing with this chemical mechanism were conducted on benzene in pyrolysis, partial oxidation and combustion conditions [22], on cyclopentadiene pyrolysis with a focus on the PAH formation [23], on heavy n-alkanes (nC_{27}H_{56}, nC_{12}H_{22}, nC_{13}H_{26}, nC_{10}H_{18}) in pyrolysis, partial oxidation and combustion conditions [16].

Soot formation, which is not described in the detailed chemical scheme, is modelled here following a simple reaction pathway including two reactions for soot inception using heavy tars C_{20}H_{16} and C_{20}H_{16} and one reaction of soot growth with C_{2}H_{2}.

\[
C_{20}H_{16} \rightarrow 20 \text{ SOOT} + 8H_2 \quad \text{(7)}
\]

\[
C_{20}H_{16} \rightarrow 20 \text{ SOOT} + 5H_2 \quad \text{(8)}
\]

\[
C_{2}H_{2} + \text{SOOT} \rightarrow 3 \text{ SOOT} + H_2 \quad \text{(9)}
\]

Each reaction is supposed to follow a first order Arrhenius law. The same activation energy as that obtained by Ziegler [24] and used by Septin [10] is used for the inception reactions, while the activation energy for soot particle growth is taken from [25]. The pre-exponential factors were adjusted to fit the experimental data and are presented in Table 6.

3.3. Gas phase reactions

Table 5

<table>
<thead>
<tr>
<th>Formula</th>
<th>Name</th>
<th>Coefficient in Eq. (6)</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>H_2</td>
<td>Hydrogen</td>
<td>x_1</td>
<td>0.753</td>
</tr>
<tr>
<td>CO</td>
<td>Carbon monoxide</td>
<td>x_2</td>
<td>2.192</td>
</tr>
<tr>
<td>CO_2</td>
<td>Carbon dioxide</td>
<td>x_3</td>
<td>0.347</td>
</tr>
<tr>
<td>H_2O</td>
<td>Steam</td>
<td>x_4</td>
<td>0.526</td>
</tr>
<tr>
<td>CH_4</td>
<td>Methane</td>
<td>x_5</td>
<td>0.566</td>
</tr>
<tr>
<td>C_2H_2</td>
<td>Acetylene</td>
<td>x_6</td>
<td>0.059</td>
</tr>
<tr>
<td>C_2H_4</td>
<td>Ethylene</td>
<td>x_7</td>
<td>0.256</td>
</tr>
<tr>
<td>C_3H_6</td>
<td>Ethane</td>
<td>x_8</td>
<td>0.0211</td>
</tr>
<tr>
<td>C_4H_8</td>
<td>Propane</td>
<td>x_9</td>
<td>0.1573</td>
</tr>
<tr>
<td>C_6H_6</td>
<td>Benzene</td>
<td>x_10</td>
<td>0.0391</td>
</tr>
<tr>
<td>C_6H_8</td>
<td>Toluene</td>
<td>x_11</td>
<td>0.0326</td>
</tr>
<tr>
<td>C_7H_8</td>
<td>Phenol</td>
<td>x_12</td>
<td>0.0243</td>
</tr>
<tr>
<td>C_8H_8</td>
<td>Styrene</td>
<td>x_13</td>
<td>0.0164</td>
</tr>
<tr>
<td>C_9H_9</td>
<td>Indene</td>
<td>x_14</td>
<td>0.0169</td>
</tr>
<tr>
<td>C_{10}H_{10}</td>
<td>Naphthalene</td>
<td>x_15</td>
<td>0.0286</td>
</tr>
<tr>
<td>C_{11}H_{12}</td>
<td>Acenaphthylene</td>
<td>x_16</td>
<td>0.0042</td>
</tr>
<tr>
<td>C_{12}H_{14}</td>
<td>Phenanthrene</td>
<td>x_17</td>
<td>0.0021</td>
</tr>
<tr>
<td>C</td>
<td>Char</td>
<td>γ</td>
<td>0.9161</td>
</tr>
</tbody>
</table>

Stoichiometric parameters of the reaction come from pyrolysis experimental result at 800 °C in the DTR. Light gas and benzene coefficients (x_i) were directly fitted to Arrhenius law parameters. Steam and tar compound coefficients (\( \beta_i \)) were adjusted to fill the mass balance, keeping the same tar mixture composition as the one determined by the tar protocol in pyrolysis experiment at 800 °C. The residue C_2H_5O, which corresponds to unidentified compounds, represents only 2.5 wt% of initial dry biomass. This residue is considered not to participate in any reaction. In this reaction char is assumed to be pure carbon. A single set of stoichiometric parameters was used for all simulations (Tables 4 and 5).

3.3. Gas phase reactions

In order to improve the prediction capability of the model, the gas phase reactions were modelled using a detailed kinetic scheme. The Ranzi and co-workers kinetic scheme [176 species, 5988 reactions] [16] was selected for gas phase modelling. It is a detailed lumped mechanism of the pyrolysis, partial oxidation and combustion of Primary Reference Fuels. It takes into account all the tars used here in the global pyrolysis reaction. It was validated in relevant experimental conditions and it predicts PAH formation up to C_{20} which is convenient for the soot formation modelling [21]. Main studies dealing with this chemical mechanism were conducted on benzene in pyrolysis, partial oxidation and combustion conditions [22], on cyclopentadiene pyrolysis with a focus on the PAH formation [23], on heavy n-alkanes (nC_{27}H_{56}, nC_{12}H_{22}, nC_{13}H_{26}, nC_{10}H_{18}) in pyrolysis, partial oxidation and combustion conditions [16].

Soot formation, which is not described in the detailed chemical scheme, is modelled here following a simple reaction pathway including two reactions for soot inception using heavy tars C_{20}H_{16} and C_{20}H_{16} and one reaction of soot growth with C_{2}H_{2}.

\[
C_{20}H_{16} \rightarrow 20 \text{ SOOT} + 8H_2 \quad \text{(7)}
\]

\[
C_{20}H_{16} \rightarrow 20 \text{ SOOT} + 5H_2 \quad \text{(8)}
\]

\[
C_{2}H_{2} + \text{SOOT} \rightarrow 3 \text{ SOOT} + H_2 \quad \text{(9)}
\]

Each reaction is supposed to follow a first order Arrhenius law. The same activation energy as that obtained by Ziegler [24] and used by Septin [10] is used for the inception reactions, while the activation energy for soot particle growth is taken from [25]. The pre-exponential factors were adjusted to fit the experimental data and are presented in Table 6.

3.4. Char gasification

Both char gasification reactions with steam and carbon dioxide are taken into account. Temperature and oxidant concentrations are supposed to be homogeneous in the whole particle. The reactions are supposed to follow Arrhenius laws and the reactive surface of char is supposed to decrease homogeneously in the particle following a Volume Reaction Model:

\[
\frac{dX}{dt} = k(T, P_{H_2O}, f(X)) + k(T, C_{O_2}, f(X)) \quad \text{(10)}
\]

\[
= \left( A_{H_2O} \exp \left( -\frac{E_{H_2O}}{RT} \right) \right) P_{H_2O}^{\beta_{H_2O}} + \left( A_{C_{O_2}} \exp \left( -\frac{E_{C_{O_2}}}{RT} \right) \right) P_{C_{O_2}}^{\beta_{C_{O_2}}} (1 - X) \quad \text{(11)}
\]
$P_i$ is the partial pressure of $i$ in the particle, $T$ is the temperature in the particle, $A_i$, $E_{a i}$, and $n_i$ are the kinetic parameters of char gasification by oxidant $i$ ($i = \text{CO}_2$ and $\text{H}_2\text{O}$), and $X$ is the char conversion (Eq. (11)).

\[
X = 1 - \frac{m(t)}{m(t = 0)} \quad (11)
\]

with $m(t)$, the mass of char at time $t$.

Gasification kinetic parameters were optimized to fit the experimental mass of char measured at the end of experiments. Experimental results used to optimize the kinetic parameters are presented in Fig. A1 in supplementary data. These experiments were conducted at 4 different temperatures – 800, 1000, 1200 and 1400 °C – in three different atmospheres: inert, and with addition of $\text{H}_2\text{O}$ and $\text{CO}_2$ (Table 3). The set of fitted parameters is presented in Table 7.

These sets of parameters are relevant compared to the results given in literature for woody biomass gasification [26]. However, gasification can be limited by the diffusive transfer in particle at high temperature [6], which is thus included in the optimized parameters.

Note that the oxidation of solids (char and soot) by $\text{O}_2$ was not taken into account in the model. This choice will be discussed in the following of the text. Moreover, soot gasification by steam and $\text{CO}_2$ is neglected. Indeed, soot gasification experiments have shown that soot gasification by steam was 2–20 times slower than char gasification between 750 °C and 950 °C [27]. Soot gasification by $\text{CO}_2$ is 2–30 times slower than char gasification at 1100 °C [28].

### 4. Results and discussion

Both experimental and simulation results are presented and discussed in this section. First, the influence of presence of steam and $\text{CO}_2$ on biomass gasification is studied by comparison with pyrolysis experiments, whose results are not detailed here as they are similar to those found in the same facility by Septien [10]. Then the influence of $\text{O}_2$ is detailed and discussed.

#### 4.1. Influence of steam or $\text{CO}_2$ addition

The influence of the addition of $\text{H}_2\text{O}$ or $\text{CO}_2$ on carbon conversion into gas, char and remaining products (soot and tar) is discussed first. Then the gas composition is detailed. At last, the simulation results are used to better understand the influence of the addition of $\text{H}_2\text{O}$ or $\text{CO}_2$ on tar and soot production.

##### 4.1.1. Distribution of carbon in products

The goal of the gasification being to produce a maximum of $\text{CO}$ and $\text{H}_2$, it is interesting to study the carbon repartition in gas, char and tar + soot in this section. We define the carbon conversion into gas as the ratio between the mass of carbon in the analyzed gas ($\text{CO}_2$, $\text{CO}$, $\text{CH}_4$, $\text{C}_2\text{H}_6$, $\text{C}_2\text{H}_4$, $\text{C}_2\text{H}_2$, $\text{C}_6\text{H}_6$, and $\text{C}_6\text{H}_5$) and the mass of carbon coming from biomass. Thus the carbon added by $\text{CO}_2$ injection in $\text{CO}_2$-experiments is subtracted. The expression of carbon conversion into gas is given in Eq. (12).

\[
\text{Carbon conversion} = \frac{m_{\text{gas analyzed}} - m_{\text{CO}_2\text{in}}}{m_{\text{biomass}}} \quad (12)
\]

The fraction of carbon in char is calculated from char yield and carbon content of char (about 80 wt%). Finally the remaining carbon, attributed to soot and tars, is calculated to fill the carbon balance.

Carbon distribution in gas, char and in tar + soot products is given in Fig. 3 as a function of temperature.

The addition of oxidant ($\text{H}_2\text{O}$ or $\text{CO}_2$) has a significant influence on carbon distribution especially at 1200 and 1400 °C. The conversion of carbon into gas in pyrolysis experiments reaches a maximum at 1000 °C (67% of carbon from initial biomass) and remains constant between 1200 and 1400 °C. In $\text{H}_2\text{O}$ and $\text{CO}_2$-experiments the maximum is reached at 1400 °C with respectively 77% and 71% of carbon from initial biomass. The simulation gives a good prediction of carbon conversion into gas even if the calculation overestimates the carbon conversion at 1000 °C.

The carbon fraction into char decreases as temperature increases. This is attributed to char gasification reaction. Addition of steam or $\text{CO}_2$ has a significant influence on char consumption at 1200 °C and 1400 °C. Altogether the model with fitted kinetic parameters allows a good representation of char consumption in function of temperature and reproduces very satisfactorily the influence of $\text{H}_2\text{O}$ and $\text{CO}_2$.

### Table 7

<table>
<thead>
<tr>
<th>Parameter</th>
<th>$\text{H}_2\text{O}$</th>
<th>$\text{CO}_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{a i}$ (J mol$^{-1}$)</td>
<td>$132.1 \times 10^3$</td>
<td>$141.3 \times 10^3$</td>
</tr>
<tr>
<td>$A_i$ (s$^{-1}$ bar$^{-n}$)</td>
<td>$254 \times 10^2$</td>
<td>$218.3 \times 10^2$</td>
</tr>
<tr>
<td>$n_i$</td>
<td>1</td>
<td>0.683</td>
</tr>
</tbody>
</table>

**Fig. 3.** Carbon distribution in gas, char and tar + soot in pyrolysis, $\text{H}_2\text{O}$ and $\text{CO}_2$-experiments – experimental results (dots) and simulation results (lines).
The fraction of carbon in soot and tars determined from experiments seems to remain constant between 800 and 1000 °C, and then increases to reach a plateau between 1200 and 1400 °C. This could be explained by two opposite phenomena: the tar yield decreases when temperature increases because of cracking reactions, whereas soot production increases with temperature. Indeed, previous works show that tar contents are very low above 1000 °C [7,10] while soot production is significant at 1000 °C and above [4]. The presence of steam or CO₂ leads to a decreasing amount of carbon in tar and soot, certainly because of soot precursors consumption, which is discussed in more details in Section 4.1.3. The simulation allows well reproducing the carbon distribution except at 1000 °C where carbon conversion into gas is overestimated and carbon in tar and soot is underestimated.

### 4.1.2. Gas species yields

The experimental and simulated gas species yields, as well as those at thermodynamic equilibrium, are shown as a function of temperature in Fig. 4 and Fig. A2 in supplementary data. Note that gas yields include the H₂O or CO₂ addition.

In all experiments, H₂, CO, H₂O, CO₂ and CH₄ are the major gas species, followed by C₂H₂, C₂H₄, C₃H₆, C₂H₆ and C₃H₈. H₂ and CO yields always increase with temperature. H₂O yield decreases as temperature increases except in CO₂ experiments in which it slightly increases. The variation of CO₂ yield depends on experimental conditions: in pyrolysis experiments CO₂ yield is steady with temperature until 1200 °C and then decreases, while in H₂O-experiments it increases until 1200 °C and then decreases; in CO₂-experiments the CO₂ yield is steady between 800 and 1000 °C and then strongly decreases.

The addition of H₂O or CO₂ on CO and H₂ yields is notable above 1000 °C. At 1200 and 1400 °C, H₂ yield decreases in CO₂-experiments and increases in H₂O-experiments while CO yield increases in CO₂-experiments.

Influence of addition of H₂O or CO₂ on CO and H₂ yields has no significant influence even at high temperature. In our operating conditions, the steam reforming and vapo-cracking reactions of light hydrocarbons are not significantly enhanced by the steam addition. In all conditions, the results of the tests and of the model are different from the predictions at thermodynamic equilibrium. The model allows reproducing the major gas yields with a very good accuracy (relative error < 20% in most cases). The maximal deviations are obtained for H₂ yields at 1000 °C and steam yield at 1400 °C in pyrolysis.

![Graphs showing gas yields as a function of temperature](image-url)
experiments. Simulation also accurately predicts CH₄ and C₂H₂ yields in the whole range of temperature (Fig. A2 in supplementary data). However, C₂H₄ production is overestimated at 1000 °C and benzene yields are systematically overestimated at 800, 1000 and 1200 °C. In spite of these slight differences the tendencies are always well reproduced, which is very satisfying.

According to the previous observations, H₂O or CO₂ addition mainly influences H₂, CO, H₂O and CO₂ yields, at 1200 and 1400 °C only. Three global phenomena could lead to this result: char gasification, tar and soot gasification, and water-gas shift reaction (Eq. (13)).

\[
\text{CO} + \text{H}_2\text{O} \rightarrow \text{CO}_2 + \text{H}_2
\]  

The two first phenomena lead to an increase of H₂ and CO yields. However, as noticed before, at 1200 and 1400 °C, in CO₂ experiments, the H₂ yield is lower than in pyrolysis experiments, while in H₂O experiments, the CO yield hardly increases compared to pyrolysis. This should be due to the third phenomenon mentioned above: the water-gas shift reaction. The water-gas shift constants calculated from experimental results and at thermodynamic equilibrium are given in Fig. 5 for each condition.

As shown in Fig. 5, the water-gas shift reaction is at thermodynamic equilibrium at 1200 and 1400 °C, whatever the atmosphere. This reaction then controls the relative H₂, CO, CO₂ and H₂O contents.

4.1.3. Soot and tar yields

The soot and tar yields cannot be precisely measured in experiments and we only have a qualitative estimation by direct observation. The GASPAR model shows a good ability to predict the gas and char yields, thus it is used here to investigate the influence of temperature and H₂O and CO₂ on soot and tar yields. Fig. 6(a) represents the soot yields calculated with GASPAR.

A high soot yield (between 0.1 and 0.17 kg/kg biomass db) is predicted at 1200 and 1400 °C as observed in experiments. The presence of steam decreases the soot yield although no soot gasification reactions are taken into account in the model. Steam or radicals derived from steam could react with soot precursors. Liu et al. [29] show that major chemical effect of addition of steam on soot formation inhibition is due to OH formation which is responsible for soot precursors decrease. The influence of CO₂ is lower but significant at 1400 °C. CO₂ only has a slight chemical effect on soot inhibition by enhancing OH formation [30]. In our case, H₂O produced by RWGS reaction is certainly also responsible for the inhibition of soot formation in CO₂-experiments.

Let us remind that the tar compounds are defined here as the compounds heavier than benzene. The GASPAR predicted tar yields are given in Fig. 6(b). The main compounds predicted by GASPAR are Polynuclear Aromatic Hydrocarbons (PAH): naphthalene, indene, styr-ene, acenaphthylene, phenanthrene, pyrene and C₂₀ species. The tar yield decreases when temperature increases because of PAH cracking and soot formation. The presence of oxidant has a very weak influence on tar yields.

In conclusion, the influence of addition of steam or CO₂ on gasification products is rather similar. This influence is only visible at 1200 and 1400 °C. The presence of oxidant then leads to a char conversion increase and a soot formation decrease, which leads to a higher carbon conversion into gas. The gas phase composition is mainly affected by the WGS reaction while light hydrocarbon yields remain unchanged. The GASPAR simulation shows a very satisfying agreement with experimental results.

![Fig. 5. Water-gas-shift constants from pyrolysis, H₂O and CO₂-experiments and at thermodynamic equilibrium as a function of temperature.](image)

![Fig. 6. Soot yields (a) and tar yield (b) as a function of different temperature calculated with GASPAR in pyrolysis, H₂O and CO₂-experiments.](image)
4.2. Influence of $O_2$

First, the influence of $O_2$ on carbon distribution is discussed. Then the char and gas yields are presented as a function of ER. At last, the simulation results are used to better understand the influence of the addition of $O_2$ on tar and soot production.

4.2.1. Distribution of carbon from biomass

The carbon distribution into gas, char and tar + soot are given at each temperature as a function of ER in Fig. 7. Note that tar + soot for experimental results is the remaining part to close the carbon balance.

According to experimental results, carbon conversion into gas increases with ER whatever the temperature and reaches a plateau at about 80–85%. At 800 and 1000 °C, this plateau is due to incomplete char conversion even at ER = 0.61. Indeed, the carbon fraction in char has a minor dependence on ER and remains almost constant at about 10–13%. At 1200 and 1400 °C, the fraction of carbon in char decreases as ER increases and reaches 0% at 1400 °C and ER = 0.61. The plateau can only be explained by production of tar and soot. We know that tars are converted at these temperatures thus the remaining part of carbon – around 20% – mainly comes from soot which are produced at high temperature. However, we have found out that only a very small quantity of soot, which cannot represent 20% of input carbon, was produced for the tests at 1400 °C with ER of 0.44 and 0.61. Additional tests were performed to check the experiment repeatability, confirming the same inconsistency. The soot yield being obtained by difference of the carbon balance, we think that this discrepancy could come from an error in the measurement of gas yields in these conditions.
The simulation results are in very good agreement with experimental ones in all conditions at 800 and 1000 °C, and for the lower values of ER at 1200 and 1400 °C. The conversion of carbon into gas predicted by the model is higher than the one determined from the experiments in the other cases.

To conclude this section, at 800 and 1000 °C the conversion of carbon into gas is limited by a significant fraction of carbon in unconverted char, while at 1200 and 1400 °C, soot seems to be responsible for the carbon gasification limitation. In the next section, the char yield evolution with ER and temperature is detailed.

4.2.2. Char yield

The char mass yields are given in Fig. 8 as a function of ER at different temperatures.

Experimental and simulation results both show that char yield decreases as temperature increases whatever the ER, because of gasification reaction enhancement. It also decreases as ER increases even if the influence of ER is smaller at 800 and 1000 °C than at 1200 and 1400 °C.

The GASPAR model gives a satisfying prediction of char yields, especially at 1200 and 1400 °C. Let us remind that it takes into account char gasification by steam and CO₂ but not direct combustion of char (Section 3.4). The kinetic parameters of the gasification reactions were not fitted with these experimental results, but only with those obtained with addition of H₂O and CO₂. The good agreement between these experimental and modelling results suggests that in our reactor and in these operating conditions, the char combustion is negligible. The O₂ and char yields profiles calculated with GASPAR along the reactor show that when char is produced – at the end of pyrolysis reaction – no oxygen remains to combust char. Thus the char yield decrease as ER increases would only be due to the increase of steam and CO₂ partial pressures, H₂O and CO₂ being produced by the combustion of a part of tars and gas species. The gas yields and gas phase reactions are discussed in the next section.
4.2.3. Gas species yields

Experimental and simulated gas species yields, together with predictions at thermodynamic equilibrium, are presented in function of ER at different temperatures in Figs. 9 and 10.

The experimental results show that at each temperature, H\textsubscript{2} yield decreases as ER increases, which is probably due to its oxidation. CO\textsubscript{2} and H\textsubscript{2}O yields increase with ER at all temperatures, because they are final combustion products. At 800 and 1000 °C, the CO yield first increases with ER to reach a maximum for ER ranging between 0.2 and 0.4, depending on the temperature. At 1200 °C and 1400 °C, the CO yield decreases first slightly, and then more rapidly as ER increases. The minimum of CO yield is always measured at ER = 0.61 whatever the temperature. This behaviour is certainly due to two competitive phenomena. At low ER, CO is produced by partial combustion of hydrocarbons while at higher ER it is also partially oxidized. All hydrocarbon gas yields decrease as ER increases because of combustion reactions.

Simulation globally gives rather good predictions of gas yields and confirms our explanations on the evolution of gas yields with ER. Except from the cases with ER equal to 0.4 and 0.61 at 1400 °C, the results of our model differ from the calculations at thermodynamic equilibrium. So in all other cases, reaction kinetics have a significant influence on the product yields at the reactor output. If we look at the gas composition along the reactor with GASPAR, both tar and gas are burnt. Indene, naphthalene, toluene, and styrene are burnt first, followed by phenanthrene, acenaphthylene and benzene and at last by methane, acetylene, di-hydrogen and carbon monoxide. Note that for temperatures of 1000 °C and higher, gas combustion kinetic rates are higher than pyrolysis one, and all hydrocarbons released in pyrolysis are burnt as soon as they are formed, as long as some O\textsubscript{2} is still present in the reactor. So soot particles can only be formed in the lower part of the reactor, where no O\textsubscript{2} remains. This confirms our choice not to consider direct oxidation of soot particles in the model, which would not have modified the results. At high temperature, CO and H\textsubscript{2} are also produced by the WGS reaction which reaches thermodynamic equilibrium at the end of the reactor. At 800 °C in presence of oxygen, light hydrocarbons yields are generally underestimated while H\textsubscript{2}.

Fig. 10. CH\textsubscript{4}, C\textsubscript{2}H\textsubscript{2}, C\textsubscript{2}H\textsubscript{4} and C\textsubscript{3}H\textsubscript{6} yields as a function of ER at 800, 1000, 1200 and 1400 °C – experimental results (dots), simulation results (continuous lines) and thermodynamic equilibrium results (dotted lines).
yield is overestimated at ER = 0.61. This seems to be due to a too long delay between pyrolysis reaction and ignition of pyrolysis gas. CO yields are not so well represented especially at 1000 °C where CO yields are underestimated and at 1400 °C where they are overestimated. To conclude, GASPAR model allows a satisfying prediction of gases above 800 °C. Based upon this, we use it to investigate soot and tar yields in the next section.

4.2.4. Soot and tar yields
Influence of oxygen on soot and tar yields is studied with simulation results given by GASPAR. Soot yields are given in Fig. 11(a) as a function of ER.

O₂ addition has a strong influence on predicted soot yield which decreases as ER increases. As expected the maximum yields are found at 1200 and 1400 °C. It was previously discussed that soot yields at 1400 °C and 1200 °C at ER = 0.61 are certainly underestimated even if the decrease in soot production was experimentally observed. As mentioned previously, no soot combustion or gasification reaction is included in the model. Decrease of soot yield can only be due to consumption of soot precursors.

Tar yield as a function of ER at different temperature is given in Fig. 11(b). As expected, increase of ER decreases the tar yield and no tar are predicted at 1400 °C. In Section 4.1, we put into evidence that addition of H₂O and CO₂ has no influence on tar yield in the model. Considering that the ranges of CO₂ and steam concentration are similar, tar yield decrease is thus only due to direct combustion of tar. Tar being soot precursors, the soot decrease is mainly due to tar burning.

5. Conclusion
Both experiments in a drop tube furnace and simulation with the GASPAR model enabled to study the influence of H₂O, CO₂, and O₂ addition on gasification products in an entrained flow reactor. The work was conducted in representative conditions of an entrained flow reactor: oxidant/biomass ratio, temperature, particle size and residence time.

The GASPAR model was improved with (1) a new pyrolysis global reaction, which takes into account the main tars produced, (2) a more efficient gas phase model which allows predicting heavy tars until C₂₀, (3) two fitted gasification laws to take into account the influence of both H₂O and CO₂ and (4) a simple soot formation mechanism. The model is validated on the whole range of experimental conditions and generally allows a good prediction of gas and char yields as well as relevant evolution of soot and tar yields. It can be used for predictive calculations taking the obvious precautions in the use of calculation results. It also enables to investigate the detailed kinetics pathways responsible for the formation of problematic species in EFR process as soot, methane, and char.

Char yield decrease as ER increases can be very well predicted accounting for char gasification by H₂O and CO₂ only. This lets us think that char oxidation by O₂ is negligible in our conditions. Similarly, soot yield decrease as ER increases is predicted by GASPAR taking into account neither direct combustion nor steam gasification of soot. The soot precursors certainly react with OH radicals which stops soot inception.

In typical process conditions in an EFR −1400 °C and ER = 0.44 – char is almost completely gasified and only a small amount of soot (2 wt% db) is produced. Considering the steam and CO₂ influence, their addition seems unnecessary. Indeed it could lower a bit the soot production but it will certainly also decrease the energy yield of the process.

Appendix A. Supplementary material
Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fuel.2015.10.046.

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