Fischer-Tropsch biofuels production from syngas obtained by supercritical water reforming of the bio-oil aqueous phase

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Abstract

A new process to produce low-temperature Fischer-Tropsch (LTFT) products from the syngas obtained by supercritical water reforming (SCWR) technology of the bio-oil aqueous phase is developed and analyzed. The process includes four sections: syngas production from SCWR, syngas upgrading by water-gas-shift and dry reforming reactors as well as PSA systems, LTFT synthesis and products refining and upgrading through a distillation columns train and hydrocracking reactor. The aim is to produce maximum biofuels and electrical power, achieving the overall energy self-sufficiency. The energy scheme also involves cogeneration (e.g., hot water for district heating) and removal of CO₂ for sequestration. Process simulations were carried out by Aspen Plus. The effect of the main operating parameters (feed concentration and composition, as well as operating conditions of LTFT reactor) on the process performance (carbon efficiency with or without refining, biofuel and electricity production) was studied by a sensitivity analysis. This way, the optimal conditions were found, so for a feeding of 60 t/h with a concentration of 35 wt.%, carbon efficiency with refining is 38.5% (4.6 t/h biofuel) and 5.3 MWe is generated. The CO₂ for sequestration is 0.50 kg/kg of organic feeding.

Keywords

Fischer-Tropsch, Supercritical Water, Bio-oil, Syngas, Reforming, Biofuel

1. Introduction

A few main routes make it possible to produce gas and liquid biofuels, such as extraction of vegetable oils, fermentation of sugars to alcohol, or gasification and chemical synthesis. Some recent studies indicate that the use of Fischer-Tropsch (FT) technology for biomass conversion to synthetic hydrocarbons may offer a promising alternative to conventional diesel and gasoline [1-3]. This synthesis was developed in the 1920s as a chemical process to produce hydrocarbons of different length (i.e. light gases, gasoline, jet fuel, diesel and wax fractions) from a mixture of CO and H₂ called syngas.

Hydrocarbon formation in the FT synthesis is comparable to a polymerization mechanism in which the chain initiation involves the adsorption and dissociation of CO reactant on the catalyst surface. This is followed by hydrogenation of surface carbon atoms to form methylene groups that act as monomer units (-CH₂-) in the polymerization. Thus, free radical species methylene ($\dot{C}H_2$) react with hydrogen to form a methyl group, which is the chain initiator. Chain propagates by sequential integration of methylene to alkyl groups thus leading to long-chain hydrocarbons. The termination step occurs by reduction or β -hydride abstraction to produce n-paraffins or α -olefins, respectively [4]. The highly exothermic reactions describing the formation of paraffins and olefins from syngas in FT synthesis are represented by reactions (R1) and (R2) [5]:

$$n \operatorname{CO} + (2n+1) \operatorname{H}_2 \leftrightarrow \operatorname{C}_n \operatorname{H}_{2n+2} + n \operatorname{H}_2 \operatorname{O}$$
(R1)

$$n \operatorname{CO} + 2n \operatorname{H}_2 \leftrightarrow \operatorname{C}_n \operatorname{H}_{2n} + n \operatorname{H}_2 \operatorname{O}$$
(R2)

Hydrocarbon production in FT synthesis depends on temperature, thus distinguishing high-temperature FT (HTFT) synthesis (300-350 °C), and low-temperature FT (LTFT) synthesis (180-250 °C) [2]. HTFT process produces mainly light hydrocarbons (LPG and gasoline) and LTFT produces heavier hydrocarbons (mainly, diesel and wax). The cobalt-

based catalyst is used on an industrial scale in either fixed-bed reactors (Shell) or slurry reactors (Sasol), but only for low temperature Fischer-Tropsch (LTFT) process, while iron-based catalyst is used for both HTFT and LTFT synthesis processes [4].

Liquid synthetic fuels (with high volumetric energy density) can be transported by the same means as oil and are free of sulfur, nitrogen and aromatics, which are typically found in normal gasoline and diesel. Thus, emissions from internal combustion engines are reduced. Liquid synthetic fuels are receiving much attention as a real alternative to reduce the petroleum dependence of the transportation sector and can be easily obtained from natural gas, coal and biomass gasification. The resulting processes are known as gas to liquid (GTL), carbon solid to liquid (CTL) and biomass to liquid (BTL), respectively [3]. CTL may involve considerable emissions of CO₂, SO_x and NO_x and particulate matter, so gas cleaning systems are necessary. Likewise, availability, transportation, handling, storage and low energy density of biomass are obstacles in the development of large scale biomass processing. For these reasons, biomass can be converted into more transportable forms such as liquid bio-oils through fast pyrolysis, which involves thermal decomposition reactions that occur in a few seconds in the absence of oxygen. Liquid biooil has a higher energy density as compared with solid biomass, so those storage space and transportation problems associated with biomass are reduced [6].

By adding water, bio-oil can be separated in a valuable oil phase and in a wasteaqueous phase, which contains organics compounds (20-30 wt.%) and may be reutilized and valorized. In this case, supercritical water (SCW) reforming is a suitable process because water does not need to be vaporized. Besides, SCW has additional advantages such as a high capability to solubilize gaseous organic molecules. Indeed, SCW reforming is an emergent technology that has being investigated the last years, as shown in our previous works about the valorization of glycerol from biodiesel production [7-11].

High energy demand along with large capital costs have been the main drawbacks of FT plants, which may call into question the economic viability of the FT process. The first issue is dealt with in this manuscript, and the second one will be treat in a future to optimize the price of biofuels. Thus, in this paper, a new process of synthetic fuels production from the combination of SCWR of aqueous phase of bio-oil and Fischer-Tropsch synthesis is designed and analyzed, achieving a novel and efficient via of valorization for the waste-aqueous phase of bio-oil. Therefore, liquid carbon from waste to liquid (WTL) is the process considered in this study. Figure 1 depicts the concept in a simple way using a block flow diagram. Aqueous phase of bio-oil is reformed under supercritical conditions and converted into syngas. Syngas from SCWR process is expanded in a turbine to generate electrical power and upgraded through water-gas shift and dry reforming reactors as well as by two PSA systems. This way, the molar flow rate of H₂ and CO in syngas are increased and a desired H₂/CO ratio is achieved at the inlet of the FT reactor, which includes a loop to recirculate the unreacted syngas, thus increasing the overall conversion of CO into biofuels. A fraction of recycle gas is purged to avoid inert gases accumulation, and FT liquid is separated in different cuts by a distillation train. In addition, high conversion of the LTFT wax can be obtained under mild hydrocracking conditions thanks to the high chemical reactivity of heavy paraffin and the absence of catalysts contaminants like sulfur or nitrogen compounds [12]. Industrially, the operation of LTFT followed by hydrocracking is industrially applied, as in the Shell Middle Distillate Synthesis Process [13]. An external fuel is not needed as a heat source, because the process is designed to be energy self-sufficient by burning the off-gas from different sections. The proposed process is designed to maximize its performance in terms of (1) biofuels production and (2) net electrical power generation, as well as (3) cogeneration water and (4) pure CO₂ for sequestration.



Figure 1. Block flow diagram of the process.

2. Process design and simulation

The flowsheet of the SCWR-FT process is illustrated in **Figure 2**, and equipment specifications are shown in **Table 1**. The process is divided into four different sections: (1) supercritical water reforming (SCWR) of the bio-oil aqueous fraction, (2) upgrading of the syngas to increase the molar flow rate of H₂ and CO and to achieve the H₂/CO molar ratio at the FT reactor inlet by water-gas-shift (WGS) and dry reforming (DR) reactors, as well as by two PSA systems, (3) Fischer-Tropsch synthesis loop, and (4) refining and upgrading of FT products through distillation columns and hydrocracking.



Lines colors: Red for flue gas from furnace, black for organic mixture and gases from SCWR, pink for gases from DR, brown for FT process, blue for cooling water, green for hydrocracking products and grey for distillation zone. Likewise, in some heat exchangers in which one of fluids is cooling water or flue gas from the furnace. the corresponding stream is simplified by mentioning "F- ..." (*"from provenance*") at the inlet. and "T-..." (*"to destination*") at the outlet.

Figure 2. Heat-integrated flowsheet of the SCWR-LTFT process.

Code	Equipment	Specifications
P1	Pump	Efficiency: 0.8; Outlet pressure: 240 bar
SCWR	Supercritical Water Reforming	Operating temperature: 800 °C; Pressure drop: 0.0 bar
TURB	Turbine	Type: Isentropic; Efficiency isentropic: 0.85; Outlet pressure: variable
HWGS	Water gas-shift reactor, type REquil (High-temperature stage)	Operating temperature: $350 ^{\circ}$ C; Pressure drop: 0.1 bar
SEP1_4	Gas-liquid separators (Flash)	Temperature: $35 ^{\circ}\text{C}$ Pressure drop: 0.1 har
PSA1	Pressure swing adsorption unit	It removes most the H ₂ (95 %) from the other gases [15]
	(First PSA system)	Outlet pressure: variable (H ₂ stream, 16-H), 1.1 bar (the rest of gases)
PSA2	Pressure swing adsorption unit (First PSA system)	CO-rich stream: 98 % CO, 1 % for CO ₂ and CH ₄ [15] Outlet pressure: 1.1 bar (top and bottom)
COMP1 and 3	Trains of compression with two	Type: Isentropic; Isentropic efficiency: 0.76; Mechanical efficiency:
	intermediate coolers and one final	0.98; intermediate cooling to 35 °C; Outlet pressure: 15 bar
DRVREE	Dry reforming reactor, type REquil	Isothermal reactor: 600 °C: Operating pressure: 1.1 bar
DRTKEF	Dry reforming reactor, type KEquit	$CH_4 + CO_2 \leftrightarrow 2H_2 + 2CO_2$
PSA1A	Pressure swing adsorption unit	It removes most the H_2 (95%) from the other gases [15]
1 5/11/1	(Second PSA system)	Outlet pressure: variable (H ₂ stream, 31-H), 1.1 bar (the rest of gases)
PSA2A	Pressure swing adsorption unit	CO-rich stream (bottom): 98 % CO, 1 % for CO ₂ and CH ₄ [15]
	(Second PSA system)	Outlet pressure: 15 bar (34 stream), 1.1 bar (35-CO stream)
PSA3A	Pressure swing adsorption unit	CO ₂ -rich stream (bottom): 90 % CO ₂ , 9 % CH ₄ , 0.5 % CO and H ₂
GOL (D2 14	(Second PSA system)	15. Outlet pressure: 1.1 bar (top and bottom)
COMP2 and 4	I rains of compression with two	Type: Isentropic; Isentropic efficiency: 0.76; Mechanical efficiency:
	intermediate coolers and one final	0.98; interinediate cooling to 55°C; Outlet pressure: variable (but 5.45
LTFT	Low-Temperature Fischer-Tropsch	Operating temperature: variable (220-240 °C)
	reactor (RStoic)	Operating pressure: variable (220-240 C)
DECANT	Liquid-liquid separator (Decanter)	Temperature: 35 °C.; Pressure drop: 0.1 bar
COMP5	Compressor	Type: Isentropic; Isentropic Efficiency: 0.7; Mechanical efficiency:
	-	0.98; Outlet pressure: 20-40 bar
COMP6	Compressor	Type: Isentropic; Isentropic Efficiency: 0.7; Mechanical efficiency: 0.98; Outlet pressure: 20-35 bar
PSA4	Pressure swing adsorption unit	It removes most the H_2 (95 %) from the other gases [15]
		Outlet pressure: same SEP4
DEST1	Distillation column 1 (heavy-light)	Operating pressure: 1.5 bar (reboiler) and 1.3 bar (condenser)
		Reflux ratio: 1.2R _{min}
		$C_{13}H_{28}$ recovery (distillate stream): 90 %
DEST2	Distillation column 2 (way diasel)	Clamping pressure: 1.5 her (rebailer) and 1.3 her (condenser)
DESIZ	Distillation column 2 (wax-dieser)	Reflux ratio: 1.2R _{min}
		C ₂₀ H ₄₂ recovery (distillate stream): 90 %
		$C_{21}H_{44}$ recovery (bottom stream): 90 %
DEST3	Distillation column 3 (jet fuel-C ₁₀₋)	Operating pressure: 1.3 bar (reboiler) and 1.2 bar (condenser)
		Reflux ratio: 1.2Rmin
		C ₉ H ₂₀ recovery (distillate stream): 95 %
		C ₁₀ H ₂₀ recovery (bottom stream): 95%
SEP5	Gas-liquid separator	Temperature: same that distillate stream in DEST3 Pressure drop: 0.1 bar
COMP7	Compressor	Type: Isentropic; Isentropic Efficiency: 0.75; Mechanical efficiency: 0.98; Outlet pressure: 35 har
P2	Pump	Efficiency: 0.7: Outlet pressure: 35 bar
HYDRCR	Hydrocracking reactor (RStoic)	Operating temperature: 360 °C: Pressure: 35 bar
	,	OUTHYD stream: 26 wt.%, C ₁₀ -C ₁₃ ; 42 wt.%, C ₁₄ -C ₂₀ ; 32 wt.% C ₂₀₊ [27]
FAN1	Fan	Type: Isentropic; Isentropic Efficiency: 0.76; Mechanical efficiency:
FURNACE	Furnace-combustor (RStoic)	Combustion of everything able to be oxidized: $1000 \ \odot \ \odot \ 1 \ ber$
I OIUMOL		Surplus heat flow to SCWR
V1-V6, V9-V11	Valves	Outlet pressure: 1.1 bar
V7	Valve	Outlet pressure: 1.5 bar
V8	Valves	Outlet pressure: variable

Table 1 (a). Specifications of the individual process units,

C 1	T • (
Code	Equipment	Specifications
HE01	Heat exchanger	Cold stream outlet temperature: variable
HE02	Heat exchanger	Hot stream outlet temperature: 700 °C
HE03	Heat exchanger	Hot stream outlet temperature: 350 °C
HE04	Heat exchanger	Cold stream outlet temperature: variable
HE05	Heat exchanger	Hot stream outlet temperature: 35 °C
HE06	Heat exchanger	Cold stream outlet temperature: variable
HE07	Heat exchanger	Cold stream outlet temperature: 600 °C
HE08	Heat exchanger	Cold stream outlet temperature: 360 °C
HE09	Heat exchanger	Cold stream outlet temperature: variable
HE10	Heat exchanger	Hot stream outlet temperature: 35 °C
HE11	Heat exchanger	Cold stream outlet temperature: variable (210-215 °C)
HE12	Heat exchanger	Cold stream outlet temperature: variable (220-240 °C)
HE13	Heat exchanger	Hot stream outlet temperature: 35 °C
HE14	Heat exchanger	Hot stream outlet temperature: 35 °C
HE15	Heat exchanger	Hot stream outlet temperature: 360 °C
HE16	Heat exchanger	Cold stream outlet temperature: 98 °C
HE17	Heat exchanger	Cold stream outlet temperature: variable
HE18	Heat exchanger	Hot stream outlet temperature: 99 °C
C1	Heat exchanger	Heat flow for endothermic heat reaction in DRYREF
C2	Heat exchanger	Sink of the heat flow coming from the condenser of the DEST3
C3	Heat exchanger	Sink of the heat flow coming from the condenser of the DEST1
C4	Heat exchanger	Sink of the heat flow coming from the condenser of the DEST2
C5	Heat exchanger	Heat flow from LTFT
H1	Heat exchanger	Heat flow towards the reboiler of DEST1
H2	Heat exchanger	Heat flow towards the reboiler of DEST2
H3	Heat exchanger	Heat flow towards the reboiler of DEST3
SP1	Splitter	Split fraction stream 10: variable (the objective is to achieve the
	•	hydrogen molar flow required to adjust the H2/CO molar ratio).
SP2	Splitter	Split fraction stream 15: variable (aimed at achieving the energy self-
		sufficiency constraint).
SP3	Splitter	Split fraction stream 17-H-C: variable on required H ₂ for
		hydrocracking.
SP4	Splitter	Split fraction stream 46: variable (aimed at maximizing CO total
an .	a. !!	conversion as much as possible).
SP5	Splitter	Split fraction stream 51-G: variable (with the goal of sending the
CD(0.1"#	required hydrogen to the hydrocracking reactor)
SPo	Splitter	Split fraction stream CH4-O: variable (in some cases, biomethane could be sold if an array solf sufficiency was achieved)
SD7	Splitter	Split fraction stream 06V: variable (with the goal of preheating the
SF /	Splitter	I TET inlet stream and producing electrical power by the steam turbine
		with the rest of steam (02V stream)
P3-P4	Pumps (streams W4, W5) -not	Efficiency: 0.75: Outlet pressure: 20 bar
	drawn in Figure 2-	
P5-P7	Pumps (streams W1, W2, W3) -not	Efficiency: 0.7: Outlet pressure: 1.1 bar
	drawn in Figure 2-	,,

 Table 1 (b). Specifications of the individual process units.

2.1. Supercritical water reforming of the aqueous fraction of bio-oil

A mixture of organic compounds representative of the bio-oil aqueous phase is pumped at 240 bar and heated as much as possible by four heat exchangers. The supercritical water reforming (SCWR) reactor operates at 240 bar and 800 °C to maximize H_2 and CO yields. SCWR reactor is simulated as a Gibbs reactor (RGibbs), where the products yields are calculated under conditions that minimize the Gibbs free energy. This reactor would be located inside of a furnace where a fraction of the product gas from SCWR and WGS reactors, light hydrocarbons gases from distillation section (separator SEP5), unconverted gases from FT synthesis and hydrocracking (PSA4) and the biomethane surplus from dry reforming reactor (PSA3) are burnt to achieve the energy self-sufficiency. The furnace, simulated by a stoichiometric reactor (RStoi), transfers the heat needed by the SCWR reactor, and the flue gas stream leaves the furnace at 1000 °C to provide heat in other parts of the process. The product gas leaving the supercritical water reformer is expanded by a turbine up to 40 bar to generate electrical power.

2.2. Upgrading of the syngas and fit of H₂/CO molar ratio for FT synthesis

A fraction of syngas from the turbine outlet enters the high-temperature water-gas shift (HWGS) reactor and the rest by-passes this unit. That fraction is calculated to increase the H₂ molar flow rate when the H₂/CO molar ratio at inlet Fischer-Tropsch reactor is insufficient, which occurs for high organic concentrations of the bio-oil aqueous phase (25-35 wt.%). The high-temperature WGS reaction is moderately exothermic and industrially performed over Fe-Cr catalyst for temperature ranging from 350 to 450 °C [14]. Thus, HWGS reactor operates at 350 °C and is simulated as an equilibrium reactor (REquil). The gas leaving the HWGS reactor is mixed with the by-pass and cooled to 35 °C in order to condense the steam, which is separated from the gas stream.

The fraction of syngas that is not sent to the furnace is processed by two pressure swing adsorption (PSA) units to obtain a pure H₂ stream (at the top of the first PSA unit), as well as a pure CO stream (at the bottom of the second PSA unit) and a CO₂+CH₄-rich stream (at the top of the second PSA unit). Details of these units are described in elsewhere [15]. PSA units are simulated as ideal separators, at 35 °C and the pressure of syngas. The depressurization steps are simulated by valves, and PSA units are assumed

to give a high purity H_2 (99 %) with a hydrogen recovery of 90 % [15], as well as CO purity higher than 90 % with a CO recovery of 98 %. A fraction of the H₂-rich stream is sent for hydrocracking and the rest is conducted to the FT synthesis loop, like the CO-rich stream after recompressing it to the operating pressure of the FT reactor.

The CH₄ and CO₂ rich stream, with minor H₂ and CO concentrations, is depressurized to 1 bar and heated to 600 °C before entering the dry reforming (DR) reactor, simulated as REquil. This is a reactor with a nickel-based catalyst where the molar flow rates of CO and H₂ increase in a H₂/CO ratio close to unity, which further justifies the previous WGS stage to achieve the H₂/CO ratio about 2.0 required for FT synthesis. DR reactor allows the increase in the flow rate of syngas fed into the FT synthesis loop and reduces the CO₂ emissions to the atmosphere. However, dry reforming is highly endothermic and temperatures of 600 °C or higher [16] are necessary to increase the conversion of the reactants and to minimize carbon deposition on the catalyst. Nevertheless, beyond 800 °C, the energy required exceed much the gains in CO₂ and CH₄ conversions, so the temperature of DR reactor was set at 600 °C to reduce the energy requirement. Likewise, a pressure of 1 bar was used to favor the CO and H₂ production, as pointed by others [17, 18] and verified by simulation by us. Under these operating conditions, CH₄ conversion

The humid gas stream leaving the DR reactor is cooled to 35 °C, so the condensate (mainly water) is removed through a separator. Afterwards, the gas is compressed to the operating pressure of Fischer-Tropsch reactor, and enters the second PSA system. As in the first system, PSA1A and PSA2A allow the separation of H₂-rich and CO-rich streams that are conducted to the inlet of the FT synthesis loop. On the other hand, the CH₄ and CO₂ rich stream enters PSA3 to obtain a stream of CO₂ with a purity higher than 95 % to

match the specification for transport and sequestration of CO_2 [19], and a CH₄-rich stream, which is sent to the furnace or even sold as biomethane if there is energy surplus.

2.3. Fischer-Tropsch synthesis loop

 H_2 and CO streams, at the operating pressure of the FT reactor, are mixed and heated to the temperature of the FT reactor. In the FT process, the probability of chain growth on the surface of the catalyst (α) is defined by the rate of chain propagation divided by the sum of propagation rates and chain termination. This parameter depends on the reaction conditions and catalyst type, and it is calculated by Eq. (1), obtained by Song et al [20] for cobalt-based catalysts used for LTFT.

$$\alpha = \left(A \frac{y_{CO}}{y_{H2} + y_{CO}} + B\right) (1 - 0.0039(T - 523)) \tag{1}$$

where A and B are coefficients with values of 0.2332 ± 0.0740 and 0.6330 ± 0.0420 , respectively. Likewise, y_{CO} and y_{H_2} are the mole fractions of CO and H₂ in the feed stream to the FT reactor, and *T* (K) is the operating temperature of the FT reactor. FT products were assumed to be composed only of paraffins and olefins, which are the main products of the LTFT synthesis [12]. Eq. (2) shows that the production of olefins (*O*) with respect to paraffins (*P*) varies exponentially with carbon number (*n*) [21].

$$O/P = exp\left(-\frac{\Delta E}{RT}n\right) = e^{-cn}$$
⁽²⁾

O/P is the olefin-to-paraffin molar ratio, and ΔE is the change in the activation energy for the olefin desorption step per every CH₂ group, caused by weak van der Waals forces. The value for ΔE is 1.1 kJ/mol C atom, which leads to a value of *c* between 0.26 and 0.28, for the temperature range of the FT reactor used in this research [21, 22]. On the other hand, product selectivity is usually described by a statistical distribution model known as the Anderson-Schulz-Flory (ASF) distribution law, which provides the molar fraction (α_{Cn}) of hydrocarbons produced by FT synthesis (Eq. (3)) and depends on the number of carbon atoms of the produced hydrocarbons (*n*) [5]. Likewise, Eq. (4) gives the weight fraction (w_{Cn}) of formed hydrocarbons.

$$\alpha_{Cn} = \alpha^{n-1} \left(1 - \alpha \right) \tag{3}$$

$$w_{cn} = \alpha^{n-1} (1-\alpha)^2 n$$
 (4)

An α -value of 0.90 was obtained using the highest values for A and B in Eq. (1). This high value increases the product selectivity towards high molecular mass linear wax and diesel [2, 4]. **Figure 3** shows the evolution of weight distribution versus carbon number, matching typical values in LTFT process [4, 23]. Larger quantities of liquid biofuels are achieved for higher α -values. In this sense, cobalt-based catalysts allow a relatively high chain growth probability and a very low activity towards the undesired water-gas shift reaction [24].



Figure 3. LTFT product weight distribution versus carbon number for different α -values.

The FT product distribution is directly related to the CO fractional conversions relative to each hydrocarbon produced in the FT reactor, which depends on the α -value and the paraffins-to-olefins ratio. Since α -value depends on temperature and H₂/CO molar

ratio, these variables also influence on CO fractional conversions and the CO conversion per pass, but this latter does not affect the CO fractional conversions. Additionally, pressure must affect both α -value and CO fractional conversions, but Eq. (1) does not consider the effect of pressure. Indeed, there is no an accepted equation that provides a relationship between pressure and α -value. Thus, reported values for different pressures were used to inspect the effect of the pressure on the performance of the FT reactor [23].

In the base case, the LTFT reactor operates at 20 bar and 220 °C, as well as with an inlet molar ratio H₂/CO of 2.0, to promote FT reactions with Co-based catalysts that allow a long-chain hydrocarbon generation [4]. Under these conditions, CO conversion per pass was assumed to be 50 %, as previously reported [4, 25], and α -value was 0.90 (computed by Eq. (1)). The stream leaving the FT reactor is cooled, so heavier hydrocarbons are condensed and separated from the gas, which is sent back to the FT reactor inlet through a recycle loop to maximize the overall CO conversion (about 87 %, as reported [26]) and to increase the specific production of liquid fuel. To avoid the build-up of inert gases, this configuration contains also a purge. On the other hand, the liquid phase leaving the loop is conducted to a decanter for separating water from heavier hydrocarbons, which are expanded and sent to the distillation section.

2.4. Refining and upgrading of Fischer-Tropsch products

Purge from LTFT process and outlet stream from hydrocracking reactor are mixed and cooled to 35 °C, thus entering a flash separator to remove the gases (unconverted CO and H₂, as well as CO₂, CH₄ and light hydrocarbons (C₂-C₅)) from the liquid, which contains traces of gaseous compounds. The liquid stream enters a distillation column (DEST1), so the light stream (at the top) is composed of jet fuel, gasoline, light hydrocarbons (C₂-C₄ with somewhat of C₅) and traces of water, H₂, CO and CO₂, and the heavy products (at the bottom) are composed of diesel and wax. These latter are separated in the second column (DEST2), where wax is conducted to the hydrocracking reactor. Likewise, the stream leaving the top of DEST1 is fed into another column (DEST3) for separating jet fuel (at the bottom) and light gas and gasoline (at the top), which are further separated in a flash separator.

Hydrocracking process of long chain wax (heavier paraffins with a boiling point temperature higher than 370 °C [27]) is an effective route to maximize the production of high-quality middle distillates (diesel and jet fuel) in LTFT synthesis. The unconverted H₂ from LTFT synthesis (and, if necessary, a fraction of the H₂-rich stream leaving PSA1) is used as reactant for hydrogenation in the hydrocracking process. In this study, only alkanes with carbon number equal or higher than 20 are considered to react and hydrocracking products are alkanes with carbon number between 10 and 15. Hydrocracking temperature must be higher than 280 °C [28] to maximize the selectivity of the C₁₀-C₂₀ fraction, for a bifunctional platinum-based catalyst [29]. The presence of H₂O and CO might poison and deactivate the noble-metal hydrocracking catalyst [30]. Likewise, olefin production rate may increase with the CO concentration fed into the hydrocracking reactor, thus reducing the liquid fuel selectivity.

The carbon number distribution of cracked products is assumed as fully symmetrical and centered at around half of the original molecule, so equal molar amounts are formed [31]. This matches the result of that the most frequent product is shifted towards a number of carbon atoms that is slightly under the medium value [32]. Therefore, the reactions are:

For even n:
$$C_nH_{2n+2} + H_2 \rightarrow 2C_{n/2}H_{2(n/2)+2}$$
 (R3)
For odd n: $C_nH_{2n+2} + H_2 \rightarrow C_{(n+1)/2}H_{2((n+1)/2)+2}C_{(n-1)/2}H_{2((n-1)/2)+2}$ (R4)

A RStoic reactor is used to simulate the hydrocracker, where CO fractional conversions were specified according to results found in the literature [27], so the outlet

stream has 32.90 wt.% of reactants (mass fraction of C_{20+}) and 67.10 wt.% (25.30 % for jet fuel and 41.80 % for diesel) of products (mass fraction of $C_{10}-C_{20}$). The hydrocracking operating conditions were 360 °C and 35 bar, and 0.06 kg H₂/kg wax, within the ranges previously reported [12, 27], to achieve a high extent of cracking reaction and to maximize the distillate fraction. Highly exothermic hydrogenation reaction and endothermic cracking reaction take place in the hydrocracking reactor, so the amount of heat released in the hydrogenation reactions might be higher or lower than the heat required for the endothermic cracking reactions. Although others have found an overall net heat slightly negative (exothermic) [33], all cases of this study presented a net heat slightly positive (endothermic), which increases as the amount of hydrogen fed into the reactor increases, because hydrogen absorbs a part of the released heat. Thus, this reactor is almost isothermal. Outlet stream from hydrocracking reactor is expanded to the pressure of purge of FT synthesis loop.

2.5. Thermal energy integration

Figure 2 includes the energy integration through heat exchangers located in the process to improve the energy use. The strategy followed for heat exchangers network is that streams close to each other at high thermal levels are the first to contact each other (the high-temperature hot streams warm up the high-temperature cold streams), and streams at low thermal levels are the last to contact each other (the low-temperature hot streams heat the low-temperature cold streams). SCWR and DR are the central points, where maximum temperatures are set. Although there could be more alternative flow sheets for this process due to possible redistributions of heat exchangers, the proposed flow-sheet provides the best heat integration performance, because minimizes the use of external utilities and maximizes heat flows into the system.

The heat required to maintain endothermic reactions in DR reactor is transferred from the hot flue gas coming from the furnace. Similarly, a heat stream exits from furnace towards SCWR to heat the feed of this reactor to 800 °C and to maintain the endothermic chemical reactions.

Likewise, the heat of highly exothermic FT reactions (about 150 kJ per mole of CO converted [2]), must be efficiently released to avoid catalyst deactivation via sintering and formation of methane, which is undesirable since selectivity of preferred products would decrease. In this study, LTFT is equipped with an evaporator where cooling water, coming from the heat exchanger located at the outlet of the FT reactor to cool down the FT products, enters as liquid at 212 °C and 20 bar and exits but as saturated steam at the same conditions. A fraction of this steam is conducted to preheat the inlet FT stream and the rest is expanded (to 1 bar and 99 °C) in a steam turbine (TURB2) to produce electricity.

In a similar way, the feed to DR reactor is heated to 600 °C by two heat exchangers. Similarly, the process stream must be heated to the specified operating temperature of LTFT by two heat exchangers. Afterwards, streams of flue gas and from the DR products transfer heat to the kettles of distillation columns DEST1 and DEST2. Lastly, all cooling water streams (at atmosphere pressure and 99 °C) are collected, so the energy scheme also involves cogeneration (e.g., hot water for district heating) thus increasing the overall efficiency of the process. Finally, the flue gas from furnace is cooled to about 120 °C before emitting it to the atmosphere.

2.6. Composition and mass flow rate of the bio-oil aqueous phase

Table 2 shows the feedstock composition of this process, representative of fastpyrolysis liquids from pine wood [34] for different total concentration (15-35 wt.%).Besides, three values of inlet mass-flow (20, 60 and 100 t/h), within the range of industrial

plants [35-37], and some changes in the concentrations of acetic acid, acetol and levoglucosan [38-40] were used in different simulations.

On the other hand, acetic acid, acetol and butanol are major constituents of the carboxylic acids, ketonic and alcohol fraction of bio-oil, and the supercritical water reforming of these model compounds, including glucose instead of levoglucosan [41], have been experimentally studied in our previous work [42]. Thus, **Table 2** also includes the feedstock used in [42], which was simulated for a total concentration of 25 wt.% to compare with the results of the more complete composition.

Compound	Formula	Case 1 (wt.%)	Case 2 (wt.%)	Case 3 (wt.%)	Case 4 (wt.%)	Case 5 (wt.%)	Case 6 (wt.%)	Case 7 (wt.%)	Case 8 (wt.%)	Case 9 (wt.%)
Acetic acid	C ₂ H ₄ O ₂	8.00	6.86	5.72	3.58	3.58	4.57	3.43	10.00	10.00
Hydroxyacetone	C ₃ H ₆ O ₂	5.00	4.29	3.58	5.72	3.58	2.86	2.14	5.00	5.00
1-Butanol	$C_4H_{10}O$	3.00	2.57	2.14	2.14	2.14	1.71	1.29	5.00	5.00
Acetaldehyde	CH ₃ CHO	1.00	0.86	0.71	0.71	0.71	0.57	0.43	-	-
Formic acid	CH ₂ O ₂	3.00	2.57	2.14	2.14	2.14	1.71	1.29	-	-
Butyric acid	$C_4H_8O_2$	2.00	1.71	1.44	1.44	1.44	1.14	0.86	-	-
Methanol	CH ₃ OH	1.50	1.28	1.07	1.07	1.07	0.86	0.64	-	-
Guaiacol	C7H8O2	1.00	0.86	0.71	0.71	0.71	0.57	0.43	-	-
Glyoxal	C ₂ H ₂ O ₂	1.00	0.86	0.71	0.71	0.71	0.57	0.43	-	-
Furfural	C5H4O2	1.00	0.86	0.71	0.71	0.71	0.57	0.43	-	-
1,4-Hydroxybenzene	C ₆ H ₆ O ₂	1.00	0.86	0.71	0.71	0.71	0.57	0.43	-	-
Propionic acid	C ₃ H ₆ O ₂	1.00	0.86	0.71	0.71	0.71	0.57	0.43	-	-
Phenol	C ₆ H ₆ O	1.50	1.28	1.07	1.07	1.07	0.86	0.64	-	-
Levoglucosan	C6H10O5	5.00	4.28	3.58	3.58	5.72	2.86	2.14	5.00	-
Glucose	C6H12O6	-	-	-	-	-	-	-	-	5.00
TOTAL	35.00	30.00	25.00	25.00	25.00	20.00	15.00	25.00	25.00	

Table 2. Case studies for different composition and total concentration of the bio-oil aqueous fractions

2.7. Simulation remarks and components from databank

Simulation was carried out using Aspen Plus version 8.0 (Aspen Technology, Inc., USA) to compute the mass and energy balances under different operating conditions. The thermodynamic methods were the predictive Soave-Redlich-Kwong (PSRK) Equation of state (EoS) for SCWR section [7], UNIQUAC for distillation train, STEAM-NBS for steam turbine (TURB2), and Peng-Robinson EoS, which is recommended when dealing

with hydrocarbons, for the rest of the plant (PSA systems, as well as DR, FT and hydrocracking reactors).

The organic compounds conversions were always 100 % (equilibrium conditions) in SCWR reactor (RGibbs) and HWGS reactor (REquil), but not in DR reactor that also operates at equilibrium conditions (REquil). The process simulation includes 66 chemical compounds, like O_2 , N_2 , H_2 , CO_2 , CH_4 , CO, glucose, water and all organic compounds shown in **Table 2**, apart from all the linear hydrocarbons from C_2 to C_{30} for paraffins and C_2 to C_{16} for olefins that represent the products of FT synthesis. As FT synthesis produces mainly linear hydrocarbons, and saturated hydrocarbons are barely representative, only linear hydrocarbons were selected to describe light (fuel) gas, gasoline, jet fuel, diesel and wax [4, 12], defined as those biofuel cuts including hydrocarbons $C_2 - C_4$, $C_5 - C_9$, $C_{10} - C_{13}$, $C_{14} - C_{20}$, and C_{20+} , respectively, similar to those previously reported [43,44]. Recovery fractions of hydrocarbons for each distillation column are presented in **Table 1(a)**, and they were chosen to achieve technical specifications, such as density, viscosity or cetane number, in liquid biofuels, thus minimizing the number of stages in each distillation column.

3. On the process optimization

The aim was to maximize the process performance in terms of carbon efficiency with or without refining (i.e., at the outlet of the FT synthesis loop or at the outlet of the process, just leaving the distillation section), as well as biofuel and electricity production, for a given mass flow-rate and total organic concentration of the bio-oil aqueous phase fed into the process. As a constraint, the process must be energy self-sufficient, so offgases with chemical energy must be sent to the furnace. The total net power is the power of the two turbines minus the consumed power in all the pumps, fans and compressors. Some potential optimization variables were fixed, such as the reforming pressure at 240 bar since it barely affects the process in the range from 200 to 300 bar [7]. The outlet pressure of the expander downstream from the SCWR reactor was 40 bar, as it is the minimum required to optimize the subsequent heat exchange that allows the maximization of biofuel and electricity production, as well as the separation to be performed in the PSA system. The optimal reforming temperature is between 750 and 800 °C to increase the CO and H₂ yields, and since higher temperatures involve slight gains of those yields but much more energy requirement, 800 °C was chosen. Similarly, the pressure and temperature of the HWGS (operating at the outlet pressure of the first expander), DR and hydrocracking reactors were the usual in these reactors for a good performance.

The purge (or recirculation) of the FT synthesis loop may be also optimized, and a minimum purge is necessary to avoid accumulation of inert gases in the FT synthesis loop. However, if the purge ratio increases, the overall CO conversion and biofuels production will decrease although, at the same time, a higher flow-rate of residual gases may be sent to furnace. Nevertheless, instead of burning a processed product (leaving the FT synthesis loop), a better option consists of burning a less processed product gas obtained upstream. In this way, a variable fraction of product gas is sent to furnace, and the recirculation-to-purge ratio is maximized. Simulations of both options (combustion of a fraction of product gas or a higher purge of FT synthesis loop) resulted in similar performance parameters. Therefore, an overall CO conversion regarding maximum recycle-to-purge ratio was 87%, which also maximizes the biofuel production in the FT synthesis loop. In this sense and to clarify, in this study, no reactor and separation systems is sized, so the effects of different operating variables must be understood without the constraint relative to the real unit size. Thus, e.g., although molar-flows of gases at inlet

LTFT stream vary in the case-studies performed in the sensitivity analysis, CO conversion per pass is assumed to be constant if the H₂/CO molar ratio, pressure and temperature are kept constant. However, if FT reactor was sized, the CO conversion per pass would be influenced by molar flow-rates of gases entering the reactor that would affect the unit performance. This will be considered in a future techno-economic assessment.

Thus, in summary, there are still some independent variables for optimization: the total organic concentration in the aqueous phase of bio-oil in the feed, and the operating conditions of the FT reactor. SCW reforming is clearly interesting when the feed has much water, so 15 to 35 wt.% are realistic feed concentration values for the aqueous phase of the bio-oil. More diluted concentrations imply higher energy demand, so energy self-sufficiency constraint may be violated. Likewise, high feed concentrations lead to a heat surplus in the overall process, and other reforming processes may be more suitable. Finally, the operating conditions of the FT reactor are referred to pressure, temperature and H₂/CO molar ratio, and the normal ranges for Co-based LTFT reactors (20-40 bar, 220-240 °C and 1.7-2.3 for H₂/CO molar ratio) are inspected.

4. Results and discussion

The base-case corresponds to an aqueous fraction of bio-oil with mass flow of 60 t/h and 25 wt.%, where the FT reactor operates at 220 °C, 20 bar, and H₂/CO molar ratio of 2.0. CO conversion per pass and α -value were 50 % and 0.90, respectively. **Table 3** shows the overall energy balance and the heat flows for all the heat exchangers. **Table 4** provides the comparison of biofuels properties with those obtained in literature and standards, so a good match is achieved. **Table 5** presents summarized simulation results for the main process streams, such as components and overall mass flow-rates, temperatures and pressures, which allows a comprehensive analysis of the process.

Table 6 illustrates the results of varying different operating parameters regarding the bio-oil aqueous phase (sensitivity analysis) and how they affect the production of LTFT products, electricity, as well as cogeneration water, which is discussed below. The main loss of carbon fed into the process is through the produced CO_2 , which is efficiently separated with high purity for sequestration, and light hydrocarbons (CH₄ and C₂-C₄), which are burnt for achieving the overall energy self-sufficiency of the process.

 Table 3.
 Overall energy balance for reforming of bio-oil aqueous phase using SCW and LTFT synthesis,

 and heat flows for all the heat exchangers (base case/case 3).

Work entering the system	(kW)	Enthalpy of the inlet st	reams (kW)	Unit	Q (kW)	Unit	Q (kW)
P1	687.52	Aqueous phase of bio-oil	-223497.54	HE01	18058.2	C1	5178.3
P2	5.56	Air	-3.60	HE02	4640.4	C2	112.3
P3-P4	22.91	W1	-859707.45	HE03	5598.8	C3	322.1
P5-P7	1.49	W2	-17635.02	HE04	1317.0	C4	138.2
FAN1	185.70	W3	-397229.11	HE05	19163.4	C5	9804.3
COMP1-COMP4 with inter. cool.	5010.27	W4	-11991.79	HE06	762.7	H1	1013.7
COMP5-COMP7	29.40	W5	-95229.16	HE07	2093.1	H2	223.9
				HE08	100.1	H3	253.0
				HE09	552.9		
TOTAL	5942.85			HE10	109.3		
Work leaving the system (kW)	Enthalpy of the outlet s	treams (kW)	HE11	2435.9		
Turbine TURBINE	-9995.65	Separated water (14-O)	-176284.47	HE12	111.8		
Turbine TURB2	-1674.77	Separated water (29-O)	-112.40	HE13	4731.7		
		Separated water (45-O)	-17344.22	HE14	626.0		
		Cogeneration water (WC)	-1353956.55	HE15	103.0		
		CO ₂ sequestered (CO ₂ -O)	-14331.79	HE16	1061.8		
		Tail gas	-54514.52	HE17	8396.7		
TOTAL	-11670.42			HE18	7642.4		

Elemental analysis and	In	this study		Typical US Refiners [51]			UNE-EN- UNE-EN-I ASTM ASTM	ISO 3675, ISO 12185, D 4052, D 1298	ASTM D 1655	Shell Bintulu LTFT refinery [43]		
properties	GasolineDieselJet FuelGasolineDieselJet FuelGasolineDiesel		Jet Fuel	Gasoline	Diesel	Jet Fuel						
C (wt.%)	85.20	85.93	86.65	85-88	84-87	87.0	-	-	-	-	-	-
H (wt.%)	14.80	14.07	13.35	15-17	16-33	12.3	-	-	-	-	-	-
O (wt.%)	0.00	0.00	0.00	0.00	0.00	0.5	-	-	-	-	-	-
H/C molar	2.25	2.13	2.18	-	-	1.70	-	-	-	-	-	-
MW (g/mol)	99.84	211.74	153.77	100	200	-	-	-	-	-	-	-
Density (g/cm ³) at 15 °C	0.70	0.81	0.76	0.75	0.85	-	0.72-0.78	0.82-0.89	0.78- 0.84	0.69	0.78	0.74
Viscosity (cP) at 15 °C	0.42	3.46	1.24	0.37-0.44	2.6-4.1	-	-	-	-	-	-	-
Aromatics (wt. %)	0	0	0	31	-	< 22	35 % (Max)	-	25 % (Max)	0	< 0.05	< 0.1
Cetane index	-	97	80	-	40-55	-	-	51 (Min)	-	-	76	58

Table 4. Comparison of biofuels properties with standards and other sources

Stream		06	07	10	14	15	16-H	17-Н-С	20-CO	22	23	30	31-Н	35-CO	T-MIX9
From		HE04	SCWR	SP1	SEP	SP2	PSA1	SP3	V4	HE07	DRYREF	COMP2	PSA1A	V6	SP6
То		SCWR	TURB	MIX1	SP2	PSA1	SP3	MIX6	MIX2	DRYREF	HE04	PSA1A	MIX3	MIX2	MIX9
H ₂ O	kg/h	45000	39926.9	39926.9	41.7	33.5	0	0	0	33.5	33.5	7.8	0	0	0
Total organic	lra/h	15000	0	0	0	0	0	0	0	0	0	0	0	0	0
compounds	Kg/II														
H ₂	kg/h	0	968.6	968.6	967.5	776.9	738.1	8.1	0	38.8	328.4	328.4	312.0	0	16.3
CO ₂	kg/h	0	12344.6	12344.6	11980.9	9620.7	0	0	96.8	9524.5	6364.1	6364.1	0	63.8	567.0
CH ₄	kg/h	0	2895.0	2895.0	2890.3	2320.9	0	0	23.2	2297.7	1145.6	1145.6	0	11.5	1089.9
СО	kg/h	0	3864.1	3864.1	3859.7	3099.3	0	0	3037.3	62.0	4084.9	4084.9	0	4003.2	81.3
Mass Flow	kg/h	60000	60000	60000	19740.5	15851.3	738.1	8.1	3157.3	11956.5	11956.5	11930.8	312.0	4078.5	1754.5
Temperature	°C	374	800	350	35.0	35.0	35.0	35.0	32.0	600	600	35.0	35.0	32.0	32.0
Pressure	bar	240	240	40.0	39.9	39.9	39.9	39.9	1.0	1.0	1.0	20.1	20.1	1.0	1.0
Stream		40	41	44	45	47-R	WAX	46-HYD	50-L	51-G	52-G	56-G	T-HE02	52	54
From		HE12	RFT	SEP3	DECANT	COMP5	P2	V8	SEP4	SP5	PSA4	MIX10	C1	V7	DEST1
То		RFT	HE13	DECANT	MIX7	MIX4	HE15	MIX5	MIX7	MIX8	MIX8	HE17	HE02	DEST1	DEST3
H ₂ O	kg/h	11.1	3925.4	3912.7	4.1	11.1	0	0	1.9	0.1	0	8.8	7376.9	6.0	6.0
H ₂	kg/h	1758.3	843.1	0.4	0.4	716.3	0	92.1	0	123.8	4.7	335.8	0	0.4	0.4
CO ₂	kg/h	660.6	660.6	71.6	71.3	500.0	0	0	1.1	49.3	37.8	3086.7	11633.4	72.4	72.4
CH4	kg/h	381.8	416.7	8.3	8.3	347.1	0	0	0.2	34.6	26.5	1728.9	0	8.5	8.5
CO	kg/h	12173.8	6087.3	48.1	48.1	5133.3	0	0	0.9	512.6	392.4	1795.9	0	49.2	49.2
$C_2-C_4^{(1)}$	kg/h	254.2	479.7	180.8	180.8	254.2	0	0	2.3	24.0	18.4	207.2	0	183.2	183.2
C5-C9 ⁽²⁾	kg/h	10.4	571.8	559.6	559.6	10.4	0	0	1.1	0.4	0.3	114.5	0	560.7	560.7
C10-C13 ⁽³⁾	kg/h	0	461.8	461.8	461.8	0	0	436.2	435.8	0.2	0.2	0.4	0	897.6	875.4
C_{14} - C_{20} ⁽⁴⁾	kg/h	0	687.1	687.1	687.1	0	15.8	685.8	685.8	0	0	0	0	1372.9	21.9
$C_{20+}(5)$	kg/h	0	1116.7	1116.7	1116.7	0	1619.7	519.2	519.2	0	0	0	0	1635.9	0
O2	kg/h	0	0	0	0	0	0	0	0	0	0	13987.2	2255.1	0	0
N_2	kg/h	0	0	0	0	0	0	0	0	0	0	45975.0	45975.0	0	0
Mass Flow	kg/h	15250.2	15250.2	7047.1	3138.2	6972.4	1635.5	1733.3	1648.3	745.0	480.3	67240.4	67240.4	4786.8	1777.7
Temperature	°C	220	220	35.0	35.0	37.1	427.4	353.5	35.0	35.0	35.0	112.1	791.3	35.7	99.1
Pressure	bar	20.0	20.0	19.9	19.7	20.0	35.0	20.0	20.0	20.0	20.0	1.0	1.0	1.5	1.3

Table 5. Simulations results for the main process streams (base-case/case 3: feed of 60 t/h; SCW reforming at 800 °C and 240 bar; dry reforming at 600 °C and 1 bar; LTFT conditions: 220 °C, 20 bar and H₂/CO ratio of 2.0).

(1) This cut has somewhat of C3; (2) this cut somewhat of C4 and C10; (3) this cut has somewhat of C9 and C14; (4) this cut has somewhat of C13 and C21; (5) this cut has somewhat of C20 (depending on separation in distillation)

Table 6. Sensitivity analysis results (biofuels and net electrical power production, carbon efficiencies, CO₂ mass flow and cogeneration water) varying aqueous phase of biooil in: (1) total concentration, (2) total mass flow rate, (3) composition.

Sensitivity analysis varying aqueous fraction of bio-oil													
		60 t/h; L	TFT condit	ions: 220 °C	; 20 bar; H ₂ /	$\mathbf{CO} = 2.00$	and C	O conversion p	oer pass: 50	%			
Aqueous fraction	Pressure	FT-diesel	FT-jet fuel	FT-gasoline	Biomethane	Carbon eff	iciency	Carbon efficien	cy Electric	ity	CO ₂ sequest	ered	Cogeneration
(wt. %)	turbine (bar)	(kg/h)	(kg/h)	(kg/h)	(kg/h)	without re	fining	with refining	(kW)	-	(kg/h)		water (t/h)
35 (Case1)	18.5	2174	1420	737	91	40.99		36.84	5441		10720		366.8
35 (Case1)	17.0	2174	1420	737	-	40.99	9	36.31	5656		10720		366.8
30 (Case2)	40.0	1818	1188	593	-	40.0	1	35.26	4030		8265		355.7
25 (Case 3)	40.0	1374	898	467	-	36.20	5	32.18	5728		5786		314.1
20 (Case 6)	40.0	824	538	268	-	27.19	9	23.95	7253		4009		301.2
15 (Case 7)	40.0	255	167	81	-	11.22	2	9.85	8682		1540		273.3
Sensitivity analysis varying total mass flow													
25 wt. %; LTFT conditions: 220 °C; 20 bar; H ₂ /CO = 2.0 and CO conversion per pass: 50 %													
Total mass flow	Pressure	FT-diesel	FT-jet fuel	FT-gasoline	Biomethane	Carbon eff	iciency	Carbon efficien	cy Electric	ity	CO ₂ sequest	ered	Cogeneration
(t/h)	turbine (bar)	(kg/h)	(kg/h)	(kg/h)	(kg/h)	without re	fining	with refining	(kW)	-	(kg/h)		water (t/h)
60.0	40.0	1374	898	467	-	36.26		32.18	5728		5786		314.1
100.0	40.0	2289	1496	779	-	36.20	5	32.18	9566		9643		502.2
20.0	40.0	458	299	156	-	36.20	5	32.18	1908		1911		108.0
		Sen	isitivity ana	lysis varying	individual c	ompounds	of aqu	leous fraction	of bio-oil				
		60 t/h; 25 wt.	%; LTFT	conditions: 2	20 °C; 20 ba	r; H2/CO =	= 2.0 ar	nd CO convers	ion per pass	: 50 %	0		
5 71 % (max)	357% (min)	Pressure	FT-diesel	FT-jet fuel	FT-gasoline	Carbon eff	iciency	Carbon efficien	cy Electric	ity	CO ₂ sequest	ered	Cogeneration
5.71 70 (IIIax)	5.57 % (iiiii)	turbine (bar)	(kg/h)	(kg/h)	(kg/h)	without re	fining	with refining	(kW)		(kg/h)		water (t/h)
AC	LV-HY	40.0	1374	898	467	36.20	5	32.18	5728		5786		314.1
LV	AC-HY	40.0	1397	913	470	36.59	9	32.41	5680		5863		320.7
HY	AC-LV	40.0	1435	938	483	37.3	1	33.06	5618		5893		325.0
	Sensitivit	y analysis val	rying indivi	dual compou	inds of aqueo	ous fraction	n of bio	o-oil (10-5-5-5	wt.% AC-B	U-HY	-LV (GL))		
		60 t/h; 25 wt.	%; LTFT	conditions: 2	20 °C; 20 ba	r; H2/CO =	= 2.0 ar	nd CO convers	ion per pass	: 50 %	6		
Case-study	Pressure	FT-diesel	FT-jet fuel	FT-gasoline	Carbon ef	ficiency	Carbon	efficiency with	Electricity	CO ₂	sequestered	Cog	eneration water
Case-study	turbine (bar)	(kg/h)	(kg/h)	(kg/h)	without r	efining	1	efining	(kW)		(kg/h)		(kg/h)
Base-case	40.0	1374	898	467	36.2	26		32.18	5728		5786		314.1
AC-BU-HY-LV	40.0	1429	933	470	38.0)4		33.57	5724		5657		321.2
AC-BU-HY-GL	40.0	1378	901	451	37.3	8		32.96	5873		5500		320.1

4.1. Effect of the concentration of bio-oil aqueous phase

The overall energy demand in the process increases significantly as the organic feed concentration decreases, so the flow rate of product gas derived to furnace must increase for achieving energy self-sufficiency. Thus, for 15 wt.%, 73.5% of product gas must be burnt; for 25 wt.% that fraction decreases to 18.9%, and for 35 wt.% there is no need of burning product gas. Consequently, at low concentrations, the specific biofuels production (in relation to feed) is clearly reduced, and carbon efficiencies after refining are low, in special for 15 wt. % where only 9.85% of total carbon fed into plant is present in biofuels leaving the process. For the base-case, carbon efficiency after refining is 32.18 %, typical of CTL-FT synthesis [43], and maximum carbon efficiency is achieved for 35 wt.% (36.84 % including pure biomethane or 36.31 % accounting for only liquid biofuels).

On the contrary, the electricity is higher for cases with more diluted aqueous fraction (8682 vs 4030 kW for 15 and 30 wt.%, respectively), despite the higher electrical power generated in the second turbine when the biofuels production is larger (at high feed concentration), because the flow rate of generated steam increases as the heat released from FT reactor is higher. As the increase in power of the first turbine for more diluted aqueous fraction is almost irrelevant, the higher net electrical power for diluted concentrations is due to the less consumed electrical power because of the reduced gas flow rates to compress in different points throughout the plant.

For 35 wt.% there is heat surplus in the overall process and no fraction of product gas must be sent to furnace at the PSA1 inlet, even despite the higher energy demand in the DR reactor and the upgrading of FT products (mainly, in distillation columns DEST1 and DEST2). Indeed, this fact limits the outlet pressure of the first turbine, in such a way that 40 bar cannot be set and pressure must be reduced to either 18.5 bar, thus obtaining

91 kg/h of biomethane and 5441 kW, or to 17.0 bar, so no biomethane is produced but more net power electricity is obtained (5656 kW). Nevertheless, the difference is not significant.

Lastly, the organic feeding concentration also influences on the tuning of H₂/CO molar ratio at the inlet LTFT because the syngas from SCWR reactor varies in both its flow rate and its composition. This way, for more diluted feed aqueous fractions, the hydrogen flow-rate leaving the FT synthesis loop through the purge is progressively lower and a higher fraction of hydrogen from the PSA1 outlet must be sent for hydrocracking. In addition, reforming gas from SCWR by-passes the HWGS reactor for 15-25 wt.% cases, because the CO flow rate, which determines the FT liquid production, is quite low. However, for 30 and 35 wt.%, a fraction of CO must be converted into H₂ by HWGS reactor, so the by-pass is reduced to 78.0 and 57.5%, respectively. Besides, for these latter two cases no H₂-rich stream leaving PSA1 is derived to hydrocracker.

4.2. Effect of mass flow rate and composition of bio-oil aqueous phase

Three real mass flow rates of bio-oil aqueous fraction were considered in this paper (20, 60 and 100 t/h). As expected, biofuels and electricity production vary in a proportional way, and carbon efficiencies with or without refining are the same for three cases. This happens because the plant is not sized, and in fact, the analysis of the effect of mass flow rate on the process performance in a sized plant is more complex. However, these results are interesting since the feed flow rates simulated were real and the plant capacity should be taken into account when applying economy of scale, that is, a larger scale may allow a relative reduction in capital costs, as will be discussed in the next work.

On the other hand, feed composition was changed by varying concentrations of the three most representative compounds (acetic acid, levoglucosan and hydroxyacetone). As can be seen in **Table 6**, higher carbon efficiencies and biofuels production (33.06% and

2856 kg/h) are possible in comparison with other cases (32.18% and 2739 kg/h, for basecase) if hydroxyacetone concentration, instead of acetic acid, is the maximum (5.71 wt.% in the base case). The reason is the higher number of carbon atoms fed into the process that leads to a higher molar flow-rate of gases (hydrogen, carbon dioxide and methane) generated in SCWR, which are processed through the PSAs and DR reactor, where H₂ and CO production is increased and, hence, syngas molar flow-rate fed into the FT reactor is higher. Electricity production and cogeneration water values are similar with respect to those for the base-case (5618 kW and 325 t/h vs 5724 kW and 314 t/h).

On the other hand, another possible aqueous fraction with the model compounds, previously studied in an experimental way [42], was simulated: 10-5-5-5 wt.% of acetic acid, hydroxyacetone, 1-butanol and levoglucosan (or glucose), respectively. Regarding the base-case, similar results were obtained in biofuels and electricity production, although slightly higher biofuels production was obtained for the case in which levoglucosan was fed into the process (**Table 6**).

4.3. Effect of the temperature of the LTFT reactor

When increasing the temperature in LTFT (from 220 °C to 230 and 240 °C), CO conversion per pass increases (assuming 60 and 70 %, respectively, as reported [4, 45, 46]) and, consequently, carbon efficiency also increases with respect to LTFT synthesis loop (carbon efficiency without refining). However, overall carbon efficiency (accounting for the FT product refining and upgrading) gradually decreases from 32.18 % (base-case) to 31.38 and 30.19 % for 230 and 240 °C, respectively (**Figure 4**). The reason is that chain growth probability decreases from 0.900 (base-case) to 0.870 (230 °C) and 0.839 (240 °C), so heavier hydrocarbon (wax and diesel) production decreases while light hydrocarbon (tail gas and gasoline) production increases. As a middle cut, jet fuel production decreases very slightly with temperature (898, 886 and 857 kg/h for 220,

230 and 240 °C, respectively). Besides, as temperature increases, cooling water flow rate increases and electricity production slightly decreases (**Figures 5-7**), thus concluding that the increase of temperature in LTFT reactor is not beneficial.



<u>Base case</u>: 220°C, 20 bar, 2.0 ratio, 25 wt.%; for any case, an operating parameter is changed. <u>MET</u>: similar to base case but in the alternative of burning methane instead of a fraction of product gas These notes must be also applied to Figures 5-7

Figure 4. Biofuels production (kg/h) for base case/case 3 (25 wt. %), FT temperature of 230-240 °C, FT pressure of 30-40 bar and FT inlet H₂/CO molar ratio of 2.30-1.70, as well as case 2 (30 wt. %) and methane derived to furnace (25-30 wt. %).



Figure 5. Net electricity production (kg/h) for base case/case 3 (25 wt. %), FT temperature of 230-240 °C, FT pressure of 30-40 bar and FT inlet H₂/CO molar ratio of 2.30-1.70, as well as case 2 (30 wt. %) and methane derived to furnace (25-30 wt. %).



Figure 6. Cogeneration water (kg/h) for base case/case 3 (25 wt. %), FT temperature of 230-240 °C, FT pressure of 30-40 bar and FT inlet H₂/CO molar ratio of 2.30-1.70, as well as case 2 (30 wt. %) and methane derived to furnace (25-30 wt. %).



Figure 7. CO_2 for sequestration (kg/h) for base case/case 3 (25 wt. %), FT temperature of 230-240 °C, FT pressure of 30-40 bar and FT inlet H₂/CO molar ratio of 2.30-1.70, as well as case 2 (30 wt. %) and methane derived to furnace (25-30 wt. %).

4.4. Effect of the pressure of the LTFT reactor

The increase in the LTFT pressure involves higher partial pressures of H_2 and CO, and, hence, an increase in the CO conversion per pass, which were estimated using Arrhenius kinetics, and orders of reaction for H_2 and CO in the rate equation of CO

consumption (+3/4 and -1/4, respectively) from Zennaro et al. (2000) [47] and Ma et al. (2014) [48]. Thus, the CO conversion per pass were 50, 54 and 57 % for 20, 30 and 40 bar, respectively. On the other hand, and as above mentioned, since Eq. (1) does not provide a relationship between pressure and α -values, these were taken from literature [23], obtaining 0.918 and 0.936 for 30 and 40 bar, which leads to a higher wax and diesel productions (1483 and 1636 kg/h at 30 and 40 bar, respectively, versus 1374 kg/h FT-diesel at 20 bar, as shown in **Figure 4**). Therefore, unlike for the temperature, an increase in the LTFT pressure leads to a higher liquid selectivity, and the chain growth probability increases. Carbon efficiency with refining increases slightly with pressure (32.18 % for base-case versus 32.28 and 32.47 for 30 and 40 bar, respectively), since more liquid biofuels are produced in LTFT. However, lower carbon efficiencies without refining (relative to the LTFT synthesis loop) were obtained, due to lower molar flow-rates of light hydrocarbons are produced in LTFT and, in consequence, a higher fraction of product gas must be sent to furnace. On the other hand, net electrical power and cogeneration water present similar results for all cases (**Figures 5** and **6**).

4.5. Effect of the H₂/CO molar ratio at the inlet of the LTFT reactor

When H₂/CO molar ratio increases in the stream entering the LTFT reactor, the equilibrium reaction between CO and H₂ is shifted to the products, thus increasing the CO conversion per pass. Values of 52.0 % and 47.5 % for CO conversion per pass were obtained (as for pressure sensitivity analysis) for 2.30 and 1.70 H₂/CO molar ratio. However, α -values computed by Eq. (1) are 0.912 and 0.888 for 1.70 and 2.30 ratio, respectively, so when increasing the H₂/CO ratio, the light hydrocarbons selectivity increases, thus decreasing the selectivity of liquid hydrocarbons. Using a ratio value of 2.30, more H₂ is needed and, consequently, more CO coming from SCWR must be converted into H₂ by HWGS. However, the fraction of product gas upstream from the

PSA1 derived to furnace is lower since the higher H₂ content and, hence, the higher chemical energy. Thus, CO entering the FT reactor increases as the H₂/CO molar ratio increases, and carbon efficiency without refining (in LTFT synthesis loop) is higher. Nevertheless, overall carbon efficiency (with refining) decreases with H₂/CO molar ratio increases (31.82 for 2.3 ratio versus 32.29 % for 1.7 ratio, as illustrated in **Figure 4**), because more light hydrocarbons (C₂-C₄) are produced in LTFT synthesis and after sent to furnace from distillation section. This reduces even more the fraction of product gas derived to furnace before entering the PSA1 unit. Likewise, lower net electrical power is obtained as the H₂/CO molar ratio increases (5478 kW for 2.3 ratio vs. 5832 kW for 1.7 ratio, as depicted in **Figure 5**), due to the higher fraction of product gas processed in both DR reactor and FT reactor, which involves higher power compression consumptions.

4.6. Optimal conditions for maximum biofuel and electricity production

After analyzing the results from sensitivity analysis, optimal conditions that maximize biofuels production (FT-diesel, FT-jet fuel and FT-gasoline) and provide the maximum net electrical power, accordingly to the biofuels production, corresponds to maximum aqueous phase concentration (35 wt.%), minimum LTFT temperature (220 °C), maximum LTFT pressure (40 bar) and minimum H₂/CO ratio in LTFT reactor (1.70), within the studied operating window.

4.7. Alternative configurations

Different configurations of the process were also studied, analyzed and compared with the proposed flowsheet. First, an alternative system consists of passing the entire product gas through the first PSA system and sending a fraction of the methane and carbon dioxide from PSA2 directly to the PSA3 unit, so pure methane is derived to furnace and pure carbon dioxide is sequestered. This way, less CH₄ and CO₂ are fed into DR reactor to produce H₂ and CO, and the flow rate at the inlet of the FT synthesis loop is reduced. A sensitivity analysis was carried out for 25 and 30 wt. %, and the results are shown in **Figure 4-7**. For 25 wt. % similar results were obtained in both options (combustion of pure CH₄ versus a fraction of product gas before entering the PSA1 unit), although the option in which a fraction of reforming gas is withdrawn before entering the PSA1 unit was chosen because higher biofuels production is obtained and, conceptually, it is better to burn a product gas as soon as possible, before further treatments. However, both net electricity and CO₂ flow rate for sequestration present higher values for the alternative option. Besides, the results also affect the equipment sizes, because lower mass flow rates are treated in PSA1, COMP1, PSA2 and PSA3 for the chosen option, but higher amounts of gases will pass through COMP2, COMP3, PSA1A, PSA2A and the DR reactor. Both options will be further studied in a future techno-economic analysis. For 30 wt.%, the difference between biofuels production is higher than for 25 wt.%, where the chosen option provides 3599 kg/h of biofuels versus 3443 kg/h in the alternative option (**Figure 4**), while net power and pure carbon dioxide (**Figures 5**, 7) are higher for this latter option (4467 kW and 9320 kg/h CO₂ versus 4030 kW and 8265 kg/h CO₂).

On the other hand, the sale of low-pressure steam (8 bar) was considered as an alternative to expand the steam in the second turbine to 1 bar, which could be interesting if the process was located close to other industrial plants demanding steam as utility. For this case, outlet pressure of second turbine is specified in 8 bar, and lower cooling water flow-rate is necessary, which is an environmental advantage. However, electrical power in second turbine is quite reduced and total benefits could be likely lower, so this was not the chosen option. This option might be more interesting for higher total organic concentrations in the feed stream (19460 and 9300 kg/h of steam are produced for 30 and 20 wt.%, respectively), although the chosen option allows maximizing the net electrical

power (for 30 wt.%, 2524 kW if saturated steam is considered versus 4030 kW when this potential utility is not produced).

4.8. Comparison with other published results

A comparison with results from other studies was performed (**Table 7**) in terms of thermal energy efficiency, defined by Eq. (5) accounting for the LHV (lower heat value) of all the FT liquids after refining, or overall energy efficiency that also considers the net electrical power, \dot{W}_{net} , by Eq. (6), with respect to the energy content in the feed compounds:

Thermal energy efficiency:
$$\eta_{Th-energy} = \frac{\sum_{i} \dot{m}_{Biofuel-i} LHV_{Biofuel-i}}{\sum_{j} \dot{m}_{Feed-j} LHV_{Feed-j}}$$
 (5)

Overall energy efficiency:
$$\eta_{Ov-energy} = \frac{\dot{W}_{net} + \sum_{i} \dot{m}_{Biofuel-i} LHV_{Biofuel-i}}{\sum_{j} \dot{m}_{Feed-j} LHV_{Feed-j}}$$
 (6)

Sensitivity analysis by varying aqueous fraction of bio-oil:													
60 t/h; LTFT conditions: 220 °C, 20 bar, H ₂ /CO of 2.0 and 50 % CO conversion per pass													
15 wt. % 20 wt. % 25 wt. % 30 wt. % 35 wt. %													
	10	20	20		(17.0 bar)	(18.5 bar)							
Overall energy efficiency (%)	34.77	47.75	55.10	56.24	58.76	59.38							
Thermal energy efficiency (%)	14.39	34.99	47.02	51.51	53.07	53.95							
Thermal liquid biofuel efficiency (%)	14.39	34.99	47.02	51.51	53.07	53.07							
Sensitivity a	nalysis at 60	t/h and 25 w	rt. % (base ca	se), by varyi	ng:	I							
LTFT temperature (23	0-240 °C), pro	essure (30-40	bar) and H ₂ /	CO molar rai	tio (1.70-2.30))							
	230 °C	240 °C	30 bar	40 bar	1.70	2.30							
Overall energy efficiency (%)	53.97	51.96	55.00	55.45	55.33	54.25							
Thermal energy efficiency (%)	45.94	44.26	47.08	47.23	47.12	46.53							
Sensitivity analysis by var	ying <i>individu</i>	ial compound	ls of aqueous	fraction of b	io-oil and des	ign:							
60 t/h; 25 wt. %; LTFT cond	itions: 220 °C	C; 20 bar; H ₂	/CO = 2.0 an	d CO conver	sion per pass	: 50 %							
	Case 4	Case 5	Case 8	Case 9	25 wt. %	30 wt. %							
			0		MET (*)	MET (*).							
Overall energy efficiency (%)	55.13	58.58	57.39	48.97	54.66	54.94							
Thermal energy efficiency (%)	47.47	50.17	48.24	41.63	46.18	49.29							

Table 7. Comparison with results of other studies

(*): 25 and 30 wt. % MET: cases where methane is derived to furnace when total aqueous phase is 25 wt. % and 30 wt. % respectively

In this study, overall energy efficiency varied mainly with the aqueous fraction concentration, so the minimum value was 34.77 % (for 15 wt.%) and the maximum was 59.38 % (for 35 wt.%). Likewise, regarding the base case (55.10 %), similar results were obtained (from 54.25 to 55.45 %) as LTFT pressure and H₂/CO ratio varied, but overall energy efficiencies were lower when LTFT temperature increased to 240 °C (51.96 %).

In a previous study by Siew et al. [6], using an aqueous solution of acetic acid, acetol and guaiacol (69.85 wt.%), and operating the FT reactor at 240 °C and 25 bar, maximum overall efficiency was 58.2 %, even considering all C₅₊ compounds in the computation of energy efficiency of the process and the recovery of the low-grade heat into useful byproducts such as hot water. In another study by Selvatico et al. [2], overall energy efficiency was 43.7 % based on a clean syngas, using 220 °C, 25 bar and a H₂/CO molar ratio of 2.0 as LTFT operating conditions. However, they did not take advantage of the generated steam from LTFT exothermic reactions and CO2 was not converted into H2 and CO to increase the syngas mass-flow. Likewise, Xiang et al. [49] proposed a process that integrated coal-to-syngas and syngas-to-olefins, where overall energy efficiency was 33.9-48.4 %. These researchers did not try to diminish CO₂. Similarly, Sudiro et al. [3] achieved an overall energy efficiency of 56.4 %, with synthetic fuels from two different feedstocks (coal and natural gas), operating the LTFT reactor at 240 °C and 15 bar. Finally, Leibbrandt et al. [50] obtained an overall energy efficiency of 59 % in the production of liquid FT and electricity by biomass gasification (at 40 bar and 1700 K), where the LTFT reactor conditions were 260 °C, 23.2 bar and 2.0 H₂/CO molar ratio.

Regarding the different liquid biofuels cuts (gasoline, diesel and jet fuel), diesel and jet fuel correspond to distillate stream in other studies, while gasoline is sometimes referred as naphta. Thus, Selvatico et al. [2] obtained component fractions of 11.49-17.96

% for naphta (C₅-C₁₀), 71.25-77.86 % (C₁₁-C₂₂) for distillate and 11.05-11.40 % (C₂₂₊) for wax. The rest was C₁-C₄. By considering the same cuts, similar results were obtained in our study although somewhat higher for distillate (78.01 %) and naphta (21.99 %). In another publication, Er-rbib et al. [26] concluded that their process could produce synthetic fuels composed of 72 % of diesel (C₁₂-C₁₈), 26 % of gasoline (C₅-C₁₁) and 2 % of C₂-C₄. By considering these cuts, very similar results were obtained when the LTFT reactor operates at 30 bar (70.48 % for diesel and 29.52 % for gasoline).

5. Conclusions

A new and comprehensive energy self-sufficient process aimed at maximizing biofuels and net power production from low-temperature Fischer-Tropsch (LTFT) synthesis using syngas obtained by supercritical water reforming (SCWR) of bio-oil aqueous phase was proposed and studied. By this process, a waste is valorized, and the produced CO₂ is separated with a high purity for sequestration, which gives added value to the proposed process. Optimal conditions were found by analyzing the effect of different operating variables regarding the bio-oil aqueous phase (concentration, composition and mass-flow in the feeding stream) and the operation of LTFT reactor (temperature, pressure and H₂/CO molar ratio) on the process performance. This was assessed by carbon efficiencies with and without refining, as well as energy efficiency that includes net electrical power. The results show that biofuels production (FT-diesel, FT-jet fuel and FT-gasoline) is maximized at maximum aqueous phase concentration (35 wt.%), minimum LTFT temperature (220 °C), maximum LTFT pressure (40 bar) and minimum H₂/CO molar ratio in LTFT reactor (1.70), within the studied operating window. Under these conditions, for a mass flow of aqueous phase of 60 t/h, biofuels production was 4596 kg/h (2804 kg/h FT-diesel, 1491 kg/h FT-jet fuel and 301 kg/h FTgasoline), a carbon efficiency with refining of 38.53 % was achieved (without refining, it

was 43.50%), and net electrical power was 5297 kWe. In addition, the separated CO_2 for sequestration is 0.50 kg/kg organic feeding and 17.5 kg hot water/kg organic feeding, which gives added value to the proposed process.

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