Solar Biomass Gasification Combined With Iron Oxide Reduction for Syngas Production and Green Iron Metallurgy

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The solar gasification of biomass with iron oxide for combined syngas and iron production was investigated. Both solar energy and biomass are promising renewable energies. The process of gasification converts solid carbonaceous feedstocks into either fuels or chemicals. However, conventional processes require partial combustion of the feedstock for energy supply and inherently suffer from high oxygen production costs and low syngas calorific value due to dilution with combustion products. Chemical looping gasification using solid oxides is an alternative option to tackle these issues. By supplying concentrated solar energy as the high-temperature heat source, it is possible to produce even more syngas from the process while enabling solar energy storage into dispatchable fuels. This work proposes to explore solar biomass gasification over iron oxide at high heating rates, representative of the conditions obtained in solar reactors. Thermodynamic equilibriums of gasification reactions between 100 and 1,500°C were calculated and experimental results obtained at 1,100°C with a specially designed induction furnace were reported for biomass gasification with iron oxide, water, or carbon dioxide as oxidizing agents. Solid products analysis showed that iron oxide can be reduced to metallic iron depending on the proportion of the oxygen carrier. These results indicate that iron oxide is an effective material for solar biomass gasification producing both syngas and iron via a novel green metallurgical process.

Keywords: solar energy, chemical looping gasification, green metallurgy, iron oxide, biomass

INTRODUCTION

Climate change due to greenhouse gases emissions is one of the main current challenges of our society. The transition from a fossil-based economy to renewable energies is ongoing. Research is very active in this field in order to develop a new energy model. Biomass gasification has been identified as a promising pathway toward the production of renewable fuels with limited greenhouse gas emissions (Kumar et al., 2009; Codina Gironès et al., 2018). The conventional autothermal process needs to burn about 1/3 of the raw biomass feedstock to provide the enthalpy of the reaction. It requires either upstream air separation units (oxy-combustion) or downstream gas separation if air is used as oxidizer or gasification agent, as the products are diluted into nitrogen. In order to supply the required heat, concentrated solar energy has been considered to
eliminate combustion and products contamination, while the high achievable temperatures and heating rates lead to high quality syngas with no CO₂ emission and solar energy is stored into chemical form (Nzihou et al., 2012; Loutzenhiser and Muroyama, 2017). Three main solar reactor concepts have been tested previously (Piatkowski et al., 2011): indirectly irradiated packed bed, directly irradiated vortex flow and indirectly irradiated entrained flow reactors. Packed bed reactors are robust but heat and mass transfer are limited even if the residence time is long. Vortex flow and entrained flow reactors show improved heat and mass transfer but the feedstock size is limited. Direct irradiation enables better heat transfer to the reactants but the challenge is to avoid any dirt on the window. The solar spouted bed reactor has recently been proposed (Boujjat et al., 2019). It can operate either in direct or indirect heating modes. Heat and mass transfer are enhanced by the gas jet that stirs the particles injected continuously in the reactor. Various types of feedstock can be treated as the solid residence time is long.

Wood biomass (overall formula can be assumed as C₆H₉O₄) can be commonly gasified either with steam or CO₂ (Billaud et al., 2016).

Steam gasification:

\[
C_6H_9O_4 + 2H_2O \rightarrow 6.5H_2 + 6CO \quad \Delta H^\circ = 800kJ/mol \tag{1}
\]

CO₂ (dry) gasification:

\[
C_6H_9O_4 + 2CO_2 \rightarrow 4.5H_2 + 8CO \quad \Delta H^\circ = 883kJ/mol \tag{2}
\]

Chemical looping can also be considered. Such process is a relatively old concept initially proposed for CO₂ separation in combustion. Then, it was applied to gasification, reforming, and more recently to chemical looping hydrogen production (Zhao et al., 2017). The objective of chemical looping gasification is to rely on a low-cost oxygen source and self-heating system, while producing high quality syngas. Iron oxide is one of the most studied solid oxygen carriers (Yu et al., 2019) while the concept of liquid chemical looping has only recently been proposed (Sarafraz et al., 2017a,b). It offers the major advantages of being a low-cost material with no environmental impact. Various types of carbonaceous feedstocks have been investigated. Wu et al. suggested to study the kinetics using cellulose as the main organic component of biomass (Wu et al., 2019). The main reported works used electric furnaces (Huang et al., 2014), or thermogravimetric analysis (Huang et al., 2015; Zeng et al., 2018) with a heating rate of about 10°C/min. A systematic review proposed by Gauthier et al. (2013) highlights a particular need to use experimental setups for which heating conditions are controlled (imposed temperature or heat flux). When using solar energy, it is possible to supply high-temperature solar heat to the gasifier and the air reactor (oxidation step) can be replaced by a water or CO₂ splitting reactor to produce additional syngas (Krenzke et al., 2017). This splitting method was proposed in the steam-iron process that uses syngas (or carbonaceous feedstocks) to reduce iron oxide. The reduced form of iron oxide is then re-oxidized with steam to form hydrogen and magnetite (Hacker et al., 2000). Solar-driven extractive metallurgy has also been investigated (Fernández-González et al., 2018). Solar chemical looping biomass gasification has only been explored over zinc oxide, zinc product being a volatile material (Chuayboon et al., 2018). In this work, the biomass gasification over iron oxide (non-volatile material) in an induction furnace was investigated. The objective was to compare biomass reactivity under high heating rates similar to the ones achieved in solar reactors (500°C/C/s) with different oxidizing agents (iron oxide, H₂O or CO₂). The overall endothermic reaction is:

\[
C_6H_9O_4 + 2/3Fe_2O_3 \rightarrow 4/3Fe + 6CO + 4.5H_2 \quad \Delta H^\circ = 865kJ/mol \tag{3}
\]

First, the biomass pyrolysis occurs leading to syngas, tars and char (carbon). Char can be further gasified by iron oxide. The Fe₂O₃ carbothermal reduction steps are as follows:

\[
C + 3Fe_2O_3 \rightarrow 2Fe_3O_4 + CO \quad \Delta H^\circ = 121.6kJ/mol \tag{4}
\]

\[
C + Fe_2O_3 \rightarrow 3FeO + CO \quad \Delta H^\circ = 207.1kJ/mol \tag{5}
\]

\[
C + FeO \rightarrow Fe + CO \quad \Delta H^\circ = 156kJ/mol \tag{6}
\]

Pyrolysis gases undergo subsequent reactions including mainly reforming and water-gas shift reactions (Dupont et al., 2007).

Since metallic Fe can be obtained, the process could support the energy transition in the metallurgical industry whose conventional processing is releasing large amounts of CO₂ in the atmosphere due to the extensive use of fossil resources (accounting for 6% of anthropogenic CO₂ emissions) (IEAGHG, 2013).

In addition, Fe can also be converted back to Fe₂O₃ by water oxidation producing hydrogen while Fe₂O₄ can be oxidized to Fe₂O₃ only with oxygen (Yang et al., 2008). The exothermic oxidation reactions can be written as:

\[
Fe + 4/3H_2O \rightarrow 1/3Fe_2O_3 + 4/3H_2 \quad \Delta H^\circ = -50kJ/mol \tag{7}
\]

\[
Fe_3O_4 + 1/4O_2 \rightarrow 3/2Fe_2O_3 \quad \Delta H^\circ = -116kJ/mol \tag{8}
\]

This last step could be an option to ensure heat supply for continuous biomass conversion at night or during transient periods with low solar power input in the case of a solar chemical looping gasification process [thermochemical heat storage (André et al., 2018)].

In this study, thermodynamic equilibrium calculations were first performed in order to determine the chemical species involved in the case of biomass gasification with H₂O, CO₂, and iron oxide. Then, the experimental equipment and procedure were described and the results were analyzed and compared for the three oxidizing agents.

**THERMODYNAMIC EQUILIBRIUMS**

**Method**

Thermodynamic equilibriums were calculated using Gibbs free energy minimization (GEMINI2 software) (Cheynet and...
This approach uses databases containing the standard Gibbs energy of the most common molecules. The overall formula of beech wood was considered as $C_{6}H_{12}O_{4}$ based on its global chemical composition. Most thermodynamic properties of gases and the different iron oxides are available in this database but there is no data for any kind of solid biomass feedstock. Thus, biomass was included into the model in the form of dioxygen, dihydrogen, and solid carbon. This model presents several advantages as it does not require to know the different reactional mechanisms and can be applied to any reactor geometry. Furthermore, it is possible to change the temperature and pressure conditions to study their impact on the composition of the mix at the equilibrium state. The equilibrium results are thermodynamic predictions for a closed system that do not take into account any kinetic aspect, especially the reactants residence time but also reaction site deactivation, sintering, and agglomeration. Anyway, this is a first approach to determine the theoretical limit of the conversion process and to identify expected products.

In the case of thermodynamic equilibrium studies, the reactants are often introduced in stoichiometric proportion which is not the case in a real gasification process. To ensure an optimal reaction rate with biomass in a real process, the oxidizing agent is often introduced in over-stoichiometric proportion. Thus, thermodynamic equilibriums were calculated with both stoichiometric and over-stoichiometric proportions of oxidizing agent in order to study the impact on the products composition. For each situation, the calculations were realized at temperatures ranging from $100^\circ C$ to $1,500^\circ C$ every $50^\circ C$ at atmospheric pressure. The results relate to the quantity of the main gases ($H_{2}$, $CO$, $CO_{2}$, $CH_{4}$, and $H_{2}O$) produced per gram of gasified dried biomass along with the LHV (Low Heating Value) of the produced gas mix ($CO$, $H_{2}$, and $CH_{4}$). The other predicted gas species, such as hydrocarbons ($C_{2}H_{4}$, $C_{2}H_{6}$), appearing in very low amounts ($<10^{-5} \text{ mmol/g}_{\text{biomass}}$) are not presented and are neglected. Calculations in over-stoichiometric proportion of oxidizing agent ($H_{2}O$, $CO_{2}$, $Fe_{2}O_{3}$) were realized with twice and three times the stoichiometric amount of oxidizing agent. Even though the results were slightly different, the obtained tendencies are similar and thus only the results of the calculations with three times the stoichiometric proportion of oxidizing agent will be presented.

**Water as Oxidizing Agent**

The gas species produced from steam biomass gasification in stoichiometric and over-stoichiometric proportions are presented in Figure 1.

Concerning the results in stoichiometric proportion, $CH_{4}$ and $CO_{2}$ productions are predominant compared to $CO$ and $H_{2}$ below $450^\circ C$. With the increase of the $H_{2}$ production starting at $250^\circ C$, a decline of the $CH_{4}$ produced occurs (as well as the remaining water). The quantity of produced $CO_{2}$ rises steadily until reaching a peak at $550^\circ C$ and then decreases to become insignificant above $900^\circ C$. All the carbon of the original biomass is found in the form of $CO$ above $900^\circ C$. The equation (1) completion is reached from $900^\circ C$ and products remain stable at higher temperatures (the gas mix LHV is then 23.39 MJ/kg$_{\text{biomass}}$). As a comparison, the LHV of a typical dry wood biomass is 18 MJ/kg, which demonstrates the upgrade of the calorific value of the feedstock. These results are in agreement with those reported for the gasification of cellulose (Hathaway et al., 2014) and carbon (Gokon et al., 2015), the main difference being the higher production of $H_{2}$ due to the higher quantities of hydrogen contained in the presently selected biomass.

When steam is present in over-stoichiometric proportion, the production of $CO_{2}$ is higher and is responsible for a lower production of CO. Even though the production of $CO_{2}$ starts decreasing at higher temperatures, $CO_{2}$ is still present at $1,500^\circ C$. The production of $H_{2}$ is also higher at any given temperature. The LHV of the gas mix at high temperature (maximum of 23.09 MJ/kg$_{\text{biomass}}$ at $1,500^\circ C$) is slightly lower than the one in stoichiometric proportion.

**$CO_{2}$ as Oxidizing Agent**

Figure 2 displays the equilibrium composition for biomass gasification with stoichiometric and over-stoichiometric $CO_{2}$ proportion (Equation 2).

As expected, the quantity of $CO$ produced from gasification with $CO_{2}$ is higher than with steam gasification whereas the $H_{2}$ quantity is lower. The completion of Equation (2) is only reached above $1,000^\circ C$ due to the fact that the Boudouard equilibrium ($C+CO_{2} \leftrightarrow 2CO$) requires such high temperatures for a complete gasification. $CO$ and $H_{2}$ are produced in low quantity at low temperature and $CH_{4}$ is produced in lower quantity compared to steam gasification. However, water is produced below $900^\circ C$. The LHV reaches a maximum value of 24.29 MJ/kg$_{\text{biomass}}$, which is slightly higher than the maximum LHV obtained with steam.

Similar to steam, calculations with $CO_{2}$ in over-stoichiometric proportion do not reach reaction completion at high temperature. After reaching a minimum at $700^\circ C$ the production of water increases with the temperature. At this temperature, the production of $CO$ also exceeds the one reached at the theoretical equilibrium in Figure 2A (55.1 mmol/g$_{\text{biomass}}$) and the $H_{2}$ production decreases. This evolution is due to the reverse water-gas shift reaction ($CO_{2} + H_{2} \leftrightarrow CO + H_{2}O$). The LHV increases very slowly from $750^\circ C$ to $1,500^\circ C$ and reaches a maximum of 25.16 MJ/kg$_{\text{biomass}}$ at $1,500^\circ C$.

**Iron Oxide as Oxidizing Agent**

Figure 3 illustrates the thermodynamic equilibria when using iron oxide as the oxidizing agent (Equation 3) either in stoichiometric proportion or in excess.

In stoichiometric proportion, the theoretical reaction completion is reached at $1,000^\circ C$ and above (at these temperatures, the syngas LHV is 20 MJ/kg$_{\text{biomass}}$). It is noticeable that the production of $CO$ is the same as with steam gasification and the production of $H_{2}$ is the same as with $CO_{2}$ gasification in accordance with reactions (1), (2), and (3). When iron oxide is introduced in over-stoichiometric proportion, the additional oxygen provided to the system results in the production of $CO_{2}$ at high temperature to the detriment of the $CO$ production and syngas LHV. The $CO_{2}$ also reacts with the $H_{2}$ to produce $H_{2}O$ (reverse water-gas shift reaction). This reaction is favored at higher temperature when the production of $H_{2}O$ becomes more
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FIGURE 1 | Quantity of gases produced at thermodynamic equilibrium per gram of dry biomass from steam gasification of wood according to temperature: (A) stoichiometric proportion \( \text{C}_6\text{H}_9\text{O}_4 + 2\text{H}_2\text{O} \); (B) over-stoichiometric proportion \( \text{C}_6\text{H}_9\text{O}_4 + 6\text{H}_2\text{O} \).

FIGURE 2 | Quantity of gases produced at thermodynamic equilibrium per gram of dry biomass from gasification of wood with carbon dioxide according to temperature: (A) stoichiometric proportion \( \text{C}_6\text{H}_9\text{O}_4 + 2\text{CO}_2 \); (B) over-stoichiometric proportion \( \text{C}_6\text{H}_9\text{O}_4 + 6\text{CO}_2 \).

FIGURE 3 | Quantity of gases produced at thermodynamic equilibrium per gram of dry biomass from gasification of wood with iron oxide according to temperature: (A) stoichiometric proportion \( \text{C}_6\text{H}_9\text{O}_4 + \frac{3}{2}\text{Fe}_2\text{O}_3 \); (B) over-stoichiometric proportion \( \text{C}_6\text{H}_9\text{O}_4 + 2\text{Fe}_2\text{O}_3 \).

significant. Figure 4 plots the quantities of iron species (\( \text{Fe}_2\text{O}_3 \), \( \text{Fe}_3\text{O}_4 \), \( \text{FeO} \), Fe) obtained at thermodynamic equilibrium for stoichiometric gasification conditions over iron oxide. Metallic iron (Fe) can be obtained for temperatures beyond 750 °C in the presence of biomass as reducing agent, whereas the direct thermo-reduction of iron oxide requires much higher temperatures (Charvin et al., 2007).

All the thermodynamic calculations reveal that the gasification reaction should be carried out at temperatures beyond 1,000 °C to maximize the biomass conversion. Thus, experiments were
performed at 1,100°C to provide additional information about the kinetics and gas products composition.

**EXPERIMENTAL SECTION**

**Feedstock**

The biomass used for the experiments consisted of beech wood particles whose characteristics are given in Table 1. They were obtained from XP CEN/TS 15104, XP CEN/TS 15105, XP CEN/TS 15107, XP CEN/TS 14775 standards. Oxygen content was deduced by difference. The LHV (dry basis) was measured with a PARR 6200 bomb calorimeter knowing humidity from NF EN 14774 standard. Granulometry was obtained using a FT4 granulometer. The biomass was first dried at 105°C overnight and then kept under vacuum to avoid humidity in the samples.

Regarding iron oxide, a fine powder of ferric oxide was used (iron oxide (III), Fe₂O₃, purity > 99.5%, size < 5 µm). This powder size allowed for an easy mix with the biomass particles with uniform coating at the biomass surface providing a good solid to solid contact. During the preparation of each sample, iron oxide was mixed with biomass in the desired proportion and a 3g sample of the mix was placed in the sample holder. The sample holder was then introduced in the Inconel tube of the experimental bench and a nitrogen flow was injected to prevent the humidification of the sample.

**Experimental Setup**

The PYRATES experimental bench was designed for fast pyrolysis and gasification of small samples using an induction furnace that allows for fast heating rates (500°C/s) (Gauthier et al., 2013). The experimental bench and its components are described in Figure 5.

The 12 kW induction batch furnace (HFP 12, EFRD induction Gmbh) heated an Inconel 600 pipe (33.5 mm diameter) with a four-coil inductor (6 cm height). The temperature at the surface of the Inconel pipe was measured and regulated with a two-color pyrometer (IMPAC infratherm, range 350–1,300°C, $\lambda_1 = 1.52 \mu m$, $\lambda_2 = 1.64 \mu m$, response time 2 ms, precision is $\pm (0.4%, T+1)$) between the two central coils. A quartz pipe provided the sealing of the furnace. A porous metallic sample carrier enabled the gas passage through the biomass particles. A 50°C temperature difference was observed between the wall temperature and the sample after 4 min heating (Gauthier et al., 2013). The evaluation of the sample temperature is considered as the main experimental uncertainty. In the following, as no temperature measurement inside the sample exists, the sample temperature ($T_{wall}$-50°C) was taken as reference. During the heating period, a $N_2$ flow of 2 NL/min [NL means Normal Liter at normal temperature (0°C) and pressure (101,325 Pa)] was maintained at the bottom of the Inconel tube to avoid biomass humidification with ambient air. Gaseous oxidizing agents were injected from the bottom. An additional flow of nitrogen (3 NL/min) was used at the hot zone exit for gas quenching and condensation of condensable gases. These gases condensed on the quartz tube wall (Figure 5, marker 6) and then a collector (Figure 5, marker 7) enabled tar recovery if large quantities were produced. The off gases passed through an electrostatic precipitator, a serpentine ice trap (0°C), two washing bottles inserted in a dry ice and isopropanol bath (−70°C) and then a cartridge filter to retain all condensable gases before analysis.

**Metrology and Gas Analysis**

Gas composition was analyzed continuously using a mass spectrometer and a Fourier Transform InfraRed (FTIR) spectrometer. A gas flow of 1 NL/min was extracted after its passage through the cartridge filter to be analyzed by the mass spectrometer (Adixen ASM1025) measuring only the H₂ concentration with a 1 s measurement interval. The rest of the gas flow was directed toward the FTIR spectrometer ( Nicolet IR 550) that continuously measured the concentrations of CO, CO₂, CH₄, C₂H₆, C₂H₂, CH₃OH, and H₂O. The gas was then stored in a sampling bag (Figure 5, marker 13) to be analyzed after the end of the experiment with a four columns gas-phase micro-chromatograph (Agilent 3000A). The columns A and B (molecular sieve 5 Å, 10m) had a 90°C inlet temperature and used, respectively, argon and helium as carrier gas. Column C (PPU, 8m) and Column D (CP wax 52, 10m) had an inlet temperature of 80°C and 40°C, respectively, and both used helium as carrier gas. The micro-chromatograph was used for the measurement of the concentration of CO, CO₂, CH₄, C₂H₆, C₂H₂, C₆H₆, and C₇H₈. A 13% discrepancy was evaluated on gas mass, based on comparisons between FTIR continuous analysis and micro-chromatograph measurements (the latter being taken as reference).

**RESULTS AND DISCUSSION**

**Effect of Oxidizing Agent**

In order to evaluate the relevance of the biomass gasification over iron oxide, a comparison with pyrolysis along with CO₂ and H₂O gasification was performed. Four runs were carried out at 1,100°C for 20 min with a total bottom gas flow of 1 NL/min to maintain same space times between runs. The pyrolysis run was carried out with 1 NL/min of N₂. In the case of H₂O and CO₂, oxidant injection was continuous. Pure CO₂ (1 NL/min) was injected for CO₂ gasification (about 20
times the stoichiometry over the run period). For H₂O, N₂ was required to limit condensation and thus a mixture of 20% of water and 80% of nitrogen (in volume) with a H₂O flowrate of 0.2 NL/min was injected (corresponding to about 4 times the oxidant stoichiometry over the run period). Finally, a mass fraction of iron oxide of 70% was set in the initial sample for solid-solid gasification (about 2 times the stoichiometry) along with a continuous N₂ flow (1 NL/min). In all cases, the oxidizing agent was in excess even if not in the same proportion; quantitative results and qualitative comparison are commented for each case. Table 2 reports the gas yields and syngas LHV for the four runs. They were obtained from the gas analysis system giving the various concentrations while N₂ tracer gas method was used to determine the total flowrate. LHV was calculated from the given gas composition. All quantities are reported with respect to the initial mass of biomass. The total gas productions (H₂, CO, CO₂, CH₄, and C₂H₄) are reported in Figure 6 while Figure 7 shows the online concentrations of H₂, CO, CO₂, CH₄ over the run duration (20 min). The productions are compared for the various oxidizing agents (H₂O, CO₂, Fe₂O₃) and for pyrolysis. CO₂ production is not reported in the case of CO₂ gasification as it comes not only from the gasification reaction but also from the CO₂ feed.

Results were compliant with the trends observed from thermodynamic calculations. H₂ production was maximum for steam gasification while CO₂ gasification showed the highest amounts of CO. The gas production was lower with iron oxide
TABLE 2 | Total gas quantities produced during gasification and pyrolysis at 1,100°C.

| Oxidizing agent | Gas production [mmol/g Biomass] | LHV (MJ/kg Biomass) |
|-----------------|---------------------------------|---------------------|
|                 | H₂ | CO | CO₂ | CH₄ | C₂H₄ | C₂H₂ | CH₃OH | H₂O | Total |
| CO₂             | 9.93 | 51.58 | – | 3.33 | 0.26 | 0.08 | 0.04 | 0.28 | 65.50 | 21.27 |
| H₂O             | 26.64 | 26.26 | 4.76 | 4.24 | 0.65 | 0.09 | 0.02 | – | 62.66 | 18.8 |
| Fe₂O₃           | 23.49 | 18.61 | 9.07 | 2.10 | 0.01 | 0.02 | 0.02 | 0.32 | 53.64 | 13.06 |
| None (pyrolysis)| 13.12 | 17.54 | 2.93 | 4.16 | 0.58 | 0.08 | 0.02 | 0.18 | 38.60 | 12.72 |

FIGURE 7 | Evolution of the gas production during gasification with CO₂, H₂O, and Fe₂O₃ and pyrolysis of the batch of biomass.

than with other oxidants but still higher than during the pyrolysis run (no oxidizing agent).

Interestingly, for the pyrolysis run, all gases were produced in < 4 min. Thus, for the other runs, any gas production occurring after 4 min should be the result of the char gasification reaction.

Concerning H₂, the production was very similar during the first 4 min for the runs with water, CO₂ or without oxidant. The peak production was about 5.5 mmol/min g⁻¹ Biomass. After 5 min, a weak H₂ production was maintained only for water as oxidizing agent, which came from the char gasification with water that also produced CO and CO₂. For the gasification over iron oxide, the H₂ production was much more pronounced with a peak production at 8.35 mmol/min g⁻¹ Biomass for a total duration of 5 min. However, the total H₂ production from steam gasification was higher but with a lower kinetic.

CO production was maximum for CO₂ gasification but the total produced quantity (51.6 mmol/g Biomass) was still well-below the equilibrium value (65.9 mmol/g Biomass). The first minutes of CO production were very similar between steam gasification and pyrolysis but steam gasification maintained a CO production after 3 min due to progressive char gasification which was not possible without oxidant (pyrolysis). CO production from pyrolysis stopped after 4 min while CO production from steam gasification only slowly decreased after 3 min. Gasification with iron oxide showed a lower CO peak production than the other cases with a total production barely exceeding that of pyrolysis. This result is mainly to be linked with the high production of CO₂ over iron oxide (two to three times the total amount obtained for water gasification and pyrolysis). This could come from the reaction of iron oxide with char (2Fe₂O₃ + 3C → 4Fe + 3CO₂). The high availability of oxygen from Fe₂O₃ was also reported in chemical looping methane reforming, leading also to a CO₂ production peak during the first cycle (Chuayboon et al., 2019).
Concerning CH₄, the gas production occurred during the first 4 min of all runs as methane was produced during the pyrolysis step. Steam gasification and pyrolysis resulted in the highest methane production peak (2.8 mmol/min g₉biomass). For pyrolysis, the large amount of CH₄ is explained by the fact that steam reforming reaction cannot occur (no water injection) while for steam gasification, the short residence time tends to hinder gas phase reactions. The lowest methane production was obtained for the gasification run over iron oxide. Indeed, reduced iron oxide has shown high selectivity toward H₂ and CO in dry reforming (Zhu et al., 2019). This last reaction is favored by the relatively high CO₂ concentration obtained with iron oxide.

As for methane, C₂H₄ and C₂H₆ productions were minimal for gasification over iron oxide. Methanol production remained very small in all cases.

Concerning the LHV of the products, CO₂ and H₂O gasification produced a syngas containing more energy than the initial biomass (18.3 MJ/kg). Solar gasification is a way to upgrade the heating value of the biomass. The Cold Gas Efficiency (CGE) permits to quantify this upgrade. This efficiency is not an energy efficiency and is sometimes called energy Upgrade Factor (F_u) in the field of solar reactors (Zgraggen and Steinfeld, 2008):  

\[ F_u = \frac{LHV_{syngas} \times m_{gas}}{LHV_{biomass} \times m_{biomass}} \] (9)

\( F_u \) superior to 1 means that the produced syngas offers a higher energy content than the initial feedstock. For CO₂ gasification, the \( F_u \) is 1.16; it is 1.03 for steam gasification while the \( F_u \) for pyrolysis is 0.7. Taking into account only the reduction step, the \( F_u \) of the gasification over iron oxide is comparable to pyrolysis (0.71). However, the hydrogen produced through the re-oxidation step should be added (not quantified experimentally) and \( F_u \) superior to one are theoretically achievable.

These results provide qualitative information about the use of iron oxide as oxygen carrier for the gasification reaction (solid-solid reaction) vs. more common solid-gas reactions (steam and dry gasification). The comparison between the two systems (solid-solid and gas-solid) is not obvious because chemical mechanisms are different but the same reactor was used. Especially, the high reactivity observed with iron oxide may be due to the fact that the oxidizing agent is already mixed with the biomass at the beginning of the run, while H₂O and CO₂ have to penetrate the biomass sample. Also, the experimental device is mainly dedicated to the pyrolysis study as the gas residence time in the heated zone is short. Anyway, the feasibility of the solid-solid reaction is demonstrated and it shows promising performance that can possibly be enhanced with continuous particles stirring. Additionally, the effect of the iron oxide proportion in the initial biomass feedstock was studied.

**Influence of the Iron Oxide Stoichiometry**

The influence of the proportion of iron oxide (in default, stoichiometric, excess) was studied for a mass fraction of 0.25 (oxide in default), 0.5 (stoichiometric proportion), and 0.66 (oxide in excess). A run without oxygen (pyrolysis) was also carried out. Samples were heated up to 1,050°C for 10 min. The main conditions and results (initial and final sample masses, gas yield, syngas LHV, carbon conversion) are recapped in Table 3. Carbon conversion is defined as:

\[ X_C = \frac{m_{C_{syngas}}}{m_{C_{biomass}}} \] (10)

Where \( m_{C_{syngas}} \) is the mass of carbon contained in the syngas and \( m_{C_{biomass}} \) the mass of carbon in the initial biomass.

For better readability, Figure 8 plots the H₂, CO, CO₂, and CH₄ yields along with the syngas LHV as a function of iron oxide initial mass fraction in the sample. Concerning the initial and final masses of the samples, when iron oxide mass fraction was 0.5 or 0.66, the final mass of the sample was lower than the initial oxide mass. This proves that oxygen was transferred from

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**TABLE 3 | Gas yields through gasification of biomass with Fe₂O₃ at 1,050°C as a function of the oxidant mass fraction.**

| Fe₂O₃ mass fraction | 0  | 0.25 | 0.5 | 0.66 |
|---------------------|--|--|--|--|
| Sample initial mass (g) | 3.184 | 2.658 | 3.04 | 3.168 |
| Sample final mass(g) | 0.504 | 1.173 | 0.913 | 1.823 |
| Tar mass (g/gbiomass) | 0.211 | 0.14 | 0.233 | 0.176 |
| Gas yield (mmol/gbiomass) | H₂ | 18.716 | 23.043 | 25.878 | 29.584 |
| | CO | 12.997 | 11.210 | 15.110 | 16.084 |
| | CO₂ | 3.639 | 5.813 | 8.163 | 10.180 |
| | CH₄ | 4.011 | 2.560 | 2.431 | 1.534 |
| | H₂O | 0.362 | 0.082 | 0.310 | 0.048 |
| | C₂H₄ | 1.111 | 0.743 | 0.558 | 0.379 |
| | C₂H₆ | 0.061 | 0.022 | 0.009 | 0.005 |
| | CO₂ | 0.312 | 0.282 | 0.188 | 0.169 |
| | C₂H₆ | 0.055 | 0.020 | 0.000 | 0.000 |
| Toluene | 0.000 | 0.088 | 0.077 | 0.084 |
| Methanol | 0.133 | 0.023 | 0.013 | 0.009 |
| Benzene | 0.000 | 0.088 | 0.077 | 0.084 |
| Syngas LHV (MJ/kgbiomass) | 13.82 | 13.04 | 14.30 | 14.55 |
| Carbon conversion rate | 55.08% | 52.76% | 65.03% | 69.11% |

**FIGURE 8 | Gas yield at 1,050°C and LHV of the produced gas according to the Fe₂O₃ mass fraction.**
the oxide to the biomass for the gasification reaction. Conversely, when the iron oxide mass fraction was 0.25, the final mass of the sample (containing unreacted char and iron oxide) was higher than the initial mass of iron oxide denoting a limited reaction extent. The mass fraction of iron oxide had a significant impact on the gas yield. H₂ and CO₂ yields increased clearly (from 18.7 to 29.6 mmol/gbiomass and 3.6 to 10.2 mmol/gbiomass, respectively) while CH₄ yield decreased (from 4.0 to 1.5 mmol/gbiomass) with the increase of the iron oxide mass fraction. The CO yield showed a minimum of 11.2 mmol/gbiomass for an oxide mass fraction of 0.25. The impact on the syngas LHV was finally limited as the minimum value was 13 MJ/kgbiomass and the maximum value was 14.5 MJ/kgbiomass. Noteworthy, the excess of iron oxide did not reduce the syngas LHV as predicted by the thermodynamic equilibrium calculations and the syngas LHV always remained lower than the biomass LHV of 18 MJ/kg. This is to be linked to the low carbon conversion obtained (maximum of 70% for an iron oxide mass fraction of 0.66). While the H₂ yield was close to the theoretical values (25.9 vs. 30 mmol/gbiomass for a stoichiometric blend), the CO yield differed from the calculations (15.1 vs. 42 mmol/gbiomass for a stoichiometric blend). This was also true for the CO₂ yield (8.2 mmol/gbiomass while no CO₂ is predicted by the thermodynamic calculations). Since gas phase reactions were minimized in the experiment (low gas residence time), it is expected that dry reforming reaction (CH₄ + CO₂ → 2CO + 2H₂) is not favored. Also, the excess of oxygen carrier can lead to partial combustion of chars and thus to additional CO₂ production.

X-ray Diffraction analysis (XRD, Panalytical XPert PRO, Cu Kα radiation, αCu = 0.15418 nm, angle range = 10–80°, 2theta) was performed to identify the phase composition of the final solid products. The identification of the crystalline phases was realized by peak indexation and comparison with reference compounds giving standard diffraction peaks (powder diffraction file PDF-2, International Center for Diffraction Data, ICDD). For each run of Table 3, XRD diffractograms are given in Figure 9. Diffraction peak corresponding to metallic iron (JCPDS n° 00-001-1262) or to iron oxides (FeO, JCPDS n° 00-002-1180; Fe₃O₄, JCPDS n° 00-001-1111; Fe₂O₃, JCPDS n° 00-024-0072) were identified. For the iron oxide mass fraction of 25% (biomass in excess) (Figure 9A), Fe was mainly identified with some traces of FeO. This proves that iron oxide was almost totally reduced and thus that particle-particle contacting was effective for an excess of biomass. Given the fact that the proportion of iron oxide was lower than the one required for complete gasification, a total iron oxide reduction was expected and experimentally confirmed. A corresponding low carbon efficiency was obtained (53%) due to the lack of oxidant species. For an oxide mass fraction of 0.5
(Figure 9B), metallic Fe was still the dominant specie in the sample which could again be expected from a stoichiometric blend. Some traces of FeO were also identified. The carbon conversion increased to 65%. Finally, for the highest iron oxide mass fraction (excess of iron oxide, Figure 9C), no trace of Fe$_2$O$_3$ was found but Fe, FeO, and Fe$_3$O$_4$ were present with Fe as the main specie. Indeed, as iron oxide was in excess with respect to reaction stoichiometry, total iron oxide reduction into Fe was not possible. Even if all Fe$_2$O$_3$ particles reacted with the biomass, the impact on the carbon conversion was limited as it reached 69%. During the initial reaction steps, the solid-solid reaction was efficient as a result of good contact between micrometric iron oxide particles and millimetric biomass particles (with the oxide particles spread at the surface of the biomass particles). However, the particles were static and during the reaction progress the contact between particles was progressively reduced leading to limited mass transfer and thus limited carbon conversion rate. Figure 10 shows that the iron and iron oxides particles remaining in the sample holder kept the shape of their surrounding biomass particles on which they were in contact before gasification. Even if a large part of the biomass was gasified, it is possible to distinguish small gray heaps mainly composed of metallic iron which corresponded initially to biomass particles covered by iron oxide. This structure was mainly visible at the top of the sample holder (Figure 10A). The remaining solid residue (char+iron oxide) collapsed at the bottom of the sample holder in the form of a dark gray pellet (Figure 10B). These observations highlight the need for stirring the particles to enhance mass transfer, carbon conversion and ultimately LHV of the syngas.

CONCLUSION

This study focused on the combined biomass gasification and iron oxide reduction for both syngas and iron production. This process can be coupled with concentrated solar energy for renewable fuel production and green iron metallurgy. Original results about beech wood biomass gasification over iron oxide (Fe$_2$O$_3$), H$_2$O, and CO$_2$ were reported. First, a theoretical study was carried out about the thermodynamic equilibrium calculations associated with the different chemical systems depending on the temperature and mass ratio of oxidizing agent. Then, the feasibility of the process was proven in an induction furnace able to reproduce high heating rates representative of concentrating solar installations. Iron oxide was successfully reduced to metallic iron at temperature above 1,000°C concomitantly with the production of hydrogen-rich syngas. A low carbon conversion (<70%) was observed because of mass transfer limitation in the developed fixed bed reactor. The results were compared with dry and steam biomass gasification along with pyrolysis in order to emphasize the remarkable performance obtained with a solid-solid gasification system involving iron oxide as oxygen carrier. In addition, when coupled with concentrated solar energy for supplying process heat, the solar gasification over iron oxide can produce green iron in a sustainable metallurgical process or additional hydrogen via chemical looping in a steam-iron process. Future work should focus on continuous biomass gasification with iron oxide under concentrated solar radiation to demonstrate the feasibility of the novel metallurgical process for high-purity syngas and Fe production. Such a solar process will represent a new promising renewable path toward sustainable solar fuels and chemical commodities.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article suplementary materials, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

QB and MG did the experimental work. SA and SR supervised the work. All co-authors analyzed the results and wrote the paper.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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