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12 ABSTRACT

13 Solar thermochemical fuel production technologies, such as biomass gasification, are confronted to 14 the intermittency of solar irradiance. The development of dynamic simulation tools is thus required 15 to design around-the-clock control strategies. An innovative model was developed here, based on 16 unsteady mass and energy conservation equations, considering gas-phase thermodynamic 17 equilibrium and heterogeneous char oxidation kinetics. The accumulation of char and gas species 18 production rates were therefore tracked throughout operation, giving insight into the reactor 19 dynamics with optimized computational cost. The model was validated via a comparison with 20 experimental results, regarding both thermal and chemical reactor performances. Simulations 21 reliably predicted the evolution of reactor temperatures and syngas production rates, under both 22 solar-only and hybridized (solar-autothermal) operation. Parametric studies regarding the impact of 23 reactants injection rates on steady-state performances were finally proposed. Steam addition (0.22 24 to 0.60 g/min) increased the syngas H_2 :CO molar ratio significantly (1.13 to 1.47). Biomass addition (1 25 to 3 g/min) boosted the solar-to-fuel efficiency (0.22 to 0.47), but altered the reactor temperature. 26 Finally, oxygen addition kept the reactor running despite fluctuations of solar power, while 27 decreasing the total H₂+CO production and cold-gas efficiency linearly. A constant H₂+CO production 28 (2.17 NL/min) could however be achieved by feeding additional biomass and oxygen during 29 hybridization, thus limiting the cold-gas efficiency decrease and improving the reactor energy efficiency (0.29 to 0.40). Such a dynamic reactor model can be further applied to hybridized 30 31 gasification process optimization and dynamic control under real fluctuating solar irradiation 32 conditions.

33 KEYWORDS

34 Solar fuels; Gasification; Hybrid solar reactor; Biomass; Char; Dynamic modelling

35

1. INTRODUCTION

With the aim to develop a green path to produce renewable syngas, a mixture of H_2 and CO for the synthesis of liquid fuels, solar gasifiers have been developed since 1980 [1,2] to carry out the high39 temperature conversion of carbonaceous feedstocks. Conventional autothermal gasifiers use in-situ 40 combustion to supply heat to the endothermal gasification reaction, leading to net feedstock losses 41 and syngas dilution in combustion products (mainly CO_2 and H_2O) [3]. Contrarily, allothermal solar 42 gasifiers enable high chemical efficiencies because all the feedstock can be converted into syngas (no 43 combustion). Numerous reactor designs have already been assessed, including packed-beds, 44 variations of fluidized-bed and spouted-bed technologies, vortex flows, and molten-salt reactors 45 [4,5]. Fluidized beds are promising solutions to ensure a continuous and homogeneous heating of 46 solid particles, but a char throughput must be managed due to incomplete solid conversion [6]. 47 Spouted beds ensure long solid residence times, and efficient solid-gas transfer rates with little 48 spouting gas requirements [7]. The application to woody biomass pyro-gasification naturally 49 emerged for the production of a fully renewable syngas fuel [8]. In several lab-scale experiments, the 50 storage of solar heat into fuels through biomass feedstock upgrading was demonstrated, as cold-gas 51 efficiencies higher than 1 were reported (1.30 in [9], 1.35 in [10]). However, the upscaling towards 52 industrial production is still compromised because of a particularly complex modelling [11] and 53 specific technological challenges. On the one hand, optimizing the particles circulation and the 54 temperatures in the reactor requires sophisticated 3D modelling techniques including radiative 55 transfer [12,13]. On the other hand, optimizing the dynamics of solar gasification through the day 56 requires strong assumptions to reduce computational costs [14,15]. The present work responds to 57 the second problematic, by proposing a model suitable to simulate the dynamic control of a solar 58 gasifier.

59 Indeed, coping with the daily fluctuations of solar power for continuous processing still requires 60 further research work [16]. Yearly solar reactor efficiencies are yet estimated using strong 61 hypotheses (steady-state modelling, hourly time steps...), leading to overestimated reactor 62 efficiencies, wrong reactor dimensioning, and approximate implementation of hybridization 63 strategies [17]. In this context, thermodynamic equilibrium modelling is often employed to assess both autothermal and allothermal gasification mechanisms [18]. As only the initial elemental 64 65 composition of the chemical system is required, it is a convenient and fast alternative to including 66 chemical kinetics [19]. Good agreements can be achieved regarding syngas (H₂, CO and CO₂) 67 production rates, as well as low-temperature (800 °C) char residuals [20]. However, using 68 thermodynamic equilibrium does not enable to describe the dynamics of char oxidation, which rule 69 the accumulation of solid particles throughout operation [21]. The present study aims at improving 70 thermodynamic equilibrium modelling, by considering heterogeneous kinetics of char oxidation and 71 thus describing more accurately the gasifier transients with reasonable computational cost.

72 This new method is applied to the pyro-gasification of beech wood, in a hybrid solar-autothermal 73 spouted-bed reactor [13,22,23]. The ideal reaction of wood gasification is provided in Equation 3. 74 However, the actual mechanism comprises the pyrolysis and devolatilization of wood feedstock 75 followed by the gasification of the solid char product [24]. Methane reforming (Equation 1) and 76 Water-Gas Shift (WGS, Equation 2) reactions rule the distribution of gaseous species during their stay 77 in the cavity. Besides, the *in-situ* injection of oxygen enables to heat the reactor chamber [25–28], 78 despite an altered syngas quality. During such autothermal hybridization, steam and wood injection 79 rates can be tuned to control the syngas quantity and quality [29]. However, the assessment of more 80 sophisticated hybridization strategies (simultaneous control of the three reactants injection rates) 81 cannot be carried out without an efficient dynamic simulation tool.

$CH_4 + H_2O_{(v)} \rightarrow CO + 3 H_2$	ΔH° = +206.1 kJ/mol	(1)

$H_2O_{(v)} + CO \leftrightarrow H_2 + CO_2 \qquad \Delta H^2 = -41.1 \text{ kJ/mol} \qquad (2)$	$H_2O_{(v)} + CO \leftrightarrow H_2 + CO_2$	ΔH° = -41.1 kJ/mol	(2)
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$$CH_{1.45}O_{0.65} + 0.35 H_2O_{(v)} \rightarrow CO + 1.07 H_2$$
 $\Delta H^\circ = +106.3 \text{ kJ/mol}$ (3)

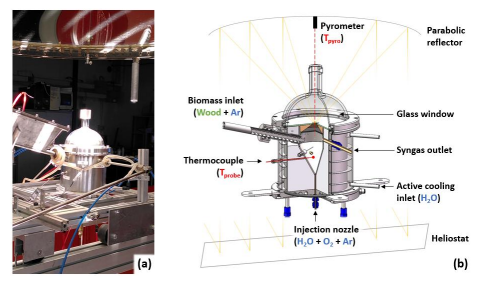
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83 A new dynamic model of a solar hybrid gasification reactor was developed and experimentally 84 validated in this work. To date, such dynamic modelling for the control and operation of hybrid solar 85 gasification systems has not been considered. The model is based on unsteady mass and energy 86 conservation equations, combined with heterogeneous char oxidation kinetics while considering gas-87 phase thermodynamic equilibrium. It is robust and flexible-enough for application to wide operating 88 conditions and reactor scales. It features a cavity receiver where the gaseous phase is assumed to be 89 at thermodynamic equilibrium. Meanwhile, the oxidation of solid char (by O_2 , H_2O and CO_2) is ruled 90 by heterogeneous kinetics featuring Random-Pore Modelling (RPM, [30]). This chemical scheme is 91 coupled to a detailed thermal model, featuring distinct temperatures for the reactor solid parts and 92 the gas phase. The resulting code enables to simulate the main dynamics of woody biomass 93 gasification, under both solar heating and in-situ oxy-combustion. Reasonable computing times are 94 achieved, making the code compatible with annual dynamic simulation and continuous reactor 95 control. After this model overview, a set of experimental validation cases was examined to verify 96 thermal responses and chemical products outcomes. Dynamic simulation revealed a satisfying 97 agreement with the observed thermal and chemical responses of the reactor under both solar and 98 solar-autothermal hybridized operations. Finally, a complete parametric analysis was performed to 99 determine the impact of reactants injection rates (steam, biomass, oxygen) on reactor temperatures, 100 char accumulation, and chemical efficiencies. Such a dynamic model is particularly well suited for an 101 application to reactor optimization and dynamic control. It can be further applied for performance 102 and environmental impact assessments of solar gasification plants under real fluctuating solar 103 irradiation conditions.

104 2. MATERIAL AND METHODS

105 2.1. Solar reactor design

106 The simulation code was conceived to model the operation of a lab-scale spouted-bed solar 107 gasification reactor [22] (Figure 1). Figure 1-a shows the whole reactor casing, topped by a 108 hemispherical glass window. It is set at the focal point of a 1.5 kW beam-down solar furnace. A 1.15 L 109 hopper contains feedstock, that is provided to the cavity through a screw driver. Figure 1-b shows 110 the 0.24 L conical cavity, made of a high-resistant, 3 mm thick FeCrAl alloy. It is surrounded by an insulation layer of 25 mm minimal thickness. Above the cavity, an alumina cap is set with a 20 mm 111 112 wide aperture, to let sunlight enter while minimizing radiative losses. The cap is covered by two 2 113 mm thick layers of zirconia felts, which restrict the aperture diameter (~18 mm) to limit thermal 114 losses. Biomass is injected in the upper half of the cavity, and solid particles are continuously stirred 115 thanks to a jet of oxidizing gases (H_2O , O_2) and argon coming from the lower injection nozzle. The 116 produced syngas is recovered through an alumina tube while outlet Venturi pumping is used to limit 117 overpressure (the cavity pressure is usually maintained around 0.87 bar, the atmospheric pressure at 118 laboratory location being 0.85 bar).



120Figure 1. Experimental setup: (a) Photograph of the solar spouted-bed gasifier during operation, (b) Scheme of the121gasifier and solar concentrator

122 Temperatures in the cavity are measured thanks to several B-type thermocouples and a pyrometer 123 (4.8-5.2 μ m, solar-blind). In particular, one thermocouple (T_{probe}) is shielded in an alumina tube and 124 inserted inside the lower half of the conical cavity. The pyrometer (T_{pyro}) is directed vertically towards 125 the cavity's inside, through a CaF₂ window.

Raw syngas pumped out of the cavity is cooled through a 1 m long steel pipe, visible in the foreground of Figure 1-a. An ice-cooled bubbler then enables to trap steam and to capture most solid residues. Two filters finally collect the remaining solid particles, before the gas stream is analysed and vented. The mole fractions of H₂, CO, CO₂ and CH₄ are measured by an online analyser (GEIT GAS 3100 SYNGAS) with a sampling period of 3 seconds.

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132 2.2. Usual operating conditions

Operation of the reactor can be performed under both direct and indirect heating. In indirect heating, a SiC-coated graphite emitter plate is added under the alumina cap (15 mm below the focal plane aperture). It absorbs concentrated sunlight, and heats the cavity by infra-red radiative transfer. In direct heating, sunlight directly enters the cavity and heats the conical region without any intermediate. A comparison of these two heating modes was already discussed [28], where higher H₂ and CO yields were observed using direct heating, during both solar-only and solar-autothermal hybrid gasification experiments.

Moisture	C (dry)	H (dry)	O (dry)	Ash (dry)	HHV (dry)	LHV (dry)
[wt%]	[wt%]	[wt%]	[wt%]	[wt%]	[kJ/g]	[kJ/g]
9.8	49.8	6.0	43.4	0.6	19.5	18.3

140 Table 1. Composition and heating value of beech wood biomass feedstock (dry basis)

The solar gasification of 1.2 g/min of millimetric beech wood particles (measured characteristics given in Table 1) is usually performed thanks to a 0.2 g/min stream of water injected at the bottom of the conical region [31]. A 0.2 NL/min stream of argon is blown along with water to stir the bulk solid, and 0.5 NL/min of argon is injected in the hopper to avoid counter current heat and mass fluxes along the screw driver. An additional 2.0 NL/min argon stream is directed towards the glass window

147 to protect it, under both direct and indirect heating modes.

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149 **2.3.** Characteristic time analysis

Upon entering the cavity, the feedstock is dried and pyrolyzed. The pyrolysis actually starts in the hot 150 screw driver, but no traces of pyrolysis oils or tars were ever spotted, showing that pyrolysis 151 152 essentially occurs in the cavity at temperatures above 800 °C [32] and produces high yields of gases. 153 The solid char particles then undergo heterogeneous oxidation by oxygen (partial combustion), 154 steam or CO_2 (gasification). The characteristic times for prevailing mechanisms are provided in Table 155 2, considering the reactor temperatures observed with CFD [26] (drying and pyrolysis at 800 °C, gas reactions at 1100 °C, char oxidation at 1300 °C). The kinetics of drying and pyrolysis are taken from 156 157 [33], where they were validated for beech wood particles with granulometries in the range 315-450 158 μm and 1120-1250 μm. As for the char oxidation kinetics, taken from [34], they were determined 159 over char particles of 44 µm average diameter. They were still successfully used to simulate the 160 gasification of larger particles (coal particles of diameters 50, 100, 200 and 500 µm in [35]), and they 161 appear to be a reasonable choice among the kinetics proposed in the literature [36].

Reaction	Rate [E _a in kJ/mol, P in MPa, C in kmol/m ³]	t _{characteristic} [s]	t _{residence} [s]	Source
Drying	dm _{H20} /dt = -5.0·10 ⁸ exp(-50/RT) x m _{H20}	5.4·10 ⁻⁷		[33]
Pyrolysis	$dm_{wood}/dt =$ -7.4·10 ⁴ exp(-70/RT) x m _{wood}	3.5·10 ⁻²		[33]
C + O ₂	$dX_{c}/dt =$ +1.36.10 ⁶ exp(-130/RT) x P ₀₂ ^{0.68} x f(X _c , Ψ =14)	3.8·10 ⁻¹	3.9·10 ⁻¹	[34]
$C + H_2O$	$dX_{c}/dt =$ +2.45.10 ⁷ exp(-214/RT) x P _{H20} ^{0.86} x f(X _c , Ψ =3)	3.1·10 ⁺¹	5.9·10 ⁺¹	[34]
C + CO ₂	$dX_{c}/dt = +6.78 \cdot 10^{4} \exp(-163/RT) \times P_{co2}^{0.73} \times f(X_{c}, \Psi=3)$	1.2·10 ⁺²	2.3·10 ⁺²	[34]
WGS (f)	dC _{H2O} /dt = -2.98·10 ⁸ exp(-163/RT) x C _{H2O} x C _{CO}	3.2·10 ⁺⁰		[26]
Methane reforming	dC _{H2O} /dt = -1.48·10 ¹¹ exp(-260/RT) x C _{H2O} x C _{CH4}	3.2·10 ⁺¹		[26]

162 Table 2. Calculation of characteristic times for the main mechanisms occurring in the gasifier

163

164 The surface function term featured in heterogeneous kinetics is detailed in Equation 4. According to 165 the RPM theory, the oxidation rate of a char particle evolves over time due to structural changes 166 (increase of porosity). This effect is modelled by a function of the particle conversion X_c (Equation 5), 167 that is called surface function. A structure parameter Ψ is also featured. In [34], Ψ equals 14 for 168 combustion and 3 for H_2O and CO_2 gasification. By integrating the surface function until a 99% 169 conversion, a residence time of char t_{residence} can be proposed for the oxidation reactions, as provided 170 in Table 2. It is almost identical to the Arrhenius-based characteristic time in the case of combustion, 171 and it becomes much higher in the case of H_2O or CO_2 gasification (1.9 times the t_{characteristic} values).

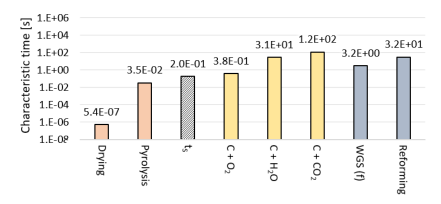
$$f(X) = (1 - X_{C})\sqrt{1 - \Psi \cdot \ln(1 - X_{C})}$$
(4)

$$X_{C}(t) = \frac{m_{C}(t=0) - m_{C}(t)}{m_{C}(t=0)}$$
(5)

172

All characteristic times are displayed in Figure 2, on a logarithmic scale, and compared with the time step that was chosen for process simulation (t_s , 0.2 s). On the one hand, the drying and pyrolysis of woody biomass are fast enough to be considered instantaneous in the model. On the other hand, the 176 combustion of char has a characteristic time near 0.4 s, so the t_s value cannot be higher than 0.2 s for

a reasonable modelling of the process. A shorter time step would make calculations longer, without
 significantly affecting the results. H₂O and CO₂ gasification reactions are much slower than
 combustion, so they will be particularly well modelled.



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Figure 2. Characteristic times of main mechanisms, compared with the simulation time step t_s

Regarding the gas phase reactions, an average residence time can be proposed by considering the total gas output flow rate. The solar gasification of 1.2 g/min of beech wood yields a syngas flow-rate (mostly H_2 and CO) around 1.6 NL/min [29], added to the 2.7 NL/min of argon injected. Under steady state, and considering that the gas temperature is 1100 °C [26], the mean gas residence time therefore lies around 0.7 seconds. The comparison with WGS and reforming characteristic times (respectively 3.2 and 32 seconds) shows that syngas composition is normally ruled by kinetics, in particular regarding the production of CH_4 .

189

190 2.4. Simulation code overview

191 The simulation code developed in this work is named DEMOSTRAR: Dynamic, Extrapolation-ready 192 Modelling and Optimization of a Solar Thermochemical ReActoR. It is written in Python 3, using the 193 library CANTERA [37] specialized in multiphasic thermodynamic calculations. CANTERA was already 194 used in the literature to investigate the thermodynamic mechanisms controlling the gasification of 195 biomass [18,38], and to identify theoretical limitations to the process. It features the thermodynamic 196 properties of numerous species as provided by the NASA [39] and GRI-MECH [40] databases. 197 DEMOSTRAR implements the heterogeneous kinetics of char oxidation in addition to the calculation of gas-phase thermodynamic equilibriums. Thus, the quantity of char present in the reactor at each 198 199 time step can be tracked, which is not the case with only thermodynamic equilibrium calculations. 200 This advance in modelling is brought without strongly affecting the computing costs, which is a key 201 advantage for the implementation of continuous day-night control algorithms [15,41].

A zero-dimensional description of the cavity is proposed to couple the heat and mass balance equations. The metal alloy wall, as well as the emitter plate (when operating under indirect heating), are assumed having uniform temperatures (T_{wall} and T_{plate}). Regarding the reactor inner volume, char particles and gases are heated by both radiative heat transfer (absorbed by char, $H_2O_{(g)}$ and $CO_{2(g)}$), and convective transfer with the walls.

As a result, the model requires around 1 minute to compute 100 minutes of operation, on a dedicated calculation machine.

- 209 The main model assumptions are summed up below:
- a) The pressure in the reactor is perfectly controlled.
- b) Solid parts of the reactor have uniform temperatures.

- c) The gas volume is perfectly stirred (homogeneous temperatures and concentrations).
- d) The gas phase only contains Ar, H₂O, O₂, H₂, CO, CO₂, CH₄ and C₂H₂ species.
- e) The gas phase is considered at thermodynamic equilibrium.
- f) No tars nor pyrolysis oils are modelled (due to high operation temperatures).
- 216 g) Solid char particles and gases share the same temperature.
- h) Solid char particles are evenly dispersed in the reaction chamber.
- i) No solid particles are entrained at the reactor outlet.

219 2.5. Thermal model

Both direct and indirect heating modes were implemented. The main modes of heat transfer considered are detailed in Figure 3. Thermal losses are dispatched between the radiative losses occurring through the upper aperture ($Q_{loss,aperture}$), the radiative losses occurring at the alumina cap upper side despite the zirconia cover felt ($Q_{loss,cap}$), and the conductive losses occurring through the reactor insulation ($Q_{loss,insul}$). The physical properties of the reactor solid parts are summed up in Table 3.

	m [kg]	ρ [kg/m³]	c _P [J/kg·K]	k [W/m·K]	ε[-]
Wall (FeCrAl alloy)	0.4	7100	800	35	0.7
Cap (99.7% Al₂O₃)	0.2	3900	795	9.1	0.8
Emitter plate (SiC covered Graphite)	0.0155	1400	710	25	0.98
Insulation (35% SiO ₂ , 65% Al ₂ O ₃)	0.6	400	800	0.22	-

226 Table 3. Thermal properties of the reactor solid parts (1000 °C)

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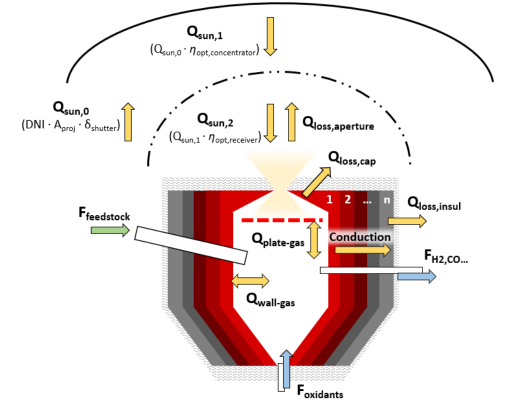




Figure 3. Heat flows of the gasifier modelled in DEMOSTRAR

The calculation of solar power input is detailed in Equation 6 and Equation 7. $Q_{sun,1}$ is the power delivered by the parabolic concentrator. It accounts for the losses occurring at the heliostat and at the parabolic reflector ($\eta_{opt,concentrator}$), and it can be controlled by a shutter obstructer ($\delta_{shutter}$). The value of $\eta_{opt,concentrator}$ was adjusted thanks to calorimetry measurements, so the relationship between the DNI and $Q_{sun,1}$ is well established. As for $Q_{sun,2}$, it accounts for the losses due to both the glass window and the radiation absorption by zirconia felts around the aperture due to spillage ($\eta_{opt,receiver}$). It thus corresponds to the actual solar power available at the cavity entrance.

$$Q_{sun,1} = DNI \cdot A_{proj} \cdot \delta_{shutter} \cdot \eta_{opt,concentrator}$$
(6)

$$Q_{sun,2} = Q_{sun,1} \cdot \eta_{opt,receiver}$$
(7)

237

The total, one-band radiative emissivity of the H₂O and CO₂-rich gas (ε_{gas}) is computed using the 238 correlation of [42] (Equation 8 and Equation 9). This correlation was established to model the heat 239 240 emitted by combustion in industrial furnaces. Based on the partial pressures of both oxidants, it 241 decreases when the gas temperature increases, consistently with conventional emissivity wide-band models. It features an effective beam length L [43,44], that differs according to the origin of 242 243 radiations (walls, emitter plate, or reactor aperture). The total gas absorptivity (α_{gas}) was taken equal 244 to half its emissivity (ε_{gas}), to achieve appropriate thermal responses. Furthermore, the radiative 245 absorptivity of the solid particles cloud (α_{char}) is taken proportional to the projected area of particles 246 across beam trajectories, including a correction regarding particles overlapping effects. No scattering is modelled here, because of restrictive calculation time constraints and incomplete knowledge of 247 248 the char particles size distribution and optical properties. Experimental data would not enable 249 validating such a radiation model, nor fitting an empirical law as proposed in [45]. Under this

approximation, α_{char} reached 3.0%, 6.0% and 11.6% for accumulated char masses equalling 0.5 g, 1 g and 2 g, respectively. The emissivity of solid particles (ε_{char}) was taken equal to their absorptivity.

$$\varepsilon_{\rm gas} = 1 - e^{-\alpha_{\Sigma} \rm PL} \tag{8}$$

$$\alpha_{\Sigma} = ((0.78 + 1.6p_{H_20})/\sqrt{PL} - C) (1 - DT)$$
(9)

252

An unsteady finite-volume conduction model was implemented to simulate the temperature distribution across the wall and insulation thickness. The alumina cap was considered to have the same temperature as the wall first layer, in accordance with CFD results [46]. The total thickness of the wall and insulant parts were 2.74 and 41.4 mm, respectively. The temperature of the inner layer enabled to compute the $Q_{cap-out}$, $Q_{wall-gas}$ and $Q_{wall-out}$ (direct heating) terms. The insulation outer area was assumed to be at 25 °C due to active water cooling (no convection is modelled).

The radiative heat transfer between the outside and the emitter plate (indirect heating) is written in 259 260 Equation 10. As seen in [47], the solar power input Q_{sun,2} is assumed to be entirely captured by the 261 solid, while radiative losses are modelled by the second term. The view factor F_{plate->out} gives the 262 fraction of light emitted by the upper plate surface that reaches the aperture. It equals only 5.5%. 263 The remaining 94.5% are transmitted towards the alumina cap. Besides, the radiative transfer 264 between the wall and the gas + particles phase is written in Equation 11. The correction of the wall 265 emissivity was proposed in [43], because the enclosure radiates towards itself and thus tends to 266 behave like a black body. A constant convective heat transfer coefficient of 50 W/m²K was applied, 267 that is a typical value for spouted beds [48]. The radiative losses attributed to the alumina cap are 268 given in Equation 12. The first term features the cap inner area A_{cap} and its view factor with the 269 aperture (only 1.0%, thus negligible with this geometry). The second term features an apparent 270 radiative area A_{app} that is adjusted to model radiative losses occurring despite the zirconia felts 271 covering.

$$Q_{\text{out-plate}} = Q_{\text{sun},2} - \sigma A_{\text{plate}} F_{\text{plate} \to \text{out}} \varepsilon_{\text{plate}} \left(T_{\text{plate}}^4 - T_{\text{out}}^4 \right)$$
(10)

$$Q_{\text{wall-gas}} = \sigma A_{\text{wall}} \frac{\varepsilon_{\text{wall}} + 1}{2} \left(\alpha_{\text{gas}} T_{\text{wall}}^4 - \varepsilon_{\text{gas}} T_{\text{gas}}^4 \right) + H_{\text{conv}} (T_{\text{wall}} - T_{\text{gas}})$$
(11)

$$Q_{cap-out} = \sigma A_{cap} F_{cap\toout} \varepsilon_{cap} (T_{wall}^4 - T_{out}^4) + \sigma A_{app} (T_{wall}^4 - T_{out}^4)$$
(12)

$$m_i c_{P_i} \frac{T_i^{t+\Delta t} - T_i^t}{\Delta t} = \sum_j Q_{j \leftrightarrow i}^{t+\Delta t}$$
(13)

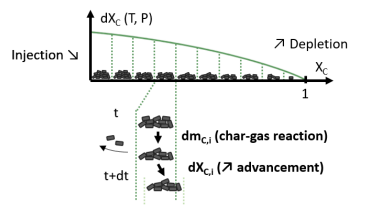
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Equation 13 illustrates the heat balance condition that is computed at each time step. All the heat transfer terms are expressed at the instant $t+\Delta t$ instead of the instant t, according to the principle of implicit resolution. This method ensures the stability of calculations, whereas an explicit resolution scheme would impose using a much lower time step duration. The heat balances written in Equation 13 thus constitute a system of non-linear equations that is solved using Newton-like root search methods [49].

279 **2.6.** Chemical model

As discussed in 2.3, the feedstock drying and devolatilization are considered instantaneous upon injection. Drying is modelled using the enthalpy of vaporization of water [33]. Then, the dry feedstock particles are split into a mass of char and a mixture of pyrolysis gases, which are brought to thermodynamic equilibrium with the other gases in the reactor. Therefore, only the elemental composition (C, H, O) of the feedstock is required to model this step. The mass of char produced is given by the yield Y_{char}, defined according to the pyrolysis conditions in the reactor. As aggregated in 286 [33], the pyrolysis of beech wood particles occuring between 900 and 1400 °C may produce between

287 $2\%_{wt}$ and $6\%_{wt}$ of char, and a pyrolysis occuring at lower temperatures can produce more than $10\%_{wt}$ 288 of char.



289 290

Figure 4. Principle of char particles dynamic classification according to their conversion

291 During operation, the char particles circulating in the cavity are dynamically classified according to 292 their respective conversion. This is necessary to model oxidation reactions using RPM, as already 293 explained in 2.3. To that extent, the particles conversion range (from 0 to 1) is segmented into ten 294 classes of equal length (Figure 4). When a new mass of char is injected, it is added to the first class 295 (conversion 0-0.1). Then, at each time step, the masses of the ten classes $m_{C,i}$ are updated because of 296 heterogeneous oxidation reactions. Ten distinct conversion rates dX_{ci} are calculated, corresponding 297 to ten mass losses dm_{ci}. These masses are subtracted from the respective m_{ci}, and the char 298 populations are partly displaced towards higher conversions according to the dX_{ci} values. Finally, 299 when particles reach the last class (conversion 0.9-1), they are integrally converted by H_2O , CO_2 and 300 O₂. A sensitivity analysis showed that increasing the number of classes did not improve the results 301 significantly.

302 3. VALIDATION OF THE SIMULATION CODE

303 **3.1**. Validation of the thermal model

Two thermal validation cases to compare the reactor thermal response with experimental data were proposed. The temperature responses of the reactor to several solar power steps were determined, in both direct and indirect heating modes, to adjust the thermal model parameters under argon heating. Two parameters were fitted, including the optical efficiency $\eta_{opt,receiver}$ (Equation 7) and the apparent area of the cap A_{app} radiating towards the outside (Equation 12).

309 3.1.1. Direct heating mode

Under direct heating, four consecutive solar power steps were performed (Q_{sun.1} equalled 310 successively 330, 610, 950, and 1035 W), and the corresponding T_{probe} and T_{pyro} responses were 311 312 plotted (Figure 5, T_{probe} and T_{pyro} lowest observable values are 200 °C and 500 °C, respectively). The 313 pyrometer pointed towards the conical region, showing temperatures 100 °C lower than the 314 thermocouple measurement during reactor heating. Actually, T_{pvro} raw measurements did not consider any surface texture, slope and emissivity, so they might be slightly under-estimated. In 315 comparison, the probe provided a representative measurement of T_{wall}, due to irradiation by the 316 317 walls during operation.

A good agreement was achieved between the simulation temperature T_{wall} (sim) and the probe measurement T_{probe} (exp), during the reactor heating and cooling. Increasing the $\eta_{opt,receiver}$ parameter tended to increase all the temperatures because of a higher solar power input, while increasing the

- 321 A_{app} value tended to decrease the highest temperatures only because of higher radiative losses.
- 322 $\eta_{\text{opt,receiver}}$ was eventually fixed to 0.65, and A_{app} to 7.0 cm², accounting for 13% of the total alumina 323 cap upper area. The difference between T_{wall} and T_{gas} in the simulation reached up to 70 °C at high
- 324 solar power input, because of the low residence time of argon in the cavity (~1 second).

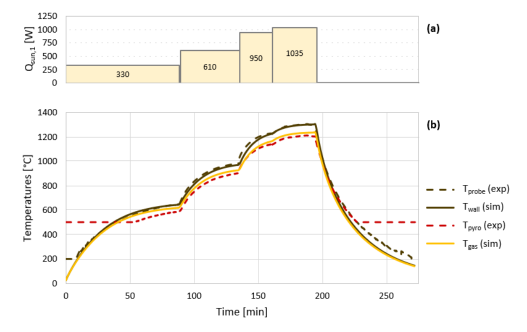


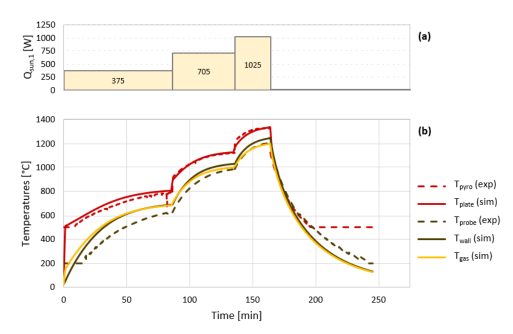


Figure 5. (a) Solar power inputs, (b) Comparison of simulation (sim) and experimental (exp) thermal responses under argon heating (direct heating)

328 3.1.2. Indirect heating mode

Under indirect heating, three consecutive solar power steps were performed ($Q_{sun,1}$ equalled successively 375, 705, and 1025 W), and the corresponding temperatures responses were plotted (Figure 6). In this case, T_{pyro} was systematically much higher than T_{probe} (120 to 350 °C of difference during reactor heating), because the pyrometer pointed towards the centre of the hot emitter plate. Higher thermal losses occurred because of this hot wall placed near the reactor aperture, thus a higher solar power input was required to let T_{probe} rise beyond 1200 °C (1025 W, versus 950 W in direct heating).

The satisfying agreement between experimental and simulation data, obtained while keeping the same parameters as in Figure 5, showed the robustness of the zero-dimensional thermal model. A higher gap was however noticed between T_{wall} (sim) and T_{probe} (exp) than in Figure 5 (it reached 120 °C at low temperatures). T_{gas} (sim) was higher than T_{wall} (sim) at low temperatures due to the high contribution of the emitter plate in the heating of the reactor chamber (T_{plate} is much higher than T_{wall} during the first 60 minutes).





343Figure 6. (a) Solar power inputs, (b) Comparison of simulation (sim) and experimental (exp) thermal responses under344argon heating (indirect heating)

345 **3.2.** Validation of solar gasification dynamics

A second set of validation cases was proposed to validate the dynamics of solar gasification (Figures 346 347 7-9). The operating conditions are summed up in Table 4. Different heating modes, feedstock compositions and steam to biomass (S/B) stoichiometries were assessed. All series were carried out 348 349 under a constant T_{probe} temperature (1300 °C), thanks to the controlled aperture of the shutter 350 obstructer. During the first part of the experiments (17-18 minutes), feedstock and steam were 351 injected, producing a H_2 and CO-rich syngas. Then, during the second part, no feedstock was injected 352 anymore and steam was maintained to gasify the remaining particles of char. The mass of char 353 accumulated during the first part of the experiments was deduced by integrating the flow rates of 354 CO_{2} and CH_{4} produced during the second part.

	Heating	Feedstock	Feedstock rate [g/min]	H₂O rate [g/min]	(S/B)/(S/B) _{st}
Case 1	Indirect	Biochar	0.23	0.39	1.13
Case 2	Indirect	Biochar	0.23	0.49	1.42
Case 3	Direct	Beech wood	1.2	0.2	1.13

355 Table 4. Operating conditions for biochar and beech wood dynamic gasification runs (Figure 7 to Figure 9)

356

357 3.2.1. Gasification of biochar

In Figure 7, the conversion of biochar was performed to validate the char gasification kinetics while minimizing the impact of devolatilization. This biochar (produced by the UK Biochar Research Centre, measured composition given in Table 5) resulted from the pyrolysis of soft-wood pellets in a 700 °C rotary kiln. The brittle centimetric cylindrical pellets were directly grinded by the screw driver before entering the cavity.

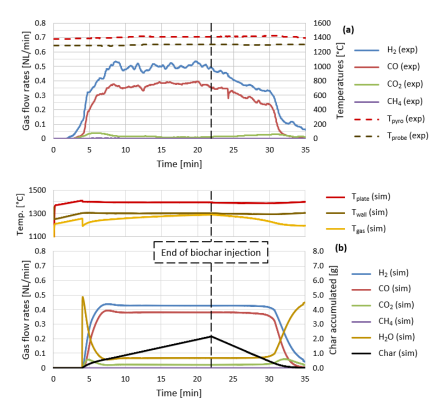




Figure 7. (a) Experimental products outcomes and temperatures, (b) Simulation products outcomes and temperatures,
 during the gasification of biochar (indirect heating). F_{biochar} = 0.23 g/min, F_{steam} = 0.39 g/min, T_{probe} = 1300 °C

366 The experiments showed that T_{probe} and T_{pyro} temperatures were successfully maintained at 1300 and 367 1400 °C, respectively. Under such conditions, the production of H_2 and CO started plateauing around time = 10 min (reaching around 0.50 and 0.38 NL/min, respectively). A H₂:CO molar ratio of 1.32 was 368 369 reached, resulting from the WGS reaction. Meanwhile, the CO₂ production stabilized around 0.01 370 NL/min, after peaking to 0.04 NL/min during the initial transient period. After the interruption of 371 biochar injection, the H_2 and CO production rates decreased by one third between time = 22 and 30 372 min, and finally dropped as char consumption was nearly complete. The CO flow rate became 373 neglectable around time = 34 min, and 10 more minutes were required to reach the end of H_2 374 production. During this transient, the CO_2 production peaked at 0.03 NL/min (time = 31 min) because of the higher fraction of H₂O available for WGS. 375

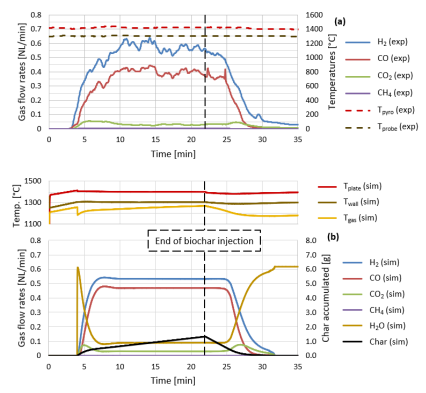
376 Regarding simulation results, the wall temperature stabilized well at 1300 °C. T_{plate} stabilized at 1400 °C, in accordance with experimental results. T_{gas} increased strongly during the first 15 minutes of 377 378 gasification and stabilized around 1300 °C. It then decreased until 1200 °C when biochar injection 379 was stopped, as the solar power input was decreased to maintain T_{wall} equal to 1300 °C. Compared with experimental data, the H₂ and CO flow rates stabilized somewhat more quickly (after 5 380 381 minutes). The H_2 :CO molar ratio equalled only 1.12, which is the value given by gas-phase 382 thermodynamic equilibrium. The accumulation of char in the cavity (black curve) increased until 2.16 383 g, and then decreased gradually after the char injection stopped. This mass was well validated by the 384 experiment, as a 1.9 g accumulation (±0.3 g, uncertainty regarding the final carbon mass balance) was determined experimentally using a mass balance on carbon (gas phase) after biomass injection. 385 386 Besides, the time necessary to gasify the char residue was similar in the experimental and simulation 387 curves, which validated the kinetics implemented.

388 Table 5. Composition of biochar pellets feedstock (dry basis)

Moisture	C (dry)	H (dry)	O (dry)	Ash (dry)
[wt%]	[wt%]	[wt%]	[wt%]	[wt%]

	1.0	90.2	1.8	6.0	1.9	
389						

Figure 8 shows the effect of a higher steam flow-rate on syngas production rates. During steady 390 state, the experiment showed a higher H_2 :CO molar ratio (1.38) because of the enhanced WGS 391 392 reaction, and a higher CO_2 production rate (0.03 NL/min) than in Figure 7. The simulation showed 393 that only 1.32 g of char was accumulated in the cavity, while a 1.0 g accumulation (±0.1 g) was determined experimentally. Less time was therefore required to gasify the residue during the second 394 395 part of the experiment than in Figure 7. Once again, a satisfying agreement was found between experimental and simulation data, and the impact of steam stoichiometry was well represented by 396 397 the model.



398

Figure 8. (a) Experimental products outcomes and temperatures, (b) Simulation products outcomes and temperatures,
 during the gasification of biochar (indirect heating). F_{biochar} = 0.23 g/min, F_{steam} = 0.49 g/min, T_{probe} = 1300 °C

401 3.2.2. Gasification of beech wood

The pyro-gasification of beech wood was then performed (Figure 9). This new validation case enabled validating the devolatilization mechanism implemented in DEMOSTRAR. Three different char yields (8%_{wt}, 12%_{wt} and 16%_{wt}) were assessed in the simulation results, corresponding to decreasing pyrolysis temperature conditions (see Section 2.6). Different pyrolysis temperatures could indeed be encountered, caused by the feedstock progressive heating in the screw driver.

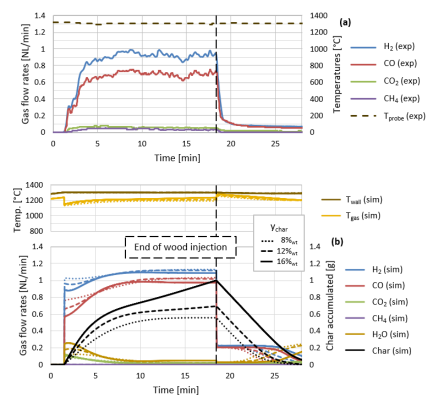




Figure 9. (a) Experimental products outcomes and temperatures, (b) Simulation products outcomes and temperatures,
 during the gasification of beech wood (direct heating). F_{wood} = 1.2 g/min, F_{steam} = 0.2 g/min, T_{probe} = 1300 °C

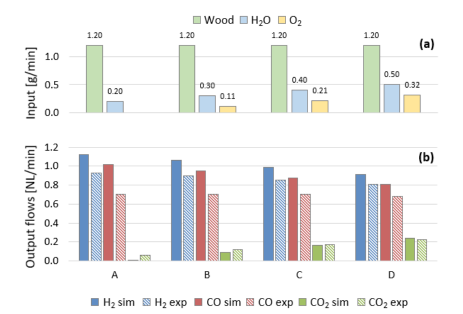
Experimental data showed the establishment of a stable syngas production regime after ~5 minutes of injection (~0.95 NL/min H₂ and 0.70 NL/min CO). When wood injection stopped, a prompt decrease of syngas production was observed as no pyrolysis took place anymore. The remaining char was gasified slowly, over more than 20 minutes. In comparison, simulation results showed the establishment of higher production rates of H₂ and CO (~1.1 and 1.0 NL/min, respectively) and lower production rates of CO₂ and CH₄. This was mainly due to the thermodynamic equilibrium hypothesis.

416 According to the simulation, the chemical and thermal transients were strongly affected by the yield 417 of char. Higher yields caused increasing char accumulations, and favoured the heating of the reactor 418 content due to higher particles radiative absorptivity. At equilibrium, T_{gas} equalled 1191, 1210 and 419 1234 °C at $Y_{char} = 8\%_{wt}$, 12%_{wt} and 16%_{wt}, respectively. Besides, higher yields led to a steeper 420 progression of H₂ and CO production rates. When the wood injection was stopped, the accumulation 421 of char equalled 0.56, 0.69 and 1.00 g at $Y_{char} = 8\%_{wt}$, 12%_{wt} and 16%_{wt}, respectively. In comparison, 422 the accumulation determined experimentally was 0.71 g (±0.08 g).

423 This comparison highlighted the role of the screw driver, that led to pyrolysis starting at 424 temperatures lower than T_{gas} . In the following, Y_{char} will be set to $12\%_{wt}$, that is the value better 425 validated by experimental data for this reactor design.

426 3.3. Validation of hybridized gasification results

A validation of the model was performed with beech wood pyro-gasification experimental results, in the case of hybridization with *in-situ* oxygen injection [29]. The aim was to control the syngas quality (H₂:CO ratio) during hybridization by conjointly increasing the steam injection rate. The results were obtained under direct solar heating, with T_{probe} being constantly maintained at 1300 °C. The flow rates of reactants and the comparison between experimental (exp) and simulation (sim) results are provided in the bar charts of Figure 10.





434 435

Figure 10. (a) Input flow rates, (b) Comparison of simulation (sim) and experimental (exp) syngas compositions, during the hybridized gasification of beech wood (direct heating, T_{probe} = 1300 °C)

436 Figure 10 shows a fair agreement between experimental and simulation data. The relative 437 discrepancy regarding H_2 , CO and CO_2 production rates was lower than 31%, with the exception of CO₂ production during run A. Both the H₂ and CO productions decreased at a comparable rate during 438 hybridization, while the production of CO₂ increased progressively because of combustion reactions. 439 440 The H₂:CO molar ratio was maintained near its initial value thanks to the injection of steam, 441 according to both the simulation data (slightly increasing from 1.10 to 1.13) and the experimental 442 data (slightly decreasing from 1.32 to 1.19). Actually, it would have decreased drastically if no steam 443 had been supplied [29]. The coupled impact of oxygen and steam injection rates was thus well 444 represented by the DEMOSTRAR model, despite an over-estimation of H₂ and CO flow rates due to 445 mainly the thermodynamic equilibrium hypothesis and to the imprecise wood injection rates over 446 long experiments [29].

447 **4. PARAMETRIC ANALYSIS**

448 After the validation step, the model was employed to assess the impact of reactants flow rates on 449 reactor temperatures, char accumulation, chemical species outputs and efficiencies.

450 **4.1.** Impact of steam input flow rate

The impact of steam injection rate on the reactor products outcome was investigated. Figure 11 illustrates the char accumulation dynamics under several steam flow rates. The input solar power was maintained constant ($Q_{sun,1} = 1200$ W, direct heating), as the gasification of 1.2 g/min of beech wood was simulated. At this feeding rate, a H₂O flow-rate of 0.165 g/min was theoretically required to achieve stoichiometry. Two kinds of responses were actually observed:

- At $F_{H2O} \le 0.2$ g/min, the quantity of char accumulated in the cavity diverged over time.
- 457 At $F_{H2O} \ge 0.3$ g/min, the quantity of char accumulated converged after 30 minutes.

458

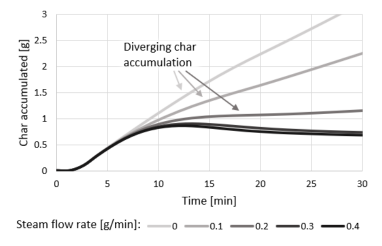
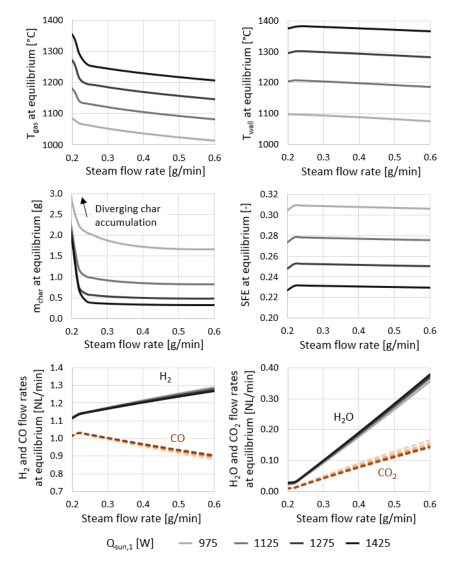


Figure 11. Char accumulation dynamics, for different steam injection rates (0-0.4 g/min). F_{wood} = 1.2 g/min, Q_{sun,1} = 1200
 W

459

462 Figure 12 illustrates more in details the impact of steam injection rate on the reactor temperatures, 463 the accumulation of char, and the reactor chemical performances. Simulations were performed with a wood feeding rate of 1.2 g/min, and solar power inputs Q_{sun,1} ranging from 975 to 1425 W. The 464 maximal steam injection rate was 0.6 g/min. All the simulated data provided here were collected 465 466 when steady state was reached, regarding both the temperatures and the accumulation of char. The 467 minimal steam flow rate ensuring char mass convergence was ~0.22 g/min, that was 1.33 times the 468 theoretical steam-gasification stoichiometry (0.165 g/min), because of kinetic limitations and short 469 steam residence times.



471Figure 12. Impact of steam flow rate on reactor temperatures, char accumulation, SFE and chemical outcome, for472different solar power inputs (Q_{sun.1}, 975-1425 W). F_{wood} = 1.2 g/min

Increasing the steam input above 0.22 g/min resulted in T_{gas} decreasing by ~100 °C, and T_{wall} 473 decreasing by ~25 °C because of higher heating requirements. Char accumulation decreased, 474 475 especially at the lowest solar power inputs (from 2.1 to 1.7 g), due to favoured steam-gasification 476 kinetics. The production of H₂+CO remained constant (2.17 NL/min), while the molar H₂:CO ratio increased strongly because of enhanced WGS (from 1.13 to 1.47 at the lowest Q_{sun.1}), as observed in 477 478 [29]. The Solar-to-Fuel Efficiency (SFE, Equation 14) decreased very slightly, as the molar LHV of H_2 479 (240.2 kJ/mol) is lower than the molar LHV of CO (282.8 kJ/mol). Meanwhile, the H_2O and CO_2 480 outputs increased linearly.

$$SFE = \frac{LHV_{H_2} \cdot \dot{m}_{H_2} + LHV_{CO} \cdot \dot{m}_{CO} + LHV_{CH_4} \cdot \dot{m}_{CH_4}}{Q_{sun,1} + LHV_{wood} \cdot \dot{m}_{wood}}$$
(14)

481

470

Increasing the solar power input naturally led to higher temperatures, and lower char accumulation
 and SFE. High solar power inputs caused the reactor walls temperature to approach 1400 °C, which
 would degrade the FeCrAl alloy constituting the reactor wall. Contrarily, low solar power inputs
 substantially increased the SFE, while decreasing T_{gas} and increasing the accumulation of char.

486 4.2. Impact of biomass input flow rate

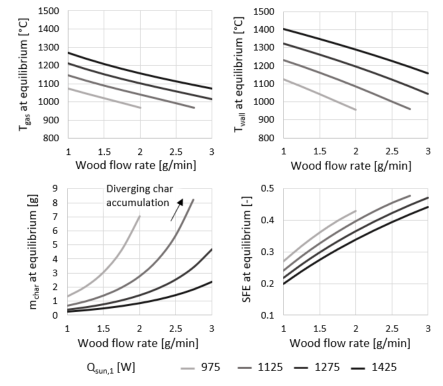
The impact of wood feeding rate (1 to 3 g/min) on the reactor temperatures, char accumulation and SFE was studied (Figure 13), under solar power inputs in the range 975-1425 W. The mass ratio between steam and injected wood (wet) was maintained equal to 0.25.

490 Because of the endothermal reaction, both the gas and wall temperatures decreased, as wood was 491 injected, by 200 to 250 °C. Converting more wood yielded a higher syngas output. The SFE thus 492 increased from 0.22 to 0.47, following the path where $Q_{sun,1} = 1275 \text{ W/m}^2$. Finally, the gasification of

493 more wood led to higher char accumulation in the reactor, that could go beyond 5 g at the lowest

494 Q_{sun,1}. The mass of char eventually diverged over time, when too much wood was provided at a given

495 $Q_{sun,1}$, so the curves of Figure 13 had to be interrupted whenever steady state could not be reached.



496

497Figure 13. Impact of wood flow rate on reactor temperatures, char accumulation and SFE, for different solar power498inputs (Q_{sun,1}, 975-1425 W). Steam/wood_(wet) = 0.25 g/g

499 Once again, increasing the solar power input increased the reactor temperatures, and thus 500 decreased the accumulation of char. Maintaining the wall temperature at 1300 °C while injecting 501 more than 2.0 g/min of biomass would require to increase $Q_{sun,1}$ above 1425 W, which is prevented 502 due to solar concentrator limitations.

4.3. Impact of oxygen input flow rate

The impact of oxygen addition in the reactor was finally assessed. The initial case was the allothermal steam gasification of 1.2 g/min of wood, under a solar power input $Q_{sun,1}$ of 1125 W. From this point, injecting oxygen alone would have caused temperatures to reach dramatically high values. Instead, the addition of oxygen was associated with a proportional decrease of the solar power input.

508 4.3.1. Hybridization at constant wood flow rate

509 In Figure 14, the oxygen flow rate was raised from 0 to 1.1 NL/min (Equivalence Ratio defined in 510 Equation 15, ER = 0.97), while $Q_{sun,1}$ was decreased from 1125 to 0 W (-102.3 W per 0.1 NL/min of O_2 511 provided). The wood flow rate was maintained constant. These specific conditions enable to range

512 from the solar gasification of biomass to its integral combustion, while maintaining a relatively stable

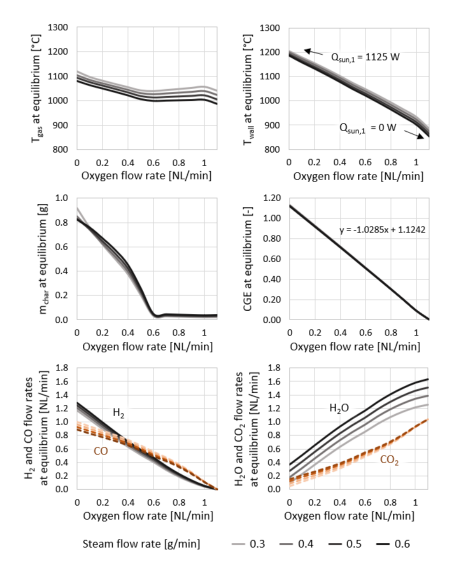
 T_{gas} temperature. In a first domain (F_{O2} < 0.6 NL/min), char accumulation was reduced because of 513 combustion until 30 mg. The gas temperature decreased down to 1000°C, as oxy-combustion did not 514 counteract the decreasing solar power input. In a second domain ($F_{02} > 0.6$ NL/min), T_{gas} increased, 515 516 reaching up to 1042 °C during autothermal operation. Besides, during the entire hybridization path, 517 the wall temperature decreased despite combustion occurring in the cavity. It went below T_{gas} 518 around F_{02} = 0.65 NL/min, and reached down to 854 °C during autothermal operation. Both the H₂ 519 and CO production rates decreased strongly because of oxy-combustion, with a declining H₂:CO 520 molar ratio, until they reached zero at F_{02} = 1.1 NL/min. Increasing amounts of CO₂ and H₂O were 521 produced due to combustion reactions. As a result, the Cold-Gas Efficiency (CGE, Equation 16) 522 decreased linearly with the oxygen input (-0.10 per 0.1 NL/min of O_2 provided), in agreement with 523 experimental results [28] (proportional relation between the O_2 :C ratio increase and the CGE 524 decrease). A CGE higher than 1 (1.12 in allothermal gasification) indicated that the fuel heat content 525 was upgraded, storing a part of the incident solar energy. A CGE lower than 1 indicated that a 526 downgrade occurred, especially because gases with high LHV were combusted.

$$ER = (n_{biomass}/n_{02})/(n_{biomass}/n_{02})_{combustion\ stoichiometry}$$
(15)

$$CGE = \frac{LHV_{H_2} \cdot \dot{m}_{H_2} + LHV_{CO} \cdot \dot{m}_{CO} + LHV_{CH_4} \cdot \dot{m}_{CH_4}}{LHV_{wood} \cdot \dot{m}_{wood}}$$
(16)

527

Increasing the steam injection rate from 0.3 to 0.6 g/min did not significantly affect the CGE. It decreased T_{gas} by 39 to 55 °C (allothermal and autothermal operation, respectively), and T_{wall} by 17 to 32 °C (allothermal and autothermal operation, respectively). It also decreased the mass of char accumulated, only under allothermal operation. Obviously, increasing the injection of steam resulted in much higher H₂O outputs, and also higher CO₂ outputs near allothermal operation because of the WGS reaction.

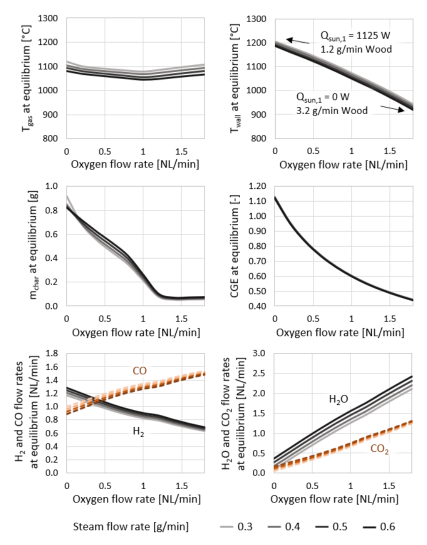


535

536Figure 14. Impact of oxygen flow rate on reactor temperatures, char accumulation, CGE and chemical outcome, for537different steam inputs (0.3-0.6 g/min) under decreasing solar power input (Q_{sun,1}, 1125-0 W). F_{wood} = 1.2 g/min

538 4.3.2. Hybridization at increasing wood flow rate

539 A similar parametric study was performed in Figure 15, but the decrease of H₂ and CO was compensated by a gradual addition of biomass during hybridization. The wood flow rate was 540 541 increased from 1.2 to 3.2 g/min, while the oxygen flow rate was increased from 0 to 1.8 NL/min (ER = 1.58). The solar power Q_{sun,1} kept on decreasing gradually from 1125 to 0 W. In this case, the gas 542 temperature remained between 1045 and 1120 °C, while T_{wall} decreased down to 920 °C. The H₂+CO 543 544 volume flow rate was successfully maintained constant (2.17 NL/min) despite hybridization, at the cost of a much higher production of H_2O and CO_2 . In this case, the CGE decreased less than in Figure 545 14, as its minimal value (autothermal operation) reached 0.44. 546



549Figure 15. Impact of oxygen flow rate on reactor temperatures, char accumulation, CGE and chemical outcome, for550different steam inputs (0.3-0.6 g/min), under decreasing solar power input (Q_{sun,1}, 1125-0 W) and increasing wood input551(1.2-3.2 g/min)

552 Thermal and mass balances were proposed in Figure 16, corresponding to the hybridization studied 553 in Figure 15 with F_{H2O} = 0.4 g/min. Four oxygen flow rates from 0 to 1.8 NL/min were assessed, with 554 decreasing solar power inputs and increasing wood feeding rates. In Figure 16-a, the sum of the solar 555 power input Q_{sun.2} and the heat delivered by combustion was provided. Against it, the sum of the 556 useful heat, as well as the radiative (rad) and conductive (cnd) heat losses was plotted. The total heat 557 required to operate the reactor decreased through hybridization, because of the decreasing heat 558 losses occurring at the wall. The reactor efficiency (useful heat over total heat source) thus increased 559 from 0.29 to 0.40. Regarding the mass balances (Figure 16-b), the increase of O_2 and wood flow rates 560 caused an increase of all the output flow rates excepting hydrogen. The CH₄ output was negligible, as 561 well as the C_2H_2 output, whatever the heating modes and operating conditions.

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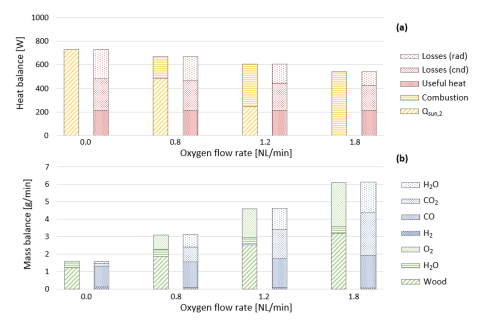




Figure 16. (a) Heat and (b) Mass balances, following the hybridization path of Figure 15. F_{H2O} = 0.4 g/min

564 **5. CONCLUSIONS**

565 A dynamic simulation code was developed to model a solar-autothermal hybrid gasifier. This code assumed that the gas phase was at thermodynamic equilibrium, while the oxidation of solid char 566 particles was ruled by detailed heterogeneous kinetics to describe the accumulation of char during 567 568 transients. This improvement enabled to unravel key dynamic phenomena regarding woody biomass 569 gasification, at a limited calculation cost (1 minute required to simulate 100 minutes of operation, 570 with a time step of 0.2 seconds). Simulations reliably predicted the reactor thermal response up to 571 1300 °C under both direct and indirect solar heating, and syngas production rates under both solar 572 and solar-autothermal hybridized operations.

- The thermal validation against experimental data was successful, showing wall temperature discrepancies below 50 °C under direct heating and 120 °C under indirect heating (the maximal temperature assessed being 1300 °C).
- The validation of solar-autothermal hybrid gasification was satisfying, with a syngas flow rate
 discrepancy explained by the gas-phase thermodynamic equilibrium assumption.
- Char accumulation usually converged after 30 minutes, when the steam input was higher 579 than 1.33 times the theoretical stoichiometry, due to kinetic limitations in the reactor.
- Providing excess steam (0.22-0.60 g/min) resulted in improved H₂:CO molar ratios (1.13-1.47 at Q_{sun,1} = 975 W), and did not decrease the H₂+CO production.
- Increasing the wood feeding rate (1-3 g/min) doubled the SFE (0.22-0.47 at Q_{sun,1} = 1275 W), but altered the reactor temperatures and caused booming char accumulation.
- Switching between solar and autothermal operation was possible, whether by keeping the wood feeding rate constant (equivalence ratio reaching 0.97, *i.e.*, total combustion of feedstock) or by increasing it to maintain a constant production of H₂+CO (equivalence ratio reaching 1.58).

The DEMOSTRAR model is therefore suitable for an application to reactor optimization and dynamic control. Accordingly, it will be specifically applied to the control of syngas production output based on various control variables and optimization schemes to identify suitable strategies for process hybridization. The main optimization goals will be to minimize the consumption of both oxygen and solid feedstock as well as the CO_2 emissions, while achieving maximum or stable syngas (H₂+CO) 593 production under variable (daily fluctuations) and intermittent solar energy input for 24/7 594 continuous operation. Further work will address the implementation of alternative hybridization 595 pathways, considering e.g., concentrated solar resource combined with external combustion, 596 electrical heating or thermal energy storage. Annual performance and environmental impact 597 assessments of solar gasification plants will finally be performed.

598 NOMENCLATURE

599 Abbreviations

- 600 CGE Cold-Gas Efficiency
- DEMOSTRAR Dynamic, Extrapolation-ready Modelling and Optimization of a Solar
 Thermochemical ReActoR
- 603 DNI Direct Normal Irradiance (W/m²)
- 604 RPM Random-Pore Model
- 605 SFE Solar-to-Fuel Efficiency
- 606 WGS Water-Gas Shift

607 **Physical variables**

608	٠	$lpha_{ ext{char}}$	[-]	Volumetric absorptivity of light by char particles
609	٠	$lpha_{\sf gas}$	[-]	Volumetric absorptivity of light by gases (H ₂ O, CO ₂)
610	٠	$\boldsymbol{\varepsilon}_{gas}$	[-]	Volumetric emissivity of light by gases (H ₂ O, CO ₂)
611	٠	$\delta_{ ext{shutter}}$	[-]	Fraction of aperture of the shutter obstructer
612	٠	Ψ	[-]	Structure parameter, used in RPM
613	٠	$\eta_{ ext{opt,concentrator}}$	[-]	Optical efficiency of the heliostat + concentrator facility
614	٠	$\eta_{ m opt,receiver}$	[-]	Optical efficiency of the window + zirconia felts restriction
615	٠	A _{proj}	[m²]	Parabolic concentrator area irradiated by sunlight (horizontal
616			nsidering the rea	actor shading effect)
617	٠	C _x	[kmol/m³]	Molar concentration of the gas component X
618	٠	F _x	[kg/s – NL/s]	Flow rate of species X
619	٠	m _c	[kg]	Mass of carbon
620	٠	P _x	[Pa]	Partial pressure of the gas component X
621	٠	Q _{loss,X}	[W]	Heat flux lost by the reactor
622	٠	Q _{sun,1}	[W]	Solar power delivered by the parabolic concentrator
623	٠	Q _{sun,2}	[W]	Solar power available at the cavity entrance
624	٠	Q _{X-Y}	[W]	Heat flux between two system components X and Y
625	•	T _{probe}	[K]	Temperature measurement by the thermocouple
626	٠	T _{pyro}	[K]	Temperature measurement by the pyrometer
627	٠	T_{gas}	[K]	Temperature of the gas + char phase in simulations
628	٠	T _{wall}	[K]	Temperature of the wall (inner layer) in simulations
629	٠	X _c	[-]	Advancement of a carbon mass oxidation
630	٠	Y_{char}	[-]	Char mass yield obtained after devolatilization (dry basis)

631 DECLARATION OF COMPETING INTEREST

The authors declare that they have no known competing financial interests or personal relationships

that could have appeared to influence the work reported in this paper.

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