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REVIEW ARTICLE



Challenges and opportunities of process modelling renewable advanced fuels

Lorenzo Testa¹ · David Chiaramonti^{1,2} · Matteo Prussi¹ · Samir Bensaid¹

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Abstract

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The Paris COP21 held on December 2015 represented a step forward global GHG emission reduction: this led to intensify research efforts in renewables, including biofuels and bioliquids. However, addressing sustainable biofuels and bioliquid routes and value chains which can limit or reverse the ILUC (indirect land-use change effect) is of paramount importance. Given this background condition, the present study targets the analysis and modelling a new integrated biomass conversion pathway to produce renewable advanced fuels, enabling the issue of indirect land-use change (ILUC) of biofuels to be tackled. The bioenergy chain under investigation integrates the decentralized production of biogas through anaerobic digestion and its upgrading to biomethane, followed by a centralized conversion to liquid transport fuels, involving methane reforming into syngas, Fischer-Tropsch (FT) synthesis, and methanol synthesis. The methodology adopted in this work stem from extensive literature review of suitable bio/thermo-chemical conversion technologies and their process modelling using a commercial flow-diagram simulation software is carried out. The major significance of the study is to understand the different modelling approaches, to allow the estimation of process yields and mass/energy balances: in such a way, this work aims at providing guidance to process modellers targeting qualitative and quantitative assessments of biomass to biofuels process routes. Beyond FT products, additional process pathways have been also explored, such as MeOH synthesis from captured CO₂ and direct methane to methanol synthesis (DMTM). The analysis demonstrated that it is possible to model such innovative integrated processes through the selected simulation tool. However, research is still needed as regards the DMTM process, where studies about modelling this route through the same tool have not been yet identified in the literature.

Keywords Biogas · Modelling · Biofuels · Biomethane · Fischer–Tropsch · Methanol · Conversion · BTL · Anaerobic digestion · Reforming · Syngas · Upgrading · Biomass · Aspen plus · Process modelling · Renewable fuels

| Abbreviati | ons | NRTL | Non-random two liquid | | |
|---|--|--------------------|--|--|--|
| AD | Anaerobic digestion | BRD | Biogas done right | | |
| MDEA | Methyldiethanolamine | NRTL-RK | Non-random two-liquid Redlich-Kwong | | |
| ADM1 | Anaerobic Digestion Model No.1 | BWR | Boiling water reactor | | |
| MEA | Monoethanolamine | PAD | Pressurized anaerobic digestion | | |
| AMP | Amino-methyl-propanol | CHP | Cogeneration or combined heat and power | | |
| MeOH | Methanol | POX | Partial oxidation | | |
| ASF | Anderson-Schulz-Flory | CIB | Consorzio Italiano Biogas | | |
| MTBE | Methyl tertiary-butyl ether | PR-BM | Peng Robinson with Boston-Mathias | | |
| ATR | Autothermal reforming | | alpha function | | |
| | | COP21 | 2015 UN Climate Change Conference | | |
| ☐ David Chi | ioromonti | PRF | Plug flow reactor | | |
| | ramonti@polito.it | CSTR | Continuous stirred tank reactor | | |
| un national e ponto.n | | PSA | Pressure swing adsorption | | |
| Galileo Ferraris" Energy Department, Politecnico Di Torino, | | CTL Coal to liquid | | | |
| Corso Due | ca Degli Abruzzi 24, 10129 Turin, Italy | PWS | Pressurized water scrubbing | | |
| | e Energy Consortium for Research | DEA | Diethanolamine | | |
| | opment (RE-CORD), Viale Kennedy 182, arperia e San Piero, Italy | RCSTR | Rigorous continuous stirred tank reactor | | |



DME Dimethyl ether

REQUIL Rigorous equilibrium reactor based on

stoichiometric approach

DMR Dry methane reforming

RGIBBS Rigorous equilibrium reactor based on

Gibbs free energy minimization

DMTM Direct methane to methanol

RK Redlich-Kwong

ELECNRTL Electrolyte non-random two liquid **RKSMHV2** Redlich-Kwong-Soave equation of state

with modified Huron-Vidal mixing rule

EOS Equation of state

RPLUG Rigorous plug flow reactor with rate-con-

trolled reactions based on known kinetics

FT Fischer-Tropsch

RSTOIC Stoichiometric reactor based on known

fractional conversions or extents of

reaction

GHG Greenhouse gases
RWGS Reverse water–gas shift

GTL Gas to liquid

RYIELD Nonstoichiometric reactor based on

known yield distribution

HPWS High-pressure water scrubbing SAF Sustainable aviation fuels

HTFT High-temperature Fischer–Tropsch SCWR Supercritical water reforming

IGCC Integrated gasification combined cycle

SMR Steam methane reforming ILUC Indirect land-use change

SRK-ML Soave-Redlich-Kwong equation of state

with T dependent

LHHW Langmuir-Hinshelwood-Hougen-Watson

UASB Upflow anaerobic sludge blanked

LPG Liquefied petroleum gas
VFA Volatile fatty acids

LTFT Low-temperature Fischer–Tropsch

WGS Water-gas shift
MB Membrane separation

1 Introduction and scope of the work

Nowadays, fossil resources still represent the main global energy source, covering about 80% of the world's energy consumption. However, their production and use cause severe impacts on the environment, as they release carbon dioxide and other greenhouse gases (GHG), responsible for global warming and climate change: according to J. G. J. Olivier and J. A. H. W. Peters [1], fossil fuel combustion accounts for 89% of all CO_2 emissions and 68% of all GHG emissions.

Climate change is today a well-recognized global concern, and the urgent need to develop and implement alternative processes for sustainable energy generation is promoting research and development worldwide.

At the international level, a major commitment to GHG emission reduction came with the adoption of the COP21 Paris Agreement in 2015, signed by 196 Parties. This binding international treaty marked the beginning of a new global effort in contrast to climate change, setting the aim of limiting global warming to well below 2 °C (possibly 1.5 °C) compared to pre-industrial levels (intended as 1850–1900) [2]. In this context, an unprecedented boost to market deployment of renewable energies is an unavoidable component of a wider climate change fighting strategy.

Along with wind and solar energy, another key solution is the use of biomass to produce transportation fuels (biofuels) replacing conventional fossil fuels, as they provide a renewable carbon-based source, being CO₂ utilized by crops and forests during the natural photosynthesis process [3]. However, biofuel production itself could induce other landrelated emissions, either directly and/or indirectly [4]. In fact, when biofuels are produced on existing agricultural land and conventional agronomic practices, the demand for food and feed crops might lead to the extension of agricultural land into areas with high-carbon stock such as forests, wetlands, and peatlands, to provide the same amount of feed/food replaced by biofuel production. If and when this happens, it may originate greenhouse gas emissions that will negatively impact on biofuels' GHG balance [5] [6], and thus on climate. This effect is known as indirect land-use change (ILUC). ILUC is a very complex phenomenon, which accounting requires the understanding of a large number of different factors. It can be anyway contrasted by adopting sustainable agricultural rotations, photosynthetic intensification in agriculture, soil carbon accumulation, improved nitrogen and carbon use efficiency, etc.

The objective of this study is to investigate a novel biomass-to-liquid fuel value chain that has the potential to address the ILUC impact of biofuels. To this aim, the upstream part of the biofuels production chain under consideration refers to the Biogas Done Right (BDR) sustainable model developed by Consorzio Italiano Biogas (CIB), being this a set of agronomic practices that links anaerobic digestion to sustainable farming [7] to produce biogas. The scheme, among other elements, involves double cropping, with a primary crop for food or feed and a secondary one through crop rotation for energy production. The model is based on cover cropping, and contributes to improve the soil quality, as it is covered during the entire year, and reduces the use of fossil fertilizers, supporting in that way also the production of the main crop. In fact, BDR contrasts soil erosion and reduces nitrogen emissions, and increases the availability and use of organic fertilizers. By doing so, the



BDR model ensures sustainable production of biogas, representing thus a valuable solution to the ILUC issue. Among the many possible applications, the biomethane produced through the BDR model could represent a suitable raw material for the production of sustainable biofuels, making it a good potential substitute to natural gas in several refinery processes and exploiting the natural gas pipeline infrastructure. For this reason, the value chain investigated in this paper employs biomethane as a basis to produce chemical products like Fischer-Tropsch (FT) liquids and methanol (MeOH), which, in this way, turn out to be low-ILUC-risk biofuels. Hence, the identified pathway includes the biogas production and upgrade to biomethane in decentralized farms, and integrated with a centralized biomethane-toliquid conversion plants as FT-liquid fuels and MeOH. As said, this is approach is possible thanks to the availability of a natural gas infrastructure, where the biomethane produced through a decentralized distributed approach is injected, and the equivalent amount is collected at a centralized biorefinery site where the gas is processed into a liquid.

The natural gas network would thus be used the same way electricity from photovoltaics uses the grid. This represents a very innovative use of renewables in the current context, valorizing the use of existing infrastructures.

In brief, the key process steps of the selected value chain would be as follows: (i) decentralized biomethane production, based on sustainable models as the Biogas Done Right, (ii) injection of biomethane into the natural gas grid, (iii)

extraction from the same volume of natural gas form the gas grid, and processing in centralized biomethane-to-liquid conversion plants, through FT synthesis and MeOH synthesis. The conceptual scheme of the whole process steps is shown in Fig. 1.

An extensive review of the conversion technologies that could be adopted in the biogas to Fischer-Tropsch fuels and MeOH production chains has been performed in this work, combined with a critical and extensive literature review on modelling the various steps identified for the selected conversion processes. The modelling approaches (assumptions, schematization, process conditions, etc.) adopted in the various papers reviewed are extensively analysed and commented. The final goal is to provide a complete set of information to study, through modelling, the efficiency, and the techno-economic feasibility of these integrated bio/ thermo-chemical processes, allowing to optimize operational conditions, determine limitations, and design the distributed + centralized scheme here proposed. The current work focuses on process modelling, while the techno-economic, potential, and regulatory modelling of distributed biomethane production and centralized conversion to advanced fuels will be the subject of a subsequent research paper.

To our knowledge, no systematic review of studies encompassing all the steps of the selected value chain and their modelling has yet been published. Besides, the elements of this paper comprehensively cover an insightful analysis of a broad spectrum of industrial and novel

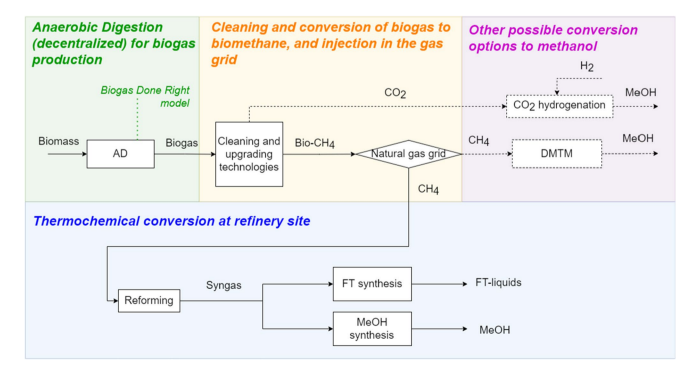


Fig. 1 FT fuels and MeOH production chain scheme (AD=anaerobic digestion; FT=Fischer-Tropsch; MeOH=methanol; DMTM=direct methane to methanol)



biogas-to-liquid conversion technologies, co-product valorization, and modelling through a commercially available simulation software.

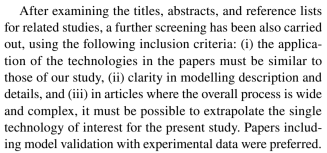
2 Materials and methods

As previously discussed, the present work investigates the state-of-the-art technology of biogas-to-liquid pathways and reviews studies modelling those technologies using a commercial simulation software by discussing essential modelling steps.

A systematic literature review of peer-reviewed journal articles and book chapters has been carried out, exploring databases such as Springer (International Journal of Energy and Environmental Engineering (IJEEE), Clean Technologies, and Environmental Policy) and Elsevier (Energy, Applied Energy, Environmental Chemical Engineering, Bioresource Technology, Energy Conversion and Management, Fuel, Chemical Engineering Research and Design, International Journal of Hydrogen Energy, Chemical Engineering & Technology, Sustainable Energy Technologies and Assessments, Fuel Processing Technology, Journal of Cleaner Production), and journals such as Energies, Chemical Engineering Transactions (CET), Energy Fuels, Fuel, Molecular Catalysis, Biomass Conversion and Biorefinery, Chemie Ingenieur Technik, Gas Processing Journal, International Journal of Chemical Reactor Engineering, Journal of Advanced Research in Biofuel and Bioenergy, and Journal of the Japan Institute of Energy. Conference papers have been also included in the research, for instance Energy Procedia, IOP Conference Series: Materials Science and Engineering, and papers of the 23rd European Biomass Conference and Exhibition. Thesis works developed within several universities have been also considered, such as Politecnico di Torino, University of Southern Denmark, University of Padua, Chalmers University of Technology, Norwegian University of Science and Technology.

Searches were limited to articles published in the last two decades, written in English, Italian, and German.

As regards the modelling papers, the first step was to choose a simulation software able to simulate all the technologies included in the identified value chain. Currently, there are lots of process modelling software packages available and used in papers in the literature, such as DWSIM, Engineering Equation Solver, COCO Simulator, Engineering Base by AUCOTEC, Aspen Plus, Aspen HYSYS, and GateCycle Software. The software chosen for this analysis is Aspen Plus, being it one of the most used process simulation tools for both industrial and research applications. The search for modelling studies has therefore been restricted to those using the selected commercial software.



The modelling studies collected have been then deeply analysed, focusing on information such as process configuration, modelling assumptions, process yield, and energy consumption of the system. Information was harmonized, i.e. data were converted to common units, to allow for effective comparisons.

3 Results

Based on the methodology described in the previous section, a comprehensive review of the conversion technologies to be applied for biogas to liquid pathways has been carried out, and 57 studies (out of 94 collected) using Aspen Plus to model them have been selected and analysed.

In this section, the findings of the present review are presented. Specifically, the paragraph is split into two main parts: in the first one, the identified biogas-to-liquid technical solutions are outlined, providing an overview of the key process parameters, whereas in the second one, the selected 57 modelling studies are described in detail, evaluating key modelling components. Challenges in process sustainability and advances are also covered.

3.1 Biogas-to-liquid pathways

3.1.1 Anaerobic digestion

Biogas is produced by the anaerobic digestion (AD) biological process. This complex microbiological path is based on the work of several groups of bacteria and archaea, working in synergy to decompose organic matter into a mixture of $\mathrm{CH_4}$ (53–70% vol), $\mathrm{CO_2}$ (30–50% vol), $\mathrm{N_2}$ (2–6% vol), $\mathrm{O_2}$ (0–5% vol), and lower fractions of $\mathrm{H_2}$, $\mathrm{H_2S}$, and $\mathrm{NH_3}$ [8], depending on the type of biomass digested and the process conditions. Substrates could be dedicated crops, agricultural residues, food and household wastes, animal manure, industrial wastes, etc.

The AD process consists in four crucial phases, i.e. hydrolysis, acidogenesis, acetogenesis, and methanogenesis. During the hydrolysis step, the complex substrates mainly composed by carbohydrates, fats, and proteins are hydrolyzed into their respective monomers, i.e. glucose, fatty acids, and amino acids. These monomers are then



turned into volatile fatty acids (VFA), i.e. valeric, butyric, caproic, iso-valeric, iso-butyric, propionic, and acetic during the acidogenesis, and subsequently converted into acetic acid, hydrogen, and carbon dioxide (acetogenesis). Finally, methanogenesis holds the conversion of these products into methane and carbon dioxide [10]. The end-products of AD also include a slurry or solid fraction consisting of what is left of the treated substrate, referred to as digestate, rich in organic carbon and nutrients [11].

The trophic chain depends on several operational factors, such as temperature, redox potential, pH, feeding procedure, mixing, retention time, type of substrates, reactor configuration, organic loading rate, and inhibitors [12].

3.1.2 Biogas cleaning and upgrading

Biogas is by far constituted by CH₄, vapour, and CO₂, with other contaminants such as H₂S, NH₃, N₂, and siloxanes, whose concentration largely depends upon the composition of the substrate digested [23]. These impurities need to be reduced, as they can cause corrosion, are toxic, can cause catalyst deactivation, and reduce gas heating value, but also to meet gas specifications and standards. Additional to the direct use in an engine for power production, in fact, biomethane can be injected into the natural gas grids. Therefore, many countries formulated standards to ensure gas quality before the injection of biomethane into the natural gas grid [24]. In Italy, the biomethane quality for injection is regulated by the Decree of the Ministry for Economic Development of 19th of February 2007, the M/475 Mandate to CEN for standards for biomethane for use in transport and injection in natural gas pipelines, and the technical report UNI/ TR 11,537:2019 (updated version of UNI/TR 11,537:2016).

Typically, the purification from contaminants is referred to as "biogas cleaning", while the CO_2 and steam removal process is called "upgrading" [25]. Although some upgrading technologies separate both impurities and CO_2 , it can be advantageous to clean the gas before upgrading; details about biogas cleaning can be found in [11].

Currently, there are several technologies for biogas upgrading, which are continually improved, while new techniques are under development [26]. The most widespread and consolidated technologies for biogas upgrading are (i) physical absorption, using water or organic solvents, (ii) chemical absorption, using amine or saline solutions, (iii) pressure swing adsorption (PSA), (iv) membrane separation (MB), or (v) cryogenic upgrading [27].

Physical absorption techniques (i) exploit the different solubility of CH₄ and CO₂ (and possibly other trace compounds) in the absorbent liquid: raw biogas meets a counter flow of liquid in an absorption column, thus the leaving liquid will contain CO₂ while the remaining gas stream will have an increased concentration of methane. The absorbent

medium can be water, i.e. pressurized water scrubbing (PWS) technology [28], or organic, in organic solvents such as polyethylene glycol, in which CO_2 is more solvent than in water. This technique is able to clean impurities such as H_2S , NH_3 , siloxanes, and halogenated components as well [20].

In chemical absorption technologies (ii), CO₂ is not only absorbed in the liquid, but also reacts chemically with the employed solutions, which can be amine, such as MEA (monoethanolamine), MDEA (methyldiethanolamine) or DEA (diethanolamine), or saline (e.g. K₂CO₃ solutions).

In the PSA process (iii), specific porous materials (usually activated carbon or zeolites) and high pressures are employed to adsorb CO_2 , but also O_2 , N_2 , and other biogas trace components, which are then released at lower pressure. When using this technique, the removal of H_2S and H_2O in the raw gas is essential, as they can damage the adsorbing material.

Membrane separation (iv) consists in using slight layers of materials that are permeable to CO_2 , H_2O , and NH_3 , but not to CH_4 , which thus gets separated from the other impurities.

Cryogenic upgrading (v) is based upon the principle that, at a fixed pressure, different gases liquefy at different temperatures. This technology has seen strong interest and development in recent years.

In Europe, 35.5% of the plants employ PWS, while the 20% MEA, the 20% MB, the 17% PSA, and the remaining 8% adopt emerging technologies [24].

After cleaning and upgrading, the gas stream is called biomethane, which is a renewable source of methane $(CH_4 > 95\%, CO_2 \text{ from 1 to 5\%})$, and can be directly used as automotive fuel or injected into the natural gas grid. [29]

3.1.3 Biomethane reforming to syngas

Both the production routes here considered to generate Fischer–Tropsch fuels and methanol are fed with syngas: therefore, in our scheme reforming of biomethane to syngas is necessary, representing a critical step of the process(Fig. 2).

Synthetic gas, or syngas, is a gaseous mixture of H₂ and CO, at different ratios, that can be used as a chemical building block for the synthesis of a variety of chemical products and carbon-based fuels. The selectivity of the final products depends upon the H₂/CO ratio [9]. Both Fischer–Tropsch and methanol synthesis requires H₂/CO ratio equal to 2 [10] [11]. Syngas is traditionally obtained from coal, natural gas, residual oils, and petroleum, but it is possible to generate syngas also from biomass, a sustainable and renewable substitute to fossil-based syngas [12].

In this work, the syngas is obtained from biomethane, the upgraded product of anaerobic digestion of organic materials [13]. Also, it can be generated from methane extracted from the gas grid, if an equivalent amount of



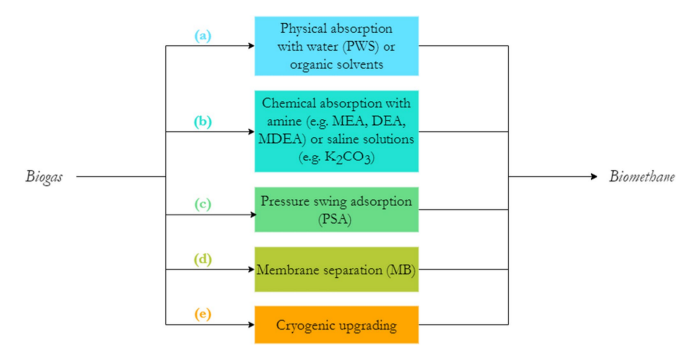


Fig. 2 Biogas cleaning and upgrading main conversion technologies

biomethane is injected into the gas pipeline elsewhere from the AD-biomethane production site (in a certified accounting mode, ensuring renewable carbon is not double counted through the use of guarantees of origin).

The main processes used to convert methane to syngas can be summarized as follows: (i) steam methane reforming (SMR), (ii) partial oxidation (POX), (iii) autothermal reforming (ATR), and (iv) dry methane reforming (DMR) ([10, 14, 15]).

SMR (i) is a well-established and large-scale technology, mostly used for hydrogen production from methane. In this route, CH₄ and steam react in a reformer over a nickel-alumina catalyst [16], at a temperature of 1073.15 to 1173.15 K and a pressure of 15 to 30 bar. The primary reaction is:

$$CH_4 + H_2O \rightarrow CO + 3H_2 \tag{1}$$

The process is strongly endothermic [17], and the resulting H_2/CO ratio is ~3, well above FT-synthesis requirements.

In typical industrial applications targeting H_2 generation via this pathway, a second reaction also occurs, producing additional hydrogen through the water gas shift (WGS) reactor. In this case, CO and water react, producing hydrogen and carbon dioxide, as in reaction (2):

$$CO + H_2O \rightarrow CO_2 + H_2 \tag{2}$$

POX (ii) instead uses oxygen to convert methane. The methane partial oxidation reaction is the following:

$$CH_4 + \frac{1}{2}O_2 \to CO + 2H_2$$
 (3)

The process is exothermic and generates H_2 -lean syngas $(H_2/CO=1 \text{ to } 1.6)$ and the reaction occurs at a high temperature (> 1473 K) if no catalyst is used [10]. The employment of a catalyst can lower the reaction temperature to ~ 1000 K. The catalysts employed could be divided into three groups: Ni, Co, and Fe, noble metal, and early transition metal carbide [18].

POX can be well combined with SMR reforming to achieve a H_2/CO ratio in the range of 1.6 to 2.6 [10]. This process is called autothermal reforming (iii). In ATR, the heat produced by the POX is used to provide the endothermic heat of the SMR reaction.

In DMR (iv), CO_2 is used as an oxidant to convert CH_4 to syngas. The technology is thus very attractive from a sustainability perspective, as it uses two types of greenhouse gases, i.e. CO_2 and CH_4 , to form a valuable product. The process is described by Eq. (3).

$$CH_4 + CO_2 \rightarrow 2CO + 2H_2 \tag{4}$$

The syngas produced is normally characterized by a H_2 /CO ratio close to 1 [10]. This could also be further adjusted for methanol and Fischer–Tropsch synthesis by reacting CO with H_2 O to produce CO_2 and H_2 in water gas shift (WGS) and partial oxidation reactions [8, 14].

Catalysts for DMR can be noble metal-based (Rh, Ru), which have good activity and stability but high cost, or



Ni-based ones (Ni/Al₂O₃), commonly used for their low cost, high H₂ yield, and fast turnover rates.

This approach is particularly interesting as it can tolerate the varying concentration of CO_2 associated with biomethane. Nevertheless, the commercialization of this technology is still in its preliminary stage [16, 19]. However, there are some drawbacks linked to these reforming routes, such as catalyst deactivation (mainly due to carbon deposition), and high energy demand, as the reforming reaction is endothermic and requires to be operated at high temperatures (1123.15 to 1273.15 K) to obtain higher conversion rates and minimize carbon deposition.

3.1.4 Fischer-Tropsch synthesis

Once syngas has been produced and purified, it can be used in the Fischer–Tropsch process to produce a mixture of hydrocarbons at different chain lengths, used as synthetic fuels (Fig. 3). These products may be used directly in a gas turbine or distilled into kerosene (C-10 to C-16), diesel-(C-14 to C-20), light naphtha (C-5 to C-6), heavy naphtha (C-6 to C-12) and waxes (C-20+).

The Fischer–Tropsch synthesis is a polymerization reaction, in which CO is hydrogenated with H₂ to the C-1 intermediate, which then grows to form different hydrocarbon chains of variable lengths. Syngas is thus converted into a variety of products, such as alcohols, aldehydes, olefins, paraffins, and especially liquid transportation fuels [20]. After the FT synthesis, the last stage is upgrading and separation of the FT syncrude in order to obtain high-quality products.

Fischer-Tropsch synthesis was developed in the early twentieth century by Franz Fischer and Hans Tropsch at the Kaiser Wilhelm Institute, with the aim of producing synthetic fuels from coal reserves in Germany during World War II. The process found only limited commercial application. This relatively well-known technology

has recently drawn renewed interest for its application to cellulosic biomass and agricultural waste, to convert them to linear- and branched-chain synthetic hydrocarbon, representing thus a very promising and sustainable solution for the production of clean fuels at competitive costs ([21–29]).

The polymerization reaction requires syngas at a H₂/CO ratio of 2–2.2 [30], which is processed over a metal catalyst (Fe or Co), at apressure range of 20 to 60 bar. Temperatures can be in the range of 473.15 to 523.15 K (low-temperature FT synthesis or LTFT), or 573.15 to 623.15 K (high-temperature FT synthesis or HTFT). In both cases, the process is highly exothermic, and therefore, a heat exchange system is necessary to cool the reactor and maintain control of the process temperature: it is also an energy-recovery opportunity for waste heat [31].

At first, reagents, hydrogen and carbon monoxide, form monomer units, which are then polymerized to yield a wide spectrum of products (mainly paraffin), ranging from C-1 to C-40 hydrocarbons.

The FT synthesis consists in four main reactions, shown in Eqs. (5) to (8), i.e.

Paraffin formation:

$$nCO + (2n+1)H_2 \to C_n H_{2n+2} + nH_2 O$$
 (5)

- Olefin formation:

$$nCO + 2nH_2 \rightarrow C_nH_{2n} + nH_2O \tag{6}$$

Alcohol formation:

$$nCO + 2nH_2 \rightarrow C_nH_{2n+1}OH + (n-1)H_2O$$
 (7)

- Carbonyl formation:

$$nCO + (2n-1)H_2 \to (CH_2)_nO + (n-1)H_2O$$
 (8)

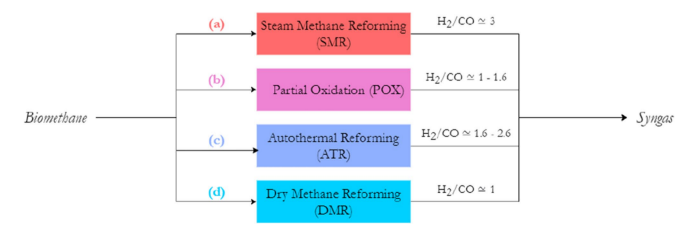


Fig. 3 Methane reforming to syngas conversion technologies

Catalysts play a crucial role in FT synthesis, as they must guarantee a good conversion yield of reactants, as well as selectivity towards products. Catalysts in FT are often supported on metal oxides, typically alumina or silica [22]. Suitable catalysts for FT synthesis are group VIII elements, in particular cobalt (Co), iron (Fe), nickel (Ni), and RUTHE-NIUM (Ru), able to chemisorb CO dissociatively (into C and O) and H₂, and have a noticeable activity. However, other elements, such as rhodium (Rh), iridium (Ir), palladium (Pd), and platinum (Pt), are also used in FT synthesis. Though the selectivity of these elements is even higher compared to Ru, Ni, Co, and Fe, they are not considered in industrial applications because of their costs [21], and only Co and Fe are used in commercial processes. Co-based catalysts are mainly used in LTFT: these are characterized by high activity, significant stability, and a tendency to produce relatively higher molar-weight hydrocarbons. On the other hand, iron-based catalysts are cheaper than Co-based ones, can be used in both HTFT and LTFT configurations, and promote a relatively higher fraction of olefins. Additionally, iron catalysts also promote the WGS secondary reaction.

There are four main different types of Fischer–Tropsch reactors: (a) fixed-bed multi-tubular reactor, (b) fluidized-bed reactor, (c) slurry-bed reactor, and (d) microchannel reactor (Fig. 4). The type of reactor influences the operational parameters of the synthesis process, the product selectivity, the product distribution with chain growth probability, the catalyst activity, and the conversion of carbon monoxide [22]. Details about the FT reactor design can be found in [32].

3.1.5 Methanol synthesis

Methanol (CH₃OH) is a valuable chemical product with a variety of uses, either as a clean fuel, mixed with other conventional fuels, or as a bulk chemical building block for the synthesis of other chemicals such as acetic acid, formaldehyde, methyl methacrylate and methyl tertiary-butyl ether (MTBE), and many others [33]. CH₃OH is extremely stable and liquid at room temperature, and this minimizes problems with storage and transportation, even if accidental release in soil and dwells can be a serious health risk ([34–37]).

Currently, the most used industrial route for methanol production is based on using syngas produced via reforming of natural gas, even if also biomethane can obviously be used. Nevertheless, there are also attractive routes that involve a single step, such as oxidative coupling of methane, e.g. methane partial oxidation to methanol (i.e. DMTM) [38], which will be discussed in a separate section of this work.

Methanol is obtained through the hydrogenation of carbon oxides over a suitable (copper oxide, zinc oxide, or chromium oxide-based) catalyst [39][39]. The conversion is exothermic and very selective, and the synthesis is followed by a distillation column to separate methanol from water, which is the by-product of the conversion [41].

The main reactions of methanol synthesis are [42]:

$$CO + 2H_2 \rightarrow CH_3OH$$
 (9)

$$CO_2 + 3H_2 \rightarrow CH_3OH + H_2O \tag{10}$$

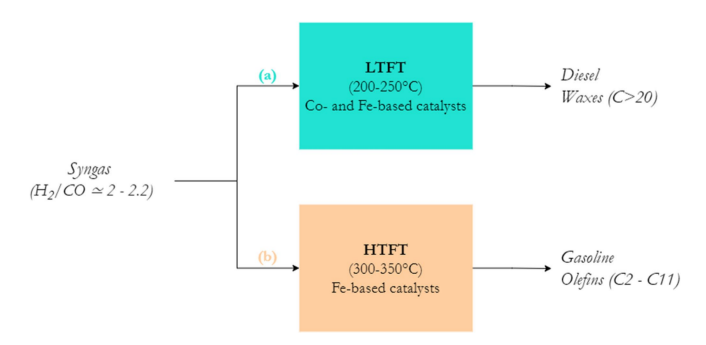


Fig. 4 Fischer-Tropsch synthesis—configurations and products



$$CO_2 + H_2 \leftrightarrow CO + H_2O$$
 (11)

Equation (9) represents the CO hydrogenation, (10) the CO_2 hydrogenation, and (11) the reverse water gas shift (RWGS) reaction. It is noted that the required H_2/CO ratio of the syngas at the inlet is equal to 2.

The typical operating conditions are in the ranges of 50 to 100 bar and 493.15 to 553.15 K, depending on the catalyst supplier.

There are several commercial types of methanol synthesis reactors, i.e. quench reactor, adiabatic rectors in series, or boiling water reactors (BWR) [43]. A detailed description of reactor types is available in [44].

3.1.6 DMTM route

As an alternative to methanol synthesis from syngas, the straight conversion of methane into methanol is also a possible and interesting route. This method allows to by-pass the very energy-intensive step required to reform CH_4 into CO and H_2 , and thus represents an economical-advantageous and environment-friendly option [45].

The technologies for the direct conversion of methane to methanol might be catalytic oxidation processes, photocatalysis technologies, plasma technologies, supercritical water oxidation technologies, membrane technologies and other methods.

Da Silva et al. [38] and Zakaria et al. [46] performed a review of the different DMTM routes.

However, to date, this method is not yet applied at full industrial and commercial scale. The process is particularly difficult, since the target product CH₃OH is more prone to oxidation than CH₄, and thus, the process needs to activate the C–H bonds on one hand, and avoid over-oxidation of CH₃OH on the other [47]. Moreover, the current technologies do not provide a relevant methanol yield [48].

4 Modelling biogas-to-liquid fuels

This review analyses and discusses the various plant configurations of the different studies, reporting, wherever possible, the yields and the consumptions of such steps that are used in the proposed pathway. Modelling approaches (assumptions, schematization, process conditions, etc.) are analysed and commented below.

4.1 Modelling anaerobic digestion

The AD process is an extremely complex process, involving numerous intermediate reaction mechanisms, such as bacterial metabolic reactions, parameters, inhibitors interactions, etc. In the literature, several models have been proposed to describe anaerobic digestion [49]. These models can be single-step models [50], involving a single bacterial population with a limited description of inhibition effects, or models of intermediate complexity [51], considering a higher number of strains of bacteria with a more accurate description of inhibition factors, or, finally, complex models [52] [53] [54] [55], entailing a high number of processes, inhibition effects, and specific bacterial populations.

Among the complex AD models, Anaerobic Digestion Model No. 1 (ADM1) [52] is considered the most comprehensive one and is widely applied for AD description [56]. ADM1, developed by the International Water Association (IWA) task group, assumes that the reaction system consists of biochemical reactions (involving enzymes) and physico-chemical reactions (involving acid-base reactions and the gas-liquid transfer). The substrate fed to the digestor is assumed to be composed of carbohydrates, proteins, and fats [57]. Another important and complete complex model is the one developed by Angelidaki et al. [55], in which the substrate is assumed to be composed of basic organic components (i.e. carbohydrates, lipids, and proteins), intermediates (i.e. volatile fatty acids and long-chain fatty acids), and inorganic components (i.e. ammonia, phosphate, cations, and anions). The model includes 2 enzymatic hydrolytic steps and 8 bacterial steps and involves 19 chemical compounds [55].

Implementing modelling schemes of AD in Aspen Plus is quite challenging, as it involves microorganisms, whose microbial activity is difficult to describe in software language. In literature, several studies simulating the anaerobic digestion in Aspen Plus have however been found and are analysed below.

Al-Rubaye et al. [56] developed a two-stage simulation model: one first step for the hydrolysis phase, and a second one for the other three phases, i.e. acidogenesis, acetogenesis, and methanogenesis. The Property Method (a collection of methods and models that the software uses to compute thermodynamic and transport properties [58]) chosen for the simulation is the non-random two-liquid (NRTL). In line with ADM1, the substrate feed rate is assumed to be made of carbohydrates, proteins, and fats, and therefore the introduction of material components is treated accordingly. The feed stream is mixed with H₂ and H₂O through a mixer and a heat exchanger is employed to model heating of this stream, necessary to maintain the required temperature in the ranges favourable for AD. The hydrolysis step is simulated using a stoichiometric reactor (RSTOIC), in which the reaction kinetics are not considered, but stoichiometry and conversion of a reactant must be specified. Thirteen chemical reactions are considered for this step, and the reaction rates have been calculated by Aspen Plus calculator blocks, using a FORTRAN code. The subsequent AD steps have been



simulated in a continuous stirred tank reactor (RCSTR), which requires the reaction kinetics to be known; thus, specifications from ADM1 and comprehensive models have been used, involving more than 33 kinetic reactions. The reaction rates have been calculated through calculator blocks, using a FORTRAN code. The RCSTR reactor releases two streams at outlet: one is the gas stream, which is the biogas and small traces of other gases, and the liquid stream. The gas stream goes through a splitter and then a flash separator, which separates the water from the biogas. Subsequently, the biogas stream passes through a gas filter, which separates the hydrogen component from the produced biogas. On the other hand, the liquid stream goes through a splitter to separate a part of it as recycle and is connected to the feed stream. The model was validated against experimental data in terms of % CH₄ in the produced biogas. Three different feed cases have been tested, i.e. (i) cattle manure, (ii) cow manure, and (iii) wastewater generated from industrial and agricultural activities. Results match the literature data; in detail, the deviation from simulation results and experimental data are 5.4% for the case of cattle manure, 8.54% for the case of cow manure, and 15.83% for the case of wastewater generated from industrial and agricultural activities. In the paper, a sensitivity analysis has also been carried out to study the effect on CH₄ yield in case of introduction of hydrogen in the process. The investigation revealed that, for H₂ feeding rates below a maximum value, there is an increase in methane gas composition in the produced biogas.

The model developed by Rajendran et al. [59] shows similarities with Al-Rubaye's one. Also, here, hydrolysis is separated from the other AD phases and has a separate reaction set (made of 13 reactions) including carbohydrates, proteins, and fats. Carbohydrates were modelled as cellulose, starch, and hemicelluloses, proteins as soluble proteins and insoluble proteins, while fats as tripalmate, triolein, palmitoolein, and palmitolinolein. Acidogenesis, acetogenesis, and methanogenesis are modelled using a different reaction set (made of 33 reactions) to calculate the kinetics of the reactions, whose constants were obtained following models such as ADM1 and comprehensive models. A FORTRAN code has been used to compute the reaction rates. Also, in this case, the simulation model uses NRTL as *Property Method*, and the reactors chosen for the hydrolysis and the other three phases are respectively a stoichiometric reactor (RSTOIC), with specified reaction extents, and a continuously stirred tank reactor (RCSTR), with specified kinetic constants. The model was validated against experimental and industrial data, using the biogas production rate as a validation parameter, for different substrates at different process conditions (7 case studies). The deviations from simulation results and experimental data span from 0.3 to 12.4% (absolute values).

Nguyen et al. [60] developed a simple one-step AD model to estimate the energy potential from the anaerobic digestion

of food waste in the municipal solid waste stream of urban areas in Vietnam. A stoichiometric reactor (RSTOIC) has been chosen to simulate the digester, in which the calculations are based on the Buswell equation, describing the overall process of anaerobic degradation. The global *Property Method* selected for the simulation is NRTL. The resulting biogas stream is separated in a flash separator, which separates the gas components and the digestate. The gaseous phase (raw biogas) is then treated to reduce the presence of H₂S and then introduced into a CHP unit, or into a boiler unit, or into an upgrading unit for biofuel production, depending on the model scenario chosen.

Scamardella et al. [61] simulated a pressurized anaerobic digestion process (PAD) using a RCSTR reactor operating at a pressure range of 1.5 to 5 bar. Reaction kinetics were taken from the ADM1 and comprehensive models. ELECNRTL (electrolyte non-random two liquid) *Property Method* is chosen here, as it allows to simulate dissociation equilibria that affect the CO₂ solubility in the liquid phase.

Peris Serrano [57] implemented the Angelidaki and the ADM1 models. The hydrolytic step is not taken into account in this simulation, and thus, only three phases are modelled. The process consists of two stages, i.e. two digesters, in which all the AD reactions occur. The reactor type selected is the RCSTR, for which total mixed flow and constant volumes are assumed, with residence time chosen as a user-defined parameter. The kinetic reactions in the model follow the power law and kinetic constants are computed in calculation blocks written in FORTRAN. The *Property Method* chosen for the simulation is NRTL.

Llanes et al. [62] developed an Aspen Plus model for the AD of vinasses, which integrates ADM1, flow pattern, and biofilm characteristics with the inclusion of sulphate reduction reactions. Vinasse is usually treated in UASB (upflow anaerobic sludge blanket) reactor types instead of completely mixed flow pattern reactors, so the authors employed two stoichiometric reactors (one for the hydrolysis stage and one for the methanogenic step) and two RCSTR reactors for the other phases. Kinetics are calculated in FORTRAN programmed blocks. Here as well, cellulose, hemicellulose, and dextrose were added as carbohydrates, proteins as soluble and insoluble, while lipids comprised of tripalmate, triolein, and palmito-olein. The *Property Method* adopted is the NRTL. The model has been validated against experimental data for three different case studies. A mean relative error lower than \pm 15% has been observed, with no significant differences between simulation results and experimental data in terms of biogas composition and methane yield.

Table 1 summarizes the main Aspen Plus models for AD reviewed in this work. The analysis has shown that the AD, though very complex to be fully described, can be simulated in Aspen Plus. Overall, the number of studies addressing AD



modelling is not very large: only six Aspen Plus simulation models have been found.

4.2 Modelling biogas upgrading

Several studies simulating biogas upgrading in Aspen Plus have been retrieved in the literature.

Ashraf et al. [8] developed a model for the PWS process: absorption and stripping columns are modelled as two RAD-FRAC distillation blocks, which is a column type designed for general vapour-liquid multistage separation. The thermodynamic method used is electrolyte non-random two-liquid (ELECNRTL): absorption of biogas components in water is accounted for by Henry coefficients, while the dissolution of H₂S and based on a first stage in which biogas is compressed to 12 bar, cooled to 313.15 K and then sent at the bottom of the absorption column, which is also fed from the top with water. The column is operated at 12 bar. Upgraded biogas leaves the column from the top of the column, and impurityrich sour water leaves from the column bottom. Sour water is then sent to a flash separator (operated at 3 bar) to remove residual CH₄, and then fed to the stripping column for regeneration at 1 bar, which uses air as a stripping medium. In the reported case study, biogas is fed at 2000 ppm of H₂S. For better removal of H₂S, activated carbon impregnated with ZnO is considered, and the resulting cleaned biogas has a composition characterized by less than 10 ppb of H₂S and NH₃, 99%, and 79% recovery of CH₄ and CO₂.

Cozma et al. used an Aspen Plus model to simulate a high-pressure water scrubbing (HPWS) system applied to biogas upgrading in [63] and [64] studies. The simulation model is characterized by operational conditions based on data taken from the literature (in particular, the work by Götz et al. [65]). The model is equilibrium-stage, and the thermodynamic method chosen for the analysis is a nonrandom two-liquid model with ideal gas and Henry's law (NRTL); the method has been chosen based on a preliminary study in which the authors compared the performance of different thermodynamic models available in the software to calculate the solubility of the main biogas components (CO₂, CH₄, H₂S, N₂, and O₂) in pure water. The simulation model assumes pressurization at 10 bar and cooling to 293.15 K of the biogas stream (60% vol CH₄, 38.9% vol CO₂, 300 ppm vol H_2S , 0.5% vol N_2 , and 0.5% vol O_2), which is then sent to the bottom of the absorber, which is also fed with water from the top. The scrubber is a RADFRAC column, working at T = 293.15 K, p = 10 bar. The number of stages and the absorbent flow rate required to achieve equilibrium have been determined through a preliminary study. The bottom stream (CO₂-enriched water) is transferred to a flash column, where the pressure is reduced from 10 to 3 bar to minimize methane loss. The gas containing CO₂, CH₄, H₂S, N₂, O₂, and water, released from the flash column is mixed with the

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|--|------------------------|-----------------|---|---|---|
| Authors | Publication year | Property method | Publication year Property method Reactor configuration in Aspen Plus | Reference model | Remarks |
| Al-Rubaye et al. [56] 2017 | 2017 | NRTL | RSTOIC (13 reactions) + RCSTR (33 reactions). Calculation blocks programmed in FORTRAN for kinetics computations | ADM1 and comprehensive models | Two-stage model (hydrolysis in RSTOIC and the other phases in RCSTR) |
| Llanes et al. [62] | 2019 | NRTL | two RSTOIC, two RCSTR, with calculation blocks programmed in FORTRAN for kinetics computations | The model integrates ADM1-flow pattern-biofilm characteristics with the inclusion of sulphate reduction reactions | Two-stage model (hydrolysis in RSTOIC and the other phases in RCSTR) |
| Nguyen et al. [60] | 2014 | NRTL | RSTOIC | Buswell equation | One-stage model |
| Rajendran et al. [59] | 2014 | NRTL | RSTOIC (13 reactions) + RCSTR (33 reactions), with calculation blocks programmed in FORTRAN for kinetics computations | ADM1 and comprehensive models | Two-stages model (hydrolysis in RSTOIC and the other phases in RCSTR) |
| Scamardella et al. [61] 2019 | 2019 | ELECNRTL | RCSTR | ADM1 and comprehensive models | PAD model (pressurized anaerobic digestion) |
| Serrano Peris [57] | 2011 | NRTL | two RCSTR in series with calculation blocks programmed in FORTRAN for kinetics computations | Angelidaki + ADM1 | The hydrolytic step is not taken in account |



raw biogas and re-circulated to the inlet of the compressor. After leaving the flash column, the rich solution is sent to the stripper, also modelled as a RADFRAC column, where it meets a counter flow of air. Here, CO₂ and H₂S are released from the water at atmospheric pressure and at a temperature of 293.15 K. Subsequently, the water is recirculated back to the top of the scrubber. Based on these conditions, it was calculated that the gas leaving the absorber contains 96.72% vol CH₄, 0.937% vol CO₂, 0.006 ppm vol H₂S, 1.1% vol N₂, and 0.976% vol O₂. The calculated energy demand for producing 309.36 Nm³/h of upgraded biogas is 171.5 kWh.

The work done by Götz et al. [65], which also described and modelled in Aspen Plus the HPWS technology for biogas upgrading, represented a reference also for the studies by Cozma et al. [63] [64]. Thus, the process conditions and scheme are nearly the same for both simulation models. The model is based on equilibrium and uses ELECNRTL thermodynamic model. The calculations take into account the gas quality requirements for biogas injection according to German law. Biogas feed has a composition of 53.7% vol CH₄, 45.2% vol CO₂, 101.8 ppmv H₂S, 0.93% vol N₂, 0.19% vol O₂, and 1.67% vol H₂O. The gas exiting the simulated process is composed by 96.8% vol CH₄, 0.47% vol CO₂, < 1 ppm vol H₂S, 2.1% vol N₂, 0.56% vol O₂, and 0.32% vol H₂O.

The PWS technology to convert biogas into biomethane has also been modelled in the thesis work by Menegon [66]. Similarly to Cozma et al. [63], the author carried out a preliminary study to select the most suitable *Property Method*, finally choosing the NRTL-RK model. The process conditions and scheme are very similar to those used by Cozma et al. [63]: biogas (45% vol CO₂, 55% vol CH₄) is compressed and cooled to 10 bar and 293.15 K and fed to an absorption column. The bottom stream is sent to a flash separator operating at 3 bar, from which gas is recirculated to the second compression stage, while liquid is sent to a stripper using air as a stripping medium. Regenerated water is sent back to the absorber. The simulation is rate-based, and the absorption and the stripping column reach a CH₄ purity of 98.7% vol. The CH₄ recovery is 99.08%.

The biogas water scrubbing technology has been also simulated in Aspen Plus by Bortoluzzi et al. [67]. The simulation scheme is similar to those cited above. The *Property Method* chosen is the Predictive Soave–Redlich–Kwong equation of state (SRK). In the simulated process, biogas is compressed to the absorption pressure of 10 bar through a two-stage intercooled compression, and water is removed via condensation. Then, biogas enters a packed column, which also receives a stream of liquid water; here, biogas upgrading occurs, thus a stream containing biomethane and a stream containing water, CO₂, H₂S, and small amounts of CH₄. This latter stream is flashed to 3 bar to recover methane; two streams exit the flash: one, containing vapour CO₂

and CH_4 , is recycled to the second compression stage, while the second one, liquid, is sent to a stripper. An air stream entering the stripper desorbs CO_2 (and H_2S) from the feed, and the solvent is then re-generated and recirculated to the absorber. The biomethane stream produced in the absorber is then dried: the CH_4 recovery of the process is 99.6%. For the base case, the molar percentage of CH_4 in the product stream is 98.7% mol.

The same modelling approach has been adopted also by Seman et al. [68]. Authors used the NRTL *Property Method*, adopting the same process conditions used in the study by Cozma et al. [64], as well as the simulation flowsheet. The specifications related to the absorption column are slightly different between the two studies, as the number of stages and the pressure is slightly higher in the work of Seman et al. [68]. This leads to a percentage of CO₂ removal and biomethane purity a little higher (97.6% mol CH₄ in the absorber product gas stream) in this latter case.

Ashraf et al. [8] also simulated the chemical absorption with MEA as absorption solvent for syngas upgrading (CO₂ removal after desulphurization). The processing scheme is similar to water scrubbing, i.e. an absorption and a stripping column, to remove CO₂ and to regenerate the solvent respectively, modelled as RADFRAC distillation columns.

Chemical absorption for biogas upgrading was simulated also by Lingelem [69], who used AMP solvent (2-amino-2-methyl-1-propanol), more specifically 30% wt AMP in aqueous solution. For the base case process, the author used the ELECNRTL thermodynamic model and RADFRAC columns (rate-based absorption, equilibrium-based desorption). The purified biogas stream is characterized by a CH₄ molar concentration of 97% mol. Six modifications of the base case have been also simulated.

Gamba et al. [70] simulated both water scrubbing and chemical absorption processes for biogas upgrading by means of a rate-based approach, according to modelling details from the by Pellegrini et al. [71]. The thermodynamic model used is electrolyte-NRTL. In the PWS simulation, biogas is treated in a one packed column at 20 bar with pure water at 298.15 K. There is no water regeneration step in the process. The inlet gas composition is 60% vol CH₄ and 40% vol CO₂. Other components have been neglected. Concerning the chemical absorption simulation, a 30% vol and 15% vol MEA aqueous solutions have been considered. The distillation column has the same characteristics as in the case of PWS, with the only differences in the packing material (metal instead of plastic) and the absorption pressure (atmospheric pressure). Both PWS and chemical absorption have been simulated find the absorbent flowrate needed for obtaining a 98% vol biomethane concentration on a dry

Gamba et al. [72] simulated water scrubbing, MEA (monoethanolamine) scrubbing, and MDEA scrubbing



when applied to obtain biomethane from municipal sewage sludge AD. Also in this case, the modelling approach is rate-based, and biogas components considered are only CH₄, CO₂, and water. For what concerns PWS, there is no water regeneration step, and the biogas is first sent to a threestage intercooled compression, then to the absorption column. Regarding the MEA chemical scrubbing, the model accounts for an absorption and a regeneration step in backed columns, operating at atmospheric pressure with a solution composed of 15% wt MEA. The MDEA chemical scrubbing case has the same process scheme of the MEA case. Here, absorption is carried out at 2.7 bar, using a 50% wt MDEA solution in water, and the regeneration at atmospheric pressure. All three simulated upgrading processes reach a biomethane purity higher than is 98% vol on a dry basis. For what matters in the PWS case, the upgraded biogas has a molar percentage of CH₄ of 98.86% mol on a dry basis, with a methane recovery to fed biogas of 95.8%. Regarding the MEA case, the upgraded biogas has a molar percentage of CH₄ of 98.71% mol on a dry basis, with a methane recovery to biogas of 99.9%. About the MDEA case, the upgraded biogas has a molar percentage of CH₄ of 98.73% mol on a dry basis, with a methane recovery to fed biogas of 99.98%.

The biogas upgrading technologies of PWS and chemical absorption with alkanolamine solutions have also been simulated in Aspen Plus by Pellegrini et al. [73]. In this study, three different biogas compositions have been tested, representing landfill gas, biogas from wastewater treatments, and gas from co-fermentation. The gases considered in these feed streams are CH₄, CO₂, N₂, and O₂. The layout of the water scrubbing process takes as a reference the one developed by Bortoluzzi et al. [67]. The biogas inlet stream is first compressed to 8 bar and cooled, then purified in a packed absorption column also fed with water. The bottom stream is then sent to a flash chamber and afterwards to a stripping column using air. Compression is here carried out in more steps. In each step separation of water from methane takes place too, using flash separators. The chemical scrubbing has been simulated referring to the process scheme of the study carried out by Gamba et al. [72]. The raw biogas is fed to the absorption column (2.7 bar), after being subjected to a single-stage compression and cooling down to 308.15 K. Both the absorption and stripping columns are packed columns (packing: metal Pall rings). The column specifications have been adjusted from the ones used in a previous work [70]. In all the case studies, the total flow rate of feed biogas is such that the volumetric flow rate of biomethane leaving the plant is 500 Sm³/h, to have a common basis for an economic feasibility comparison.

Similarly, Worawimut et al. [74] used Aspen Plus to simulate and compare the processes of water scrubbing and chemical scrubbing with diethanolamine (DEA) solution with regeneration and recirculation. The studies by Cozma

et al. [64] (PWS) and Niu et al. [75] (chemical absorption) have been taken as references to set process conditions and for results validation. The NRTL property method has been selected and RADFRAC distillation columns without condenser and reboiler have been used to model the absorber and the desorber, both set as equilibrium based. Biogas from swine farm wastes was used in this work, with a composition of 68% vol CH₄, 24% vol CO₂, 3000 ppm vol H₂S, 2% vol N₂, 0.1% vol O₂ and 5.6% H₂O. The biogas flow rate of the plant is 1000 kmol/h. Both water scrubbing and chemical scrubbing were simulated to find the absorbent flow rate needed to obtain at least 96% v/v biomethane purity. Total water flow rate of the plant is 16,000 kmol/h, which is the same amount of the total DEA solution flow rate of the plant. The product gas of the PWS process is characterized by a composition of 96.005% vol CH₄, 0.32% vol CO_2 , < < 0.001% vol H_2S , 3.098% vol N_2 , 0.154% vol O_2 , 0.424% vol H₂O. The methane recovery is 89.96%, while the energy consumption of the process is 11309 kW. The product stream of the chemical absorption with DEA has a composition of: 96.026% vol CH_4 , 0.310% vol CO_2 , < 0.001% vol H_2S , 3.114% vol N_2 , 0.155% vol O_2 , 0.395% vol H_2O . The process has a methane recovery equal to 89.47%, with an energy consumption of 11,331 kW.

Gangadharan et al. [16] also simulated the technology of chemical scrubbing with DEA in Aspen Plus for acid gas removal from natural gas. The simulation is rate-based, the thermodynamic property method is ELECNTRL, and also here the main blocks of the flowsheet are the absorber and stripper (RADFRAC distillation columns), but no solvent recirculation is included. The cleaned gas then enters 20-stages distillation column, operating at 44.6 bar, where methane gets separated from C-2 to C-3 components. A 99.5% mole recovery of methane is obtained from the process.

Membrane separation technique has been simulated by Scamardella et al. [61]. The authors adopted a user-defined model (user 2 block), interfacing the block with an Excel file. The *Property Method* selected is ELECNRTL. The model refers to Fick's law with diffusive model assumptions. In the model, the output of the membrane are two streams, i.e. a CH₄-rich gaseous stream, a CO₂-rich gaseous stream, and the off-gas of the process. The resulting biomethane can reach high purity percentages (> 95% vol) for operating pressures higher than 3 bar.

As Aspen Plus is a steady state calculator, no dynamic options are available in the software. Therefore, since PSA is a dynamic process, examples of PSA process simulations have been found mainly on Aspen Adsorption, e.g. Menegon et al. [66], Abdeljaoued et al. [76]. Anyway, some studies simulating the PSA process in Aspen Plus have been found and are reported below.



Gamero et al. [34] simulated the PSA process to clean the outlet gas of gasification to obtain high quality syngas and simultaneously capture the greenhouse gases. The system consists of four units, composed by ideal column separators operating at pressure and temperature conditions (30.6 bar and 308.15 K), to separate H_2 (first PSA unit), CO (second PSA unit), CO_2 (third PSA unit), and CH_4 (fourth PSA unit). The components obtained are then mixed, to obtain the H_2/CO ratio required for the downstream utilization. Multistage compressors and valves are also used in the process. The Peng-Robinson with Boston Mathias function method was selected for the simulation. As a result, about 80% of the CO_2 and 95% of CH_4 fed in the PSA system were sequestrated.

Similarly, Ortiz et al. [77] simulated the PSA process to clean syngas obtained by supercritical water reforming of glycerol. The PSA system is composed by three units: in the first unit, highly pure H₂ is separated as a non-adsorbed stream; in the second one, CO is separated as the adsorbed component; in the third one, separation of CO₂ and CH₄ occurs. The obtained gas streams are then mixed to obtain the H₂/CO ratio required for downstream utilization. Distillation columns for purification and valves for depressurization are involved in the process. NRTL property method has been selected. The system reaches 95% H₂ recovery, 98% CO recovery, and 90% CO₂ as well as CH₄ recovery.

Table 2 summarizes the Aspen Plus simulation models reviewed in this work for biogas upgrading. The analysis revealed that upgrading processes have been largely simulated in literature: more than 15 studies have been found, most of which addressing the processes of physical and chemical absorption.

4.3 Modelling methane reforming

Gangadharan et al. [16] simulated dry reforming and steam reforming for syngas production from natural gas. The simulation scheme consists of a first step dedicated to acid gas removal (H₂S and CO₂) from natural gas through chemical absorption with DEA, followed by methane separation from higher hydrocarbons in a distillation column, steam production in heat exchanger and finally SMR. Methane exiting the acid gas removal step is mixed with steam in a mixer, which uses heat from the SMR reactor output stream. The mixture is then sent to another heater and then to the plug flow reactor (RPLUG), where the SMR and WGS reactions take place over a Ni/Al₂O₃ catalyst at a constant temperature of 890 K. The output stream is then sent to a heat exchanger, used to generate steam, and then to a flash separator, where syngas and water are obtained. The thermodynamic property method selected is Peng-Robinson, and the convergence criteria have been relaxed due to issues in the PFR convergence with the default criteria. The rate expression for the catalytic reactions occurring in the PFR reactor have been modelled using the Langmuir-Hinshelwood-Hougen-Watson kinetics formulation (LHHW), obtained from the work of Xu and Froment [78]. The produced syngas is characterized by a composition of 72.22% mol H₂, 21.71% mol CO, 3.56% mol H₂O, 1.77% mol CO₂, 0.48% mol CH₄, and 0.26% mol N₂. The authors also simulated a combination of SMR and DMR. In this process, the syngas generated by SMR is sent to a heat exchanger. Here, the syngas is cooled and passed through a CO₂ membrane separator, separating CO_2 from the syngas mixture, which contains CO_2 , H_2 , CO_2 , H₂O, and unreacted CH₄. Then, the stream is sent to a flash separator, where the separation of syngas and water takes place. The CO₂ separated by membrane filtering is sent to the dry reformer, where the methane reacts with CO₂ for increased production of syngas. As for the SMR process, LHHW kinetic expressions are used to determine the rate of reaction of the DMR process. The resulting syngas has a composition of 73.61% mol H₂, 23.85% mol CO, 1.20% mol H_2O , 0.51% mol CO_2 , 0.55% mol CH_4 , and 0.27% mol N_2 .

Giwa et al. [79] simulated the SMR for hydrogen production. The authors modelled two different versions of the process, i.e. with and without feed ($\rm CH_4$ and $\rm H_2O$) mixer. For reforming, an equilibrium reactor was chosen, in which the stoichiometry of the reaction was specified, i.e. the reforming reaction (1) and the water–gas-shift reaction (2). Several case studies have been simulated. In the case of the reactor operating at 1173.15 K temperature and 1 bar pressure, the syngas compositions obtained in both versions of the model were characterized by the same molar composition, i.e. 62.56% mol $\rm H_2$, 16.06% mol $\rm CO$, 17.77% mol $\rm H_2O$, 3.59% mol $\rm CO_2$, and 0.02% mol $\rm CH_4$.

The SMR process of natural gas has been also modelled by Amran et al. [80], using a kinetic-based approach with the Redlich-Kwong-Soave equation of state with modified Huron-Vidal mixing rules (RKSMHV2) thermodynamic model. Natural gas and steam are first mixed, fed to a heat exchanger and then to a RPLUG reactor to model the methane reforming reaction, and finally to another RPLUG reactor for WGS reaction. Both reactors follow a rearranged LHHW kinetic model. It is assumed that natural gas does not contain H₂S and CO₂. The modelling approach was validated against data from other published studies, showing a good agreement with the literature. A sensitivity analysis of the reaction performance has also been performed.

Gopaul et al. [81] simulated the syngas production from biogas through dry reforming. In particular, three different cases have been simulated, i.e. (i) DMR alone, (ii) DMR and POX, (iii) DMR and hydrogen oxidation (HOX). The target H₂/CO ratio is 1.6–1.7, for downstream Fischer–Tropsch synthesis. The study also compares different types of biogas in terms of H₂ and CO yield: landfill, corn cob, whole stillage, and combined cob and stillage. The compositions of



 Table 2
 Reviewed Aspen Plus models for upgrading processes

| Authors | Publication year Technology | Technology | Property method | Property method Main simulation blocks | Process yield | Remarks |
|-------------------------|-----------------------------|---------------------------------|--------------------------------|---|---|--|
| Ashraf et al. [8] | 2015 | PWS | ELECNRTL | RADFRAC distillation columns for absorption and stripping | CH₄ recovery of about 99% | Biogas upgrading |
| | | Chemical absorption with MEA | ELECNRTL | RADFRAC distillation columns for absorption and stripping | Not specified | Syngas upgrading |
| Bortoluzzi et al. [67] | 2014 | PWS | SRK | Distillation columns for absorption and stripping | 98.97% mol, with a CH ₄ recovery of $99.6%$ | Natural gas cleaning |
| Cozma et al. [63] [64] | 2013, 2015 | PWS | NRTL | RADFRAC distillation columns for absorption and stripping (equilibrium-based) | About 96.72% vol CH_4 in the product stream | Biogas upgrading |
| Gamba et al. [70] [72] | 2013, 2015 | PWS | ELECNRTL | RADFRAC distillation columns for absorption and stripping (both rate-based) | About 98.86% mol CH ₄ on a dry basis in the product stream, CH ₄ recovery of about 95.8% | Biogas upgrading |
| | | Chemical absorption with MEA | | | About 98.71% mol CH ₄ on a dry basis in the product stream, CH ₄ recovery of about 99.9% | |
| | | Chemical absorption with MDEA | | | About 98.73% mol CH ₄ on a dry basis in the product stream, CH ₄ recovery of about 99.98% | |
| Gamero et al. [34] | 2018 | PSA | PR-BM | Ideal column separators, multi-stage compressors and valves | about $80\% \text{ CO}_2$ and $95\% \text{ CH}_4$ from the feed stream captured | Cleaning the outlet gas of gasification to obtain syngas |
| Gangadharan et al. [16] |] 2012 | Chemical absorption with DEA | ELECN- RTL+RK for vapour | 2 RADFRAC distillation columns for absorption and stripping | CH ₄ mole recovery of about 99.5% | Natural gas cleaning |
| Götz et al. [65] | 2011 | PWS | ELECNRTL | Distillation columns for absorption and stripping | About 96.8% vol CH_4 in the product stream | Biogas upgrading |
| Lingelem [69] | 2016 | Chemical absorption with AMP | ELECNRTL | RADFRAC distillation columns for absorption and stripping (rate-based absorp- tion, equilibrium-based desorption) | 97% mol on the base case | Biogas upgrading |
| Menegon [66] | 2017 | PWS | NRTL-RK | Distillation columns for absorption and stripping (equilibrium-based first, then rate-based) | 98.7% vol, with a CH ₄ recovery of 99.81% | Biogas upgrading |



| Table 2 (continued) | | | | | | |
|------------------------------|-----------------------------|------------------------------|-----------------|--|--|---|
| Authors | Publication year Technology | Technology | Property method | Property method Main simulation blocks | Process yield | Remarks |
| Ortiz et al. [77] | 2012 | PSA | NRTL | Ideal column separators, multi-stage compressors and valves | 95% H ₂ recovery, 98% CO recovery, 90% CO ₂ recovery, 90% CH ₄ recovery | Cleaning syngas obtained by supercritical water reforming of glycerol |
| Pellegrini et al. [73] | 2015 | PWS | No information | Distillation columns for | Not specified | Biogas upgrading |
| | | Chemical absorption with MEA | No information | absorption and stripping | Not specified | |
| Scamardella et al. [61] 2019 | 2019 | Membrane separation | ELECNRTL | User-defined model | >95% vol | Biogas upgrading |
| Seman et al. [68] | 2019 | PWS | NRTL | Distillation columns for absorption and stripping | 97.6% mol | Biogas upgrading |
| Worawimut et al. [74] | 2018 | PWS | NRTL | RADFRAC distillation columns for absorption and | 96.005% vol, with a CH ₄ recovery of $89.96%$ | Biogas upgrading |
| | | Chemical absorption with DEA | | stripping (equilibrium-based) 96.026% vol, with a CH ₄ recovery of 89.47% | 96.026% vol, with a CH ₄ recovery of $89.47%$ | |
| | | | | | | |

the analysed biogas types do not include H₂S or NH₃. Biogas is not upgraded to biomethane prior to reforming, thus the reforming reactor is directly fed by the biogas stream. About the thermodynamic model, the Property Method chosen here is IDEAL, which uses both Raoult's law and Henry's law. The DMR alone case was simulated using a RGIBBS reactor, i.e. an equilibrium reactor whose output is computed following the method of Gibbs free energy minimization at specified operating conditions (pressure, temperature, flowrates). Biogas is fed into the reforming reactor at 1123.15 K and 1.01325 bar. The reactor operates at 1223.15 K and 1.01325 bar. The DMR + POX case is similar to the previous one in terms of reactor configuration and operating conditions. However, in this case, a second feed stream containing oxygen at 1.01325 bar and 473.15 K is used, and therefore the exothermic partial oxidation reaction satisfies the energy demand of the endothermic DMR process. The amount of oxygen required was determined using the Design Specification function available in Aspen Plus, taking also into account the desired syngas H₂/CO ratio of 1.6–1.7. The case of DMR+HOX modelling comprises two RGIBBS reactors, one for DMR and one for H₂ combustion to provide energy to the DMR process. Biogas feed and DMR reactor have the same conditions of the other two cases. The HOX reactor is fed under stoichiometric excess conditions of O₂ at 473.15 K and 1.01325 bar, H₂ at 1123.15 K and 1.01325 bar, and combustion occurs at 1273.15 K and 1.01325 bar. Also in this case, the required H2 and O2 feed rates were determined through the Design Specification function in Aspen Plus. The optimal process conditions to maximize syngas yield and quality were determined through a sensitivity analysis on the DMR case with landfill biogas type, which turned out to be similar also for the other biogas types. The analysis showed that, however, the desired 1.6–1.7 H₂/CO ratio is found at temperatures and pressure ranges for which syngas quality is low. Therefore, other values of H₂/CO ratio, slightly outside the desired range have been accepted in favour of a better CH₄ conversion (from 96 to 100%) and syngas quality (meaning, for high-quality syngas, a syngas composed mainly of H₂ and CO, with a minimal amount of by-products). Authors also performed an energy analysis of the processes, which are both exothermic and endothermic. DMR reactor heat duty ranges from $+ 14.88 \text{ to} + 28.74 \text{ kW}_{th}$ kmol, while the DRM + HOX process values of -23.93and – 4.58 kW_{th}/kmol depending on biogas type (resulting in exothermicity); the combined DMR + POX process was able to counterbalance the high energy demand of DMR, achieving thermal-energy neutrality.

The DRM process has been modelled in Aspen Plus also by Ashraf et al. [8]. In this case, the RGIBBS reactor block was used, fed by cleaned biogas, steam, and air (to lower coke formation and external energy demand). A sensitivity study to find the optimum process conditions has also been



carried out. As a result, a syngas stream characterized by a H₂/CO ratio of about 1.58 was obtained, with a methane conversion of about 99%.

Er-rbib et al. [82] simulated a combination of DMR and SMR processes to produce syngas from natural gas. The reforming unit is composed by two parts: a pre-reformer and a reformer. In the pre-reformer, a complete conversion of the higher hydrocarbons of natural gas into methane occurs over a nickel catalyst at 823 K and 5 bar. Then, in the reformer, the primary SMR reaction (1), the primary DMR reaction (4), and the RWGS reaction (11) take place. The chosen reactors are equilibrium reactors, and thus no kinetic models are considered, whereas the thermodynamic model used is the Peng Robinson with Boston-Mathias alpha function (PR-BM).

Table 3 summarizes the main Aspen Plus models on upgrading processes reviewed in this work. A large number of studies were identified, most of them dealing with SMR. The analysis however revealed that there is also a good number of papers addressing the DMR, also in combination with SMR, as well as POX, while a smaller number of articles addressing ATR was found.

4.4 Modelling Fischer-Tropsch synthesis

Modelling the Fischer–Tropsch process is particularly challenging due to the high number of species existing in equilibrium, the variety of reaction products, the complexity of the CO catalyst chemistry, and the large number of process parameters relevant to the process [12]. Indeed, the identification of a plausible mechanism, as well as the formulation of a representative expression addressing the consumption rate of the primary component CO and an accurate description of the product distribution are crucial and complicated steps in modelling the Fischer–Tropsch synthesis [84] [85]. A comprehensive review of the FT kinetics has been carried out by Van Der Laan and Beenackers [86]. The Anderson-Schulz-Flory (ASF) model is normally used to represent the FT product distribution, based on one parameter, namely the chain probability factor α , which describes the addition of carbon atoms into the molecule chain [87]. However, in most cases, the real Fischer-Tropsch product selectivity does not obey the ideal ASF distribution [88], and deviations (essentially higher selectivity to CH₄, and lower to C₂H₄ than expected in the model) are well documented in the literature [84]. Dependence of the chain probability factor α on process conditions (pressure, temperature, composition, catalyst type, etc.) has been largely studied and correlations have been formulated, e.g. [89–91].

In this section, different models found in literature simulating the Fischer-Tropsch synthesis in Aspen Plus are analysed.

Ashraf et al. [8] simulated the FT synthesis in a slurry reactor from bio-syngas. A RYIELD reactor block with CO conversion of 80% has been selected. This block does not require exact information about the stoichiometry or kinetics, but models a reactor by specifying the reaction yields of each component [58]. The product distribution follows the ASF distribution, with α values computed according to Kruit et al. [91] parameters. After the synthesis, the FT syncrude is then sent to a distillation column (RADFRAC) to separate products according to the following classification: C-1 to C-4 lights, C-5 to C-9 naphtha, C-10 to C-16 kerosene, C-17 to C-22 diesel, and C-22 + waxes. As the selectivity of FTcrude products depends on the reaction temperature and feed syngas (H₂/CO ratio), a sensitivity study has been carried out, and these parameters optimized to maximize kerosene and diesel fraction using the solver function in MS Excel. In order to achieve 80% CO conversion and maximize kerosene and diesel yield fractions, the optimal values have been estimated between 1.6 and 2 for the H₂/CO ratio, and between 473.15 and 573.15 K as regards the reaction temperature. Considering the whole process (biogas to liquid fuel conversion process using pressurized water scrubbing, dry methane reforming, and FT-synthesis), the overall carbon conversion efficiency reaches 45%, while the energy efficiency is 30%.

Adelung et al. [30] simulated the production of synthetic hydrocarbons (in particular kerosene and diesel) from syngas derived from captured CO2 and H2 obtained through water electrolysis. In the proposed approach, syngas is generated through a reverse water-gas-shift (RWGS) reaction and then converted through the FT reaction into a broad range of hydrocarbons. Product separation is performed downstream hydrocarbon production: long-chain hydrocarbons are sent to a hydrocracker to increase the yield at the desired chain length for transport fuel production (< C-22). Gas products and unreacted species are recycled to increase the carbon efficiency of the process. Operational parameters are optimized to maximize energy efficiency. In Aspen Plus, the thermodynamic method here selected is the Peng-Robinson with Boston-Mathias modifications (PR-BM), H₂, CO, CO₂, H₂O, N₂, O₂ and alkanes are the chemical species considered in this model. It is assumed that carbon is not a possible product. The FT reactor is a tubular fixed bed: it has been simulated as a stoichiometric reactor (RSTOIC) with Co catalyst, without considering reaction kinetics. The reactor operates at 493.15 K and 25 bar, while the H₂/CO ratio is set at 2. CO conversion is assumed equal to 40%, and inert gas share is fixed at 50%. Under these conditions, the chain growth probability factor α is equal to 0.839, calculated using the expression proposed by Vervloet et al. [90]. The stream from the reactor is first sent to a flash separator, where heavy hydrocarbons are separated, and then to a hydrocracker. The product from the hydrocracker is then subject to further separation through 8 different flashes into



Syngas production from Hydrogen production Hydrogen production from natural gas Study objectives from methane natural gas natural gas natural gas biogas biogas $96\% \div 100\%$ depending $0.96 \div 2.80$ depending $0.96 \div 2.80$ depending 0.96 ÷ 2.80 depending Resulting syngas H₂/CO Not reported for speon the feed type on the feed type on the feed type cific conditions Not specified Not specified 1.58 3.09 3.90 3.33 96% ÷100% depending 96% ÷ 100% depending Methane conversion reactor length of Can reach 30% for on the feed type on the feed type on the feed type Not specified about 10 m Not specified Not specified Not specified Not specified About 99% Two RGIBBS reactors Two REQUIL reactor RPLUG reactor with RPLUG reactor with RPLUG reactor with LHHW kinetics LHHW kinetics LHHW kinetics REQUIL reactor Main simulation blocks RGIBBS reactor RGIBBS reactor RGIBBS reactor RGIBBS reactor Not specified for this part of the process Thermodynamic Not specified PENG-ROB PENG-ROB **RESMHV2** PR-BM IDEAL IDEAL model combined SMR and
 Table 3
 Reviewed Aspen Plus simulation models for reforming
 SMR (Ni/Al₂O₃ DMR process DMR + SMR DMR+HOX DMR + POX Publication year Technology catalyst) DMR SMR DMR SMR ATR 2015 2012 2012 2013 2015 2017 2008 Gangadharan et al. Gopaul et al. [81] Amran et al. [80] Er-rbib et al. [82] Ashraf et al. [8] Giwa et al. [79] Hao et al. [83] Authors



hydrocarbons, water, and recycling gases. The carbon efficiency of the overall process is 88%, thanks to the recycles, while the power-to-liquid efficiency for the base case is 38.7%.

Campanario et al. [77] simulated the production of lowtemperature Fischer-Tropsch products from syngas obtained by supercritical water reforming of the bio-oil aqueous phase. The overall process is composed of four different sections, i.e. (i) supercritical water reforming (SCWR) of the bio-oil aqueous fraction, (ii) upgrading of the syngas to increase H₂ and CO molar flow rate and to achieve the desired H₂/CO ratio through water-gas-shift and dry reforming reactors and PSA systems, (iii) Fischer-Tropsch synthesis loop, and (iv) refining and upgrading of FT products by means of distillation columns and hydrocracking. Focusing on the FT synthesis section, the selected reactor is a stoichiometric one (RSTOIC) operating in a temperature range of 493.15 to 513.15 K and 20 to 40 bar, fed by syngas at $2 H_2$ / CO ratio. FT products were assumed to be composed only of olefins and paraffins, and the probability parameter of chain growth propagation, α , has been computed using the expression obtained by Song et al. [89]. The stream leaving the FT reactor is then cooled to condense heavier hydrocarbons and separate them from the gas. The gas stream is recycled back to the FT reactor to maximize the overall CO conversion and to increase the production of liquid fuel, while the liquid phase is sent to a decanter for separating H₂O from heavier hydrocarbons, which are first expanded through a valve and finally sent to the distillation section. The thermodynamic methods used are UNIQUAC for the distillation train and Peng-Robinson EOS for the FT section. The effect of the main operating parameters on the process performance, such as feed composition and operating conditions of the Fischer-Tropsch reactor, was studied by a sensitivity analysis. Optimal conditions were identified; thus, for a mass flow of aqueous phase of 60 t/h with a total organic concentration of 35% wt, biofuel production was estimated equal to 4596 kg/h (2804 kg/h FT-diesel, 1491 kg/h FT-jet fuel, and 301 kg/h FT-gasoline), the carbon efficiency with refining to 38.53% (without refining, it was estimated at 43.50%), while net electrical power was 5297 kWe.

Niassar et al. [92] simulated the FT synthesis in the context of development and optimization of an Integrated Process Configuration for IGCC Power Generation Technology, with Fischer–Tropsch fuels from coal and biomass. Basically, the process consists in a first part related to syngas generation from gasification, which is then split and sent into the FT unit, where it is converted to fuel, and the combined power cycle generates electricity and power. Focusing on the Aspen Plus model section related to the Fischer–Tropsch unit, the reactor chosen by the authors is a stoichiometric reactor (RSTOIC) in which 31 reactions have been considered. The property method selected is Peng-Robinson. The

feed syngas is characterized by H₂/CO ratio of about 2, and the process is carried out at about 513.15 K and 20 bar. The syngas is thus mainly converted to C-1 to C-30 hydrocarbons and water. Chemical reactions have been defined up to C-30 as the database of software does not contain hydrocarbons that are heavier than the C-30. The products of the Fischer–Tropsch reactor undergo downstream separation under gradual cooling in the three separators. Lightweight and heavyweight hydrocarbon liquids are the main products of the process, while gases are sent to the power plant for power generation. A sensitivity analysis has been carried out, and the simulation results compared with an experimental work [93, 94], indicating that the difference in results is about 4%.

In the context of assessing biogas-to-liquid processes for bagasse utilization, Michailos et al. [95] developed an Aspen Plus model to simulate the Fischer-Tropsch synthesis of bio-syngas. The studied production route included a gasifier unit, syngas quenching and cleaning, a FT synthesis reactor, product recovery and separation, and finally a heat and power generation system. The biomass, i.e. sugar cane bagasse, undergoes a pre-treatment constituted by bagasse crushing to small particles and drying before entering the system. With regards to Fischer-Tropsch modelling, authors used a product distribution reactor (RYIELD), following the Anderson-Schultz-Flory distribution model ($\alpha = 0.9$), through which the mass yield of the products of the synthesis were determined. The reactor module interfaces with an Excel Spreadsheet where these calculations are carried out. The feed syngas is characterized by a H₂/CO ratio of 2.05. The product stream exiting the FT unit is then sent to a flash to separate the hydrocarbons from the unconverted syngas, which is recycled back. The hydrocarbon stream is then sent to a purification zone, consisting of four distillation columns and a hydrocracking unit for waxes, with a conversion efficiency of 88%. The Property Method chosen for conventional components is the Redlich-Kwong-Soave cubic equation of state with Boston-Mathias alpha function (RKS-BM). For a feed consisting of 100 t/h of sugar cane bagasse (before being subjected to the pre-treatment process), the product flow rates of the system is 9100 kg/h diesel, 6050 kg/h gasoline, and 1175 kg/h LPG. The energy efficiency of the process is about 68%.

Hao et al. [83] used Aspen Plus to simulate a gas-to-liquid (GTL) process involving syngas generation through ATR and Fischer-Tropsh synthesis. The FTS has been simulated based on detailed kinetic models considering two kinds of industrial catalysts, i.e. iron and cobalt. The authors tested two different Aspen Plus reactor blocks, i.e. PRF (plug flow reactor) and CSTR (continuous stirred tank reactor). The detailed kinetic models for the two different catalyst types have been programmed in FORTRAN and compiled as user-defined functions for the simulation software. The authors



performed a sensitivity analysis for both the catalysts to understand the performances of the two models (PFR and CSTR), by varying operating conditions and H₂/CO ratio of the syngas. Different recycling options for the FT tail to the ATR have been simulated to find the optimal flowsheet structure, which was selected according to the overall thermal efficiency to crude products, the overall carbon efficiency to crude products, and the energy value of the purge gas. The study concluded that the thermal efficiency to crude products for the cobalt-based catalyst is about 60%, while for the iron-based catalyst, it is in the range of 49–55%. Additionally, FT synthesis with Fe-based catalyst generates CO₂; its carbon efficiency (61–68%) turns out to be lower with regards to the cobalt-based catalyst (73–75%).

Er-rbib et al. [82] developed an Aspen Plus model to describe the production of synthetic gasoline and diesel fuels. The process consists of four different stages: (i) production of syngas from the combination of dry reforming and steam reforming of natural gas, (ii) Fischer-Tropsch synthesis to produce long chains of hydrocarbons, (iii) separation of fuel and wax hydrocracking, and (iv) recovery of hydrogen. The FT synthesis was modelled using a stoichiometric reactor (RSTOIC), specifying 42 reactions for which information about selectivity and efficiency has been found in the literature. The operating conditions are 513 K and 20 bar. In these conditions, the conversion of synthesis gas was estimated at 87%. The reactor products are cooled and separated from water and oxygen compounds, and then sent to a distillation column for the separation of heavy and light components. Waxes are finally converted into high-quality diesel through a hydrocracking unit, which has been simulated as a RYIELD reactor. The Property Method used for the reactors and the distillation columns is Peng Robinson with Boston-Mathias alpha function (PR-BM). The results show that the overall process can produce synthetic fuels composed by 72% of diesel, 26% of gasoline, and 2% of LPG.

Sudiro et al. [96] used Aspen Plus to simulate synthetic fuel production through LTFT synthesis of syngas obtained from coal and natural gas. Three processes have been simulated, i.e. (i) gas to liquid (GTL), (ii) coal to liquid (CTL), and (iii) a hybrid process coupling features of both CTL and GTL. Focusing on the Fischer–Tropsch section of the model, a RYIELD reactor block has been chosen, with syngas conversion assumed equal to 87% at 513.15 K and 15 bar operating conditions. Selectivity values (ratio between moles produced and moles of syngas converted) have been specified, taking data from the literature. Forty-four reactions of type (6) and (7), respectively olefin formation and alcohol formation, have been used for all components from CH₄ to C₆₀H₁₂₂ and ethanol. Product distribution on a weight basis is gasoline (C-5 to C-11) 25.6%; diesel (C-12 to C-18) 40.3%; waxes (C-19 to C-60) 31.6%; light gases 1.6%; and oxygenated compounds 1%. The products are then subject to hydrocracking, separation, water treatment, and recycling. The *Property Method* used for the process parts involving reactors, distillation columns, and two-phase separators is the Peng-Robinson equation of state with Boston-Mathias alpha function, while for separations involving three phases the NRTL equation was applied. Simulated product yields for three cases are 66.7% for GTL, 32.5% for CTL, and 44.4% for the hybrid process., on a weight basis. The estimated thermal efficiency, i.e. ratio between the energy contents in the products and in the feedstock, is 54.2%.

Bao et al. [97] simulated the FT synthesis in the context of process design optimization of a GTL plant. In the study, the authors assumed a feed $\rm H_2/CO$ ratio of ~2 and that the process follows the ASF product distribution, with a fixed chain growth probability factor α equal to 0.95. The syncrude is fed into a distillation column to separate LPG, naphta, and wax. NRTL-RK is the property method used, while the reactor configuration is not specified. The simulated plant converts 900,000 kg/h of natural gas into 118,000 BDP of products.

Cinti et al. [98] used Aspen Plus to model the FT synthesis as a part of a study addressing the production of synthetic green fuels through a system integrating a solid oxide electrolyzer and the Fischer-Tropsch process. The plant is divided into two main sections, i.e. (i) the electrolyzer unit and (ii) the liquid fuel synthesis unit. In (i), H₂O and CO₂ are converted into H₂ and CO (syngas) via co-electrolysis, whereas in (ii), syngas conversion into hydrocarbons occurs. In the process modelled, the FT synthesis occurs at 20 bar and 503.15 K, and only a first FT crude separation is considered, which divides purge water from hydrocarbons and light refinery gases. The main Aspen Plus simulation blocks considered for the FT modelling part are a splitter, a stoichiometric reactor, a mixer, and a flash separator. The splitter divides the feed syngas ($H_2/CO = 2.1$) into two streams: one enters the RSTOIC block, while the other bypasses the reactor and is mixed with the FT products. The FT reactor (Co-based) accomplishes several reactors, and the product distribution is assumed to follow the ASF products distribution model with a chain growth probability factor α equal to 0.94. The synthesis of alcohols, aromatics, and other oxygenated compounds is neglected; only alkanes and alkenes are considered products. The syncrude is then sent to a flash separator performing the separation of light gases, liquid phase, and water. For a given syngas feed characterized by a molar composition equal to 51.2% mol H₂, 24.4% mol CO, 24.3% mol CO₂, and 0.1% mol CH₄, the products at FT reactor outlet (without any separation) is composed by 8.16% mol H₂, 40.78% mol H₂O, 3.58% mol CO, 43.94% mol CO₂, 0.4% mol CH₄, 0.24 mol C₃H₆, 0.38% mol C₃H₈, 0.06% $mol C_6H_{12}, 0.32\% mol C_6H_{14}, 0.06\% mol C_8H_{16}, 0.57\% mol$ C_8H_{18} , 0.01% mol $C_{16}H_{32}$, 0.94% mol $C_{16}H_{34}$, and 0.53%



mol C_{30+} . The total energy efficiency of the FT system is 52.57%; while considering the FT products deprived of light gases as a valuable product, the energy efficiency is 40.95%.

Also Pondini et al. [99] developed an Aspen Plus model considering low-temperature Fischer-Tropsch to simulate synthetic fuels production from biomass-derived syngas. The FT reactor (Co-based catalyst) has been modelled using a RSTOIC reactor, for which the fractional conversion of each reaction is imposed as calculated in an integrated Excel file according to an estimated chain-length distribution. Hydrocarbons (olefins and paraffins) with carbon numbers up to 30 are considered, and the Song et al. [89] correlation for the chain growth probability factor α is considered. The mole fraction calculations for products with carbon numbers C-1 to C-4 have been adjusted to take into account the ASF deviation (i.e. higher methane selectivity) with reference to Rane et al. [100]. Different operating conditions (H₂/CO ratio, reactor pressure, temperature, CO conversion) have been tested.

Marchese et al. [101] modelled the Fischer-Tropsch synthesis in the context of analysing different power-to-liquid options, in which the FT section is integrated into a complete carbon capture and utilization from a biogas upgrading unit producing about 1 ton/h of CO₂. The recovered CO₂ is turned into syngas through either a reverse water gas shift reactor or to a solid oxide electrolysis unit operating in co-electrolysis mode; the produced syngas is fed to a Fischer-Tropsch reactor operating at 25 bar and 501 K, whose products are then separated into light gas, naphtha, middle distillates, light waxes, and heavy waxes. The process model implements a detailed kinetic model developed in the author's previous study [102] based on real experimental data, which accounts for deviations from the Anderson-Schulz-Flory (ASF) distribution (i.e. higher methane and lower ethylene formation). For the implementation in Aspen Plus, a plug flow reactor (RPLUG) was selected, integrated with an external kinetic subroutine for rates definition up to C-80 for paraffins and C-40 for olefins. The Property Method chosen for the FT unit was the RKS-BM. The simulation results show that, for the case of the solid oxide electrolyzer to produce syngas, the best model configurations can reach a plant efficiency of 81.1%, while for the reverse water gas shift option, the plant efficiency reaches 71.8%.

The same modelling approach was used also in another study [103] addressing the energy and economic analysis of plant configuration integrating the direct air capture technology for CO_2 recovery and the Fischer–Tropsch synthesis. In this case, the carbon number of alkanes and alkenes considered spans from C-1 to C-70.

In another study [104], aimed at analysing the technoeconomic feasibility of a biomass-to-X plant, Marchese et al. modelled the FT synthesis using a different approach. The synthesis of paraffin was described up to C-40 and for olefins up to C-19, using the ASF distribution with α dependent over the temperature and syngas composition according to Song et al.'s [89] correlation. CH₄ yield was assumed equal to 20%mol, in order to account for the ASF deviation for this compound. Moreover, a 90% internal recirculation for unconverted syngas was considered. The reactor configuration in Aspen Plus is not specified in the paper, while the thermodynamic method used for the FT section of the overall process is Redlich-Kwong-Soave with Boston-Mathias modification (RKS-BM) EoS.

Gabriel et al. [105] modelled the FT synthesis in the context of a GTL process composed of three sections, i.e. synthesis gas production from natural gas and conditioning, FT reaction, and FT product upgrading and separation. Different plant configurations have been evaluated; changes are limited to the syngas production technologies and conditioning sections. Focusing on the part of the process model in Aspen Plus assessing the Fischer–Tropsch synthesis, the authors used a RSTOIC reactor with a per pass conversion of 70%. The products follow the ASF distribution, with a constant $\alpha = 0.92$, used along to reverse calculate the stoichiometric coefficients of the produced hydrocarbons from C-1 to C-100. Only paraffins are considered, and the stoichiometric coefficients are adjusted for the C-30+lumping assumption. The syncrude is then sent to a refining section.

Hamad [106] developed an Aspen Plus model for the FT synthesis as part of an analysis of the solvent selection for supercritical Fischer–Tropsch synthesis reactors, in order to provide a basis for future supercritical phase simulations. The method used for this research is derived from the ASF distribution and the calculations of the stoichiometric coefficients are done in an Excel spreadsheet. For the calculation of α as temperature-dependent, the Levenberg–Marquardt algorithm was used. Products are assumed to be composed only by paraffins. The Aspen Plus reactor type chosen is not specified, as well as the *Property Method* used.

Dahl [107] modelled the FT process in the framework of a study evaluating a power and biomass to liquid (PBtL) process concept, which consists of biomass gasification to produce syngas, hydrogen addition to the syngas to increase its H₂/CO ratio, and FT synthesis to produce hydrocarbons, which are then separated and the longer hydrocarbons cracked. The author developed two Aspen Plus models for the Fischer-Tropsch reactor: a conversion-based model, whose operating conditions were varied, and a kinetic-based model, using a plug-flow reactor (RPLUG) in which the conversion of CO is studied. The Property Method used is Peng Robinson-Boston Mathias (PR-BM). The conversion-based involves two stoichiometric reactors (RSTOIC). In the first reactor, the three main FT reactions occur (paraffin, olefin, and alcohol formation), modelled according to the ASF distribution with a method of lumping high-weight hydrocarbons described in Hillestad et al. [108]. The products



are assumed to be composed by paraffins (up to C-20 with a C-21 + lump), olefins (up to C-10 with a C-11 + lump), and alcohols (up to C-5 with a C-6+lump). Each FT product is characterized by its specific probability function α : in the case of paraffins and olefins, α is assumed to be temperature- and pressure-dependent and formulated through Todic et al.'s correlation [109], while for the oxygenates, α is assumed to be constant (i.e. equal to 0.5). To figure out the ASF underestimation of methane in the products, a standalone reaction describing the CH₄ formation from CO and H₂ is added into the reactor, as well as an additional reaction for CH₄ among the olefins formation. A total CO fractional conversion has been set to 60%, which is split between the four reactions involved in the first RSTOIC reactor; its values were adjusted for each reaction in order to yield approximately the same carbon selectivities found in the experimental study carried out by Shafer et al. [110]. The second RSTOIC reactor considers the ethylene deviation, referring to Pandey et al. [84]. Computations have been performed in Aspen Plus calculation blocks. The kinetic-based model uses Langmuir-Hinshelwood (LHHW) adsorption kinetics and the consorted vinylene mechanism, a modified ASF distribution model. The model considers also ASF deviation and the effect of water over the reaction rate. The oxygenates are not included in this model. The product separation flowsheet is the same for both reactor configurations. After the Fischer-Tropsch synthesis, the liquid and gaseous FT products are separated in a FLASH at the temperature of the FT reactor outlet; in fact, at this temperature (more than 450 K), the hydrocarbons of 17 carbons or higher are liquid and unreacted syngas and hydrocarbon of 16 or lower are gaseous. The gaseous stream is then cooled in a countercurrent heat exchanger, which recovers the heat to warm the syngas feed, and then further cooled to 283.15 K. The cooled gaseous stream enters a three-outlet FLASH separator, which gives (i) a C1-C5 hydrocarbons gaseous stream, (ii) a C6-C16 hydrocarbons stream, and (iii) a water stream. The C6-C16 hydrocarbons stream undergoes further separation, involving a pressure decrease to 5 bar through a VALVE, a temperature increase to 483.15 K, in order to evaporate the C6-C7 hydrocarbons, and then a separation of the C6–C7 hydrocarbons from C8–C16 hydrocarbons in a second FLASH. Different operating conditions have been tested (i.e. 493.15 K, 27.6 bar, $H_2/CO = 2$; 483.15 K, 25 bar, $H_2/CO = 1.95$; 483.15 K, 25 bar, $H_2/CO = 1.60$; 483.15 K, 20 bar, $H_2/CO = 1.95$; 493.15 K, 25 bar, $H_2/CO = 1.95$) for both reactor configurations, and the simulations results showed that, at the same operating conditions, the conversion-based reactor results in higher selectivity towards lower-weight hydrocarbons. For both models, carbon selectivity increases with carbon number and has a peak around C-13.

Table 4 summarizes the selected research works modelling Fischer–Tropsch synthesis in Aspen Plus. Seventeen different studies have been identified.

4.5 Methanol production from syngas modelling

In the literature, several studies for simulating methanol production in Aspen Plus can be found; these describe the conversion of syngas and CO₂, to commercial-grade methanol.

4.5.1 Modelling MeOH production from syngas

Trop et al. [111] studied methanol production from a mixture of torrefied biomass and coal. In the process, gasification of biomass and coal, synthesis gas purification, and methanol synthesis from syngas have been simulated.

Authors modelled methanol synthesis as a series of plugflow reactions occurring in a stoichiometric reactor. Products are then cooled to 303.15 K and the condensed crude methanol is sent to a flash separator and a purification system, consisting of four distillation columns, where methanol is separated from water, small amounts of ethanol, and dissolved reactants. The composition (mass fractions) of the final products is made of 0.9998 methanol, 0.0001 ethanol, and traces of $\rm H_2O$ and $\rm CO_2$. For the methanol production section of the process, the Peng-Robinson thermodynamic method was chosen.

Gamero et al. [34] developed an Aspen Plus model for methanol synthesis from syngas obtained through gasified biomass and then gas cleaning through PSA. In the model, the cleaned syngas stream is compressed and heated up to the operating pressure and temperature. Then, the stream is introduced into the methanol synthesis reactor, which has been simulated as an equilibrium reactor (REQUIL). This kind of reactor requires the stoichiometry to be specified. The equations involved in the synthesis are (9) and (11), with a conversion of 36% for CO and 17% for CO₂. The selected catalyst is Cu/ZnO. The reaction product is then depressurized and cooled down, and then sent to a column separator, in order to condense and separate methanol from the gas phase, getting at the bottom pure methanol as the final product. The H_2/CO in the feed stream is about 2.4–2.5. As regards the thermodynamic method, the Peng-Robinson with Boston Mathias function was selected as appropriate for the process application, in particular for high-temperature gasification. Operating conditions have been varied, evaluated, and optimized through a sensitivity study. The optimal conditions were fixed at 493.15 K and 55.7 bar, yielding 32 kg/h methanol produced from a biomass feed rate of 100 kg/h.

Chein et al. [112] modelled methanol synthesis in Aspen Plus from syngas produced from biogas. After being compressed and cooled to the operating conditions for methanol



 Table 4
 Aspen Plus simulation models reviewed for FT synthesis

| Adelung et al. [30] | Publication year 2020 | Publication year Inermodynamic method | Main simulation blocks | Kererence model(s) | Kemarks |
|------------------------|-----------------------|---|---|--|--|
| Adelung et al. [30] | 2020 | | | | |
| | | PR-BM | RSTOIC reactor with CO conversion of 40%, flash separator, hydrocracker unit | Vervloet et al. expression for the calculation of α | synthetic kerosene and diesel from syngas derived from captured CO ₂ and H ₂ obtained through water electrolysis |
| Ashraf et al. [8] | 2015 | Not specified for this part of the process | RYIELD reactor with CO conversion of 80%, RADFRAC distillation column for product separation | ASF distribution model with α parameters from Kruit et al | FT liquids from biogas |
| Bao et al. [97] | 2010 | NRTL-RK | Reactor block type not specified | ASF distribution model with constant α | Optimal process design of a GTL plant |
| Campanario et al. [77] | 2017 | PENG-ROB for FT, UNIQUAC for distilla- tion train | RSTOIC reactor, cooler, decanter, distillation section | ASF distribution model, Song et al. correlation for α calculation | Low-temperature Fischer-Tropsch products from syngas obtained by supercritical water reforming of bio-oil aqueous phase |
| Cinti et al. [98] | 2015 | Not specified | RSTOIC reactor, splitter and mixer blocks, flash separator | ASF distribution model with constant α | Synthetic green fuels produced by a system integrating a solid oxide electrolyzer and the Fischer-Tropsch process |
| Dahl [107] | 2020 | PR-BM | RSTOIC RPLUG with LHHW kinetics | ASF distribution, and deviations Hillestad et al. [108], Shafer et al. [110], Pandey et al. [84], Todic et al.'s correlation [109] | Evaluate a power and biomass to liquid (PBtL) process concept |
| Gabriel et al. [105] | 2014 | Not specified | RSTOIC reactor | ASF distribution model with constant α | Gas-to-liquid (GTL) process involving syngas generation from natural gas and Fischer–Tropsch synthesis |
| Hao et al. [83] | 2008 | Not specified | RPLUG reactor with detailed kinetic models programmed in FORTRAN | User-defined detailed kinetic models | Gas-to-liquid (GTL) process involving syngas generation through ATR and |
| | | Not specified | RCSTR reactor with detailed kinetic models programmed in FORTRAN | User-defined detailed kinetic models | Fischer–Tropsch synthesis |
| Hamad [106] | 2011 | Not specified | Not specified | ASF distribution model with Levenberg–Marquardt correlation for α calculation | Analysis of Solvent Selection for Super- critical Fischer–Tropsch Synthesis Reactors |
| Er-rbib et al. [82] | 2012 | PR-BM | RSTOIC reactor with 42 reactions for FT, distillation columns, RYIELD reactor for hydrocracking | Specified equations | Synthetic gasoline and diesel fuels from syngas |
| Marchese et al. [101] | 2020 | RKS-BM | RPLUG with LHHW formulation | ASF distribution model and ASF deviations | Power to Liquid routes analysis (carbon capture and utilisation from biogas upgrading) |
| Marchese et al. [104] | 2021 | RKS-BM | Not specified | ASF distribution model, Song et al. correlation for α calculation | Biomass-to-X-plant: FT synthesis from digestate gasification |
| Marchese et al. [103] | 2021 | RKS-BM | RPLUG with LHHW formulation | ASF distribution model | CO ₂ from direct air capture as feedstock for FT synthesis |
| Michailos et al. [95] | 2017 | RKS-BM | RYIELD reactor interfacing an Excel Spreadsheet, flash separator, hydroc- racking unit | ASF distribution model | FT fuels from bio-syngas |



| • | | | | | |
|---------------------|------------------|--|--|---|---|
| Authors | Publication year | Publication year Thermodynamic method Main simulation blocks | Main simulation blocks | Reference model(s) | Remarks |
| Niassar et al. [92] | 2018 | PENG-ROB | RSTOIC reactor with 31 reactions, coolers, separators | Specified equations | Development and optimization of an integrated process configuration for IGCC power generation technology with a Fischer–Tropsch fuels from coal and biomass |
| Pondini et al. [99] | 2013 | not specified | RSTOIC reactor | ASF distribution model, Song et al. correlation for α calculation, Rane et al. [100] | FT crudes from biomass-derived syngas (gasification) |
| Sudiro et al. [96] | 2009 | PR-BM, NTRL | RYIELD reactor, distillation columns, RYIELD reactor for hydrocracking | Selectivity values from Mulder H | Synthetic fuels production through LTFT synthesis of syngas obtained from coal and natural gas |

synthesis (5 MPa and 523.15 K), the syngas enters the methanol reactor, modelled as an equilibrium reactor. This kind of reactor can simulate thermodynamic equilibrium reactions with good accuracy. Products (methanol and water) are expanded and then separated in a flash unit. Recycling unreacted syngas to improve the methanol yield is also performed by using a splitter with a recycle ratio. In addition, this study also estimated the performance of a green process for methanol synthesis using captured $\rm CO_2$ as a feedstock. Optimized conditions for obtaining 25.48% methanol yield have been found.

Ortiz et al. [39] developed an Aspen Plus simulation model for methanol synthesis from syngas obtained by supercritical water reforming of glycerol. In the model, the syngas is compressed and heated to the MeOH synthesis operating conditions (86.13 bar and 523.15 K). Then, the stream is sent to the methanol reactor, modelled as a stoichiometric (RSTOIC) gas-phase reactor, with specified CO conversion of 20% and CO₂ conversion of 3%. According to the authors, this kind of reactor can better represent an industrial reactor, as an equilibrium reactor (REQUIL) would lead to a CO conversion and CO₂ conversion of about 76.8% and 9.6% respectively (thermodynamic limits). In the study, it is assumed that no by-product formation occurs. The reactor effluent is then cooled down to condensate the crude methanol, and thus separated from the gas phase, which is then recompressed and recycled to increase the overall CO conversion to methanol. A fraction of the recycled gas is purged to prevent the accumulation of inert gases and sent to the furnace to support energy self-sufficiency of the process. Finally, crude methanol is sent to a distillation column, in which H₂O is separated from CH₃OH, with a recovery of 99%.

De María et al. [113] simulated methanol production from syngas in Aspen Plus to investigate a kinetic model developed by the authors. For this matter, an external model of the reactor was integrated into the simulation flowsheet instead of using a reactor block already available in Aspen. The reactor model was developed in Matlab and integrated into the Aspen Plus flowsheet using CAPE OPEN standard. The simulated process is constituted by a first part dedicated to syngas compression from 1 to 110 bar in a two-stage intercooled (311.15 K) compression train. Then, syngas is mixed with recycled streams, preheated in a feed-effluent heat exchanger, and then sent to a distillation column for raw methanol separation. The property method chosen for the whole process is the RK-Aspen (Redlich-Kwong), with the only exception of the distillation column, for which the NRTL-RK was set. The composition of the feed syngas is equal to 6.9% mol CO₂, 23% mol CO, 0.2% mol H₂O, 67.5% mol H₂, 0.3% mol N₂, and 2.2% mol CH₄. The product stream is characterized by 98.1% mol CH₃OH, 0.2% mol CO_2 , and 1.6% mol H_2O .



Table 4 (continued)

4.5.2 Modelling MeOH production from CO₂

Suhada et al. [114] developed an Aspen Plus simulation model to convert the CO_2 separated from biogas to methanol. The methanol reactor, which is also fed by a stream of H_2 obtained from an electrolysis unit, has been modelled as an equilibrium reactor type (REQUIL).

Another simulation approach for modelling methanol synthesis from captured CO₂ has been developed by Atsonios et al. [115]. Two different reactor types have been investigated: a tubular catalytic reactor and a zeolite membrane reactor. The tubular catalytic reactor is composed by three main units, i.e. methanol synthesis, gas separation, and product purification. The inlet gas is constituted by H₂ and CO₂, with a H₂/CO₂ ratio of 3.0, heated to 423.15 K, and then sent to the methanol reactor. The authors do not specify which Aspen Plus reactor block has been chosen for the simulation. However, the process stoichiometry and kinetics is considered. In particular, the reactions involved are (9), (10), and (11), and the process kinetics follows the study of Graaf et al. [116], developed for a commercial Cu/Zn/Al catalyst. The composition of the stream exiting the reactor is equal to 14.5% mol CO₂, 63.3% mol H₂, 6.9% mol H₂O, 1.8% mol CO, and 7.5% mol CH₃OH. The crude methanol is then subject to a refining step, made of two flash separators and a distillation column, leading to a final product composition of 99.3% mol CH₃OH, 0.1% mol H₂O, and 0.6% mol CO₂.

About the membrane reactor, it has been modelled as a series of equilibrium reactors with the intermediate interpolation of split separators. The split fraction of the vapours (only water and methanol) that are assumed to permeate the membrane is specified by a determined separation factor. The methanol purity of the product exiting the process is about 99.4%.

Van-Dal et al. [117] also simulated in Aspen Plus the methanol synthesis from captured CO₂ via hydrogenation. In the simulation model, CO₂ (1 bar, 298.15 K) is compressed to 78 bar in a series of intercooled compressors, while H₂ (30 bar, 298.15 K) is compressed to 78 bar in a single stage. The two gases are mixed and then re-mixed with the recycle stream, heated to 483.15 K, and finally injected into the RPLUG reactor for methanol synthesis, which is a fixed bed adiabatic reactor. The stream leaving the reactor is then split into two streams, one used to heat the fresh feed and the other in the reboiler and to heat the feed of the distillation column. These streams are then re-mixed, cooled to 308.15 K by water, and then sent to a knock-out drum, where condensed water and methanol get separated from the non-reacted gases, which are partially purged to minimize the accumulation of inerts and by-products in the reaction loop. The crude methanol obtained (composed of CH₃OH, H₂O, and residual dissolved gases) is expanded to 1.2 bar through two expansion valves, fed into a flash separator where residual gases are almost completely removed, heated to 353.15 K, and finally sent to a distillation column (RAD-FRAC). Here, water and methanol are separated, and the resulting CH₃OH stream, in gaseous form, contains 69 ppm wt. of H₂O and some unreacted gases. The RPLUG reactor is packed with a fixed bed of Cu/ZnO/Al₂O₃ catalyst. The model of Bussche and Froment [118] describing the reactions of methanol production and the RWGS reaction with this catalyst has been chosen, with readjusted parameters of Mignard and Pritchard [119]. The kinetic constants follow the Arrhenius law, while the equilibrium constants are provided by the study of Graaf et al. [116]. In Aspen Plus, the LHHW (Langmuir-Hinshelwood-Hougen-Watson) kinetics has been selected. Achieved CO2 conversion was 33%. About the used thermodynamic method, the Redlich-Kwong-Soave equation of state with modified Huron-Vidal mixing rules (RKSMHV2) was used for streams at high pressure (> 10 bar), while for low pressure streams the NRTL-RK model was employed.

In the thesis work developed by Mantoan [120], MeOH production from CO2 hydrogenation was simulated following the model developed by Fortes et al. [121], and adopts the similar approach seen in Van-Dal et al. [117]. Also, in this case, the methanol synthesis process involves a first step, in which the feed gases are compressed up to reactor feed pressure through different intercooled compression stages; a second process step, in which the pressurized feed stream gets is heated up and sent to the reactor; and a third process step, in which MeOH is finally separated from H₂O in a distillation column. As in the study of Van-Dal et al. [117], the kinetic model used is that of Bussche and Froment [118] with readjusted parameters of Mignard and Pritchard [119], the kinetic constants follow the Arrhenius law, and the equilibrium constants are given by Graaf et al. [116]. The thermodynamic models used are the RKSMHV2 for highpressure streams (> 10 bar) and NRTL-RK for low pressure. The reactor type selected is an adiabatic ideal plug flow reactor (PFR), following Eqs. (10) and (11) and reaction rates implemented in CHEMCAD®. The CO₂ conversion into CH_3OH in the reactor is ~21%. The distillation column has been designed to yield a high MeOH purity (>99.9% wt). The stream fed into the reactor has a composition of 13% wt H₂, 75% wt CO₂ and 12% wt CO. The product stream exiting the MeOH reactor has a composition of 12% wt CH₃OH, 7% wt H_2O , 11% wt H_2 , 58% wt CO_2 and 12% wt CO. The product methanol stream leaving the system has a composition of: 99.96% wt CH₃OH, 0.01% wt H₂O and 0.03% wt CO₂.

Calogero et al. [122] also simulated methanol synthesis through CO₂ hydrogenation, referring to the models developed by Atsonios et al. [115] and Van-Dal et al. [117]. The process consists of two main parts, i.e. a preparation section, in which the reactants are brought to the process conditions, and a processing section, in which the synthesis reaction



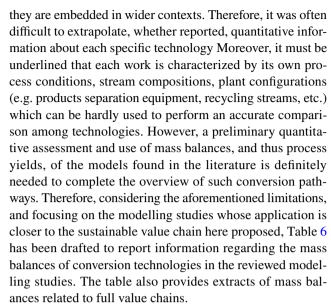
takes place and the separation of products and recirculation occurs. The process conditions and scheme follow those of the study developed by Atsonios et al. [115], with the aim to obtain a product methanol purity equal to 99.9% mol.

Kiss et al. [123] used Aspen Plus to simulate a process for methanol synthesis from CO₂ and wet hydrogen by-product from chlor-alkali production. In the simulated process, reactants are first brought to the required temperature and pressure, mixed, and then fed to the reactor, with is simulated by a plug flow reactor (PFR) using the LHHW kinetics. The reactor outlet contains products (CH₃OH and H₂O) as well as unconverted reactants (CO_x and H₂), and then this gaseous mixture is cooled and flashed to separate the condensable products from the non-condensable reactants, which are recycled. The condensed components are then separated in a distillation column, to get into lights (dissolved CO_x and minor light impurities), MeOH and water. The authors included a stripping unit in the process, in which wet hydrogen flows in counter-current mode with the condensed water mixture from the high-pressure low-temperature separator after the reaction. In this way, there is a complete recycle of CO₂ as CO/CO₂ is removed from the methanol-water mixture, and at the same time, by removing water from the wet hydrogen, there are no negative impacts on the reaction equilibrium conversion. The chosen Property Methods are the Soave–Redlich–Kwong EOS and NRTL with Henry components. The feed stream entering the PRF reactor is composed of 2.96% mol CO, 25.14% mol CO₂, 71.39% mol H₂, and 0.51% mol CH₃OH. The product stream exiting the PRF reactor has a composition of 3.24% mol CO, 22.78% $mol\ CO_2,\,63.95\%\ mol\ H_2,\,4.74\%\ mol\ H_2O,\,and\,5.29\%\ mol$ CH₃OH. The methanol stream exiting the system is characterized by a molar concentration of 0.01% mol H₂, 0.02% mol H₂O, and 99.98% mol CH₃OH. The H₂ conversion is 18.17%, while the CO₂ conversion is 17.20% and the MeOH yield of the overall process is 99.83%. In this process, all the carbon from the CO₂ feed is converted into MeOH product, whereas only two-thirds of H2 are converted into MeOH product, while the rest is converted to the water by-product.

Table 5 summarizes the reviewed studies on modelling methanol synthesis. The analysis revealed that the process (either from syngas or CO₂ as main feedstock) has been largely simulated in Aspen Plus. On the other hand, no studies simulating the process of direct methane to methanol (DMTM) conversion in Aspen Plus have been found in literature, probably because the technology is still in the developing phase.

5 Preliminary quantitative assessment

In most of the reviewed modelling studies, the technologies suitable for the value chain that we investigate in the current paper were not simulated as stand-alone units, but



The study developed by Ashraf et al. [8] is of particular interest, as the pathway encompassed is extremely coherent to that proposed in the present article. In this research, authors considered a relatively small-scale biogas to FT liquid conversion route, including biogas upgrading through PWS, biomethane reforming through DMR, syngas cleaning through chemical absorption with MEA, FT synthesis, and product upgrading in distillation columns. The study showed that, for 10,000 Nm³/h of dry biogas, the process requires 7.08 MW of power in addition to 35 of heating and 185 GJ/h for cooling, and from 4000 kg/h of methane in the biogas feed, 1602 kg/h of FT products could be produced. The process investigated is thus characterized by a carbon conversion efficiency of 45% and energy efficiency of 30%.

Another article that could be taken as a reference to study the sustainable biofuel production chain proposed is that of Bao et al. [97], even though the feedstock considered in the process is natural gas. The simulated plant converts 900,000 kg/h of natural gas to 118,000 bpd of products, through a conversion route involving reforming, FT reaction, and product upgrading.

6 Discussion

The aim of the research was to analyse a sustainable value chain, producing low-ILUC-risk biofuels (FT liquids and MeOH) and investigate the possibility to model it with a commercial software. The low-ILUC nature of the value chain is ensured by the implementation of the Biogas Done Right (BDR) model for the production of biomethane in a farm-scale decentralized configuration; the biomethane produced through this sustainable pathway serves as a basis for the production of synthetic fuels and chemicals in



 Table 5
 Studies reviewed modelling MeOH synthesis in Aspen Plus

| | 0 | , | 7 | | | |
|-----------------------|----------------------------|--|--|--|--------------------------------|---|
| Authors | Publication year Feedstock | r Feedstock | Thermodynamic method | Main simulation blocks | MeOH % in the products Remarks | Remarks |
| Atsonios et al. [115] | 2015 | CO ₂ | Not specified | Reactor type not specified (reaction kinetics considered), flash separators, distillation column | 99.3% mol | Convert captured CO ₂ into MeOH |
| Calogero et al. [122] | 2018 | CO_2 | Not specified | RPLUG reactor with LHHW kinetics | 99.9% mol | Convert captured CO ₂ into MeOH, based on Atsonios et al. and Van-Dal et al. studies |
| Chein et al. [112] | 2021 | Syngas and syngas + captured CO ₂ | Not specified | REQUIL reactor, flash separator | Not specified | Methanol production from biogas |
| De María et al. [113] | 2013 | Syngas | NRTL-RK | User-defined reactor developed in Matlab, distillation column | 98.1% mol | Methanol production from syngas to investigate a kinetic model developed by the authors |
| Gamero et al. [34] | 2018 | Syngas | PR-BM | REQUIL reactor, distillation column | Not specified | Methanol production from biomass |
| Kiss et al. [123] | 2015 | CO ₂ | SRK, NRTL | RPLUG reactor with LHHW kinetics, distillation columns, flash separator | 99.98% mol | MeOH from CO ₂ and wet hydrogen by-product from chlor-alkali production |
| Mantoan [120] | 2019 | CO_2 | NRTL-RK for low pressures (<10 bar), RKSMHV2 for high pressures (>10 bar) | RPLUG with reaction rates implemented in CHEMCAD, flash separator, distillation column | 99.96% wt | Convert captured CO_2 into MeOH. Follows Van-Dal et al.'s simulation model |
| Ortiz et al. [39] | 2012 | Syngas | PSRK | RSTOIC reactor with CO conversion 99% MeOH recovery per pass of 20% and CO2 conversion per pass of 3%, distillation column | 99% MeOH recovery | Methanol production from syngas obtained by supercritical water reforming of glycerol |
| Suhada et al. [114] | 2020 | CO_2 | Not specified | REQUIL | Not specified | Used to convert the CO ₂ separated from biogas to methanol |
| Trop et al. [111] | 2014 | Syngas | PENG-ROB | RPLUG rectors, flash separator, distillation columns | 99.98% mass | Methanol production from a mixture of torrefied biomass and coal |
| Van-Dal et al. [117] | 2013 | CO ₂ | NRTL-RK for low pressures (<10 bar), RKSMHV2 for high pressures (>10 bar) | RPLUG reactor with LHHW kinetics, RADFRAC distillation column, flash separator | Not specified | Convert captured CO ₂ into MeOH |
| | | | | | | |



 Table 6
 Mass balances reported in the reviewed modelling studies

| Technology | Authors | Process feed | | | Process output | | | Remarks |
|---------------------|---------------------------|-----------------------------------|----------|--------------------|-----------------------------------|---------------------------|-------------------------|--|
| | | Stream | Amount | Unit | Stream | Amount | Unit | |
| Overall value chain | Ashraf et al. [8] | Raw biogas | 10,000 | Nm ³ /h | Syncrude | 1602 | kg/h | Biogas to FT liquids conversion (PWS, DMR, MEA, FT, and upgrading) |
| | Bao et al. [97] | CH ₄ | 854,961 | kg/h | Diesel LPG Naphta | 139,170 6310 62,210 | gal/h gal/h gal/h | NG to FT liquids conversion (reform- ing, FT reaction, and upgrading) |
| AD | Scamardella et al. [61] | Biomass | 2 | t/day | Biogas | 157 | Nm ³ /d | Feed is composed of fruit waste |
| Upgrading—PWS | Bortoluzzi et al. [67] | Biogas | 1.60E-01 | kg/s | Biomethane | 4.81E-02 | kg/s | |
| | Cozma et al. [124] | Biogas | 604.558 | kg/h | Biomethane | 229.11 | kg/h | |
| | Gamba et al. [70] | Biogas | 49.5 | kmol/h | Biomethane | 29.2 | kmol/h | |
| | Menegon et al. [66] | Biogas | 500 | Nm ³ /h | Biomethane | 276.05 | Nm ³ /h | |
| | | Biogas | 1000 | Nm ³ /h | Biomethane | 649.23 | Nm ³ /h | |
| Upgrading—MEA | Gamba et al. [70] | Biogas | 51.9 | kmol/h | Biomethane | 32.8 | kmol/h | |
| Upgrading—MDEA | Gamba et al. [70] | Biogas | 50.2 | kmol/h | Biomethane | 32.8 | kmol/h | |
| Reforming— | Er-Rbib et al. [19] | CH ₄ | 122.326 | t/h | Syngas | 715.826 | t/h | H ₂ /CO ratio about 2 |
| DMR + SMR | | CO_2 | 330 | t/h | | | | _ |
| | | H_2O | 263.5 | t/h | | | | |
| FT synthesis | Er-Rbib et al. [19] | Syngas | 715.826 | t/h | Diesel | 67.1 | t/h | |
| | | | | | Gasoline | 25 | t/h | |
| | | | | | LPG | 0.3 | t/h | |
| | | | | | Other chemicals | 0.6 | t/h | |
| | Cinti et al. [98] | Syngas | 1532 | mol/h | Gasoline | 0.15 | bbl/day | H2/CO = 2.1 |
| | | | | | Diesel | 0.43 | bbl/day | |
| | | | | | WAXC30 | 0.36 | bbl/day | |
| | | | | | Total FT products | 1 | bbl/day | |
| | Campanario et al. | Syngas | 15,250.2 | kg/h | FT diesel | 1374 | kg/h | C5-C9 |
| | [77] | | | | FT jet fuel | 898 | kg/h | C10-C13 |
| | | | | | FT gasoline | 497 | kg/h | C14-C20 |
| | Sudiro et al. [96] | Syngas | 268 | t/h | Gasoline | 28.41 | t/h | $H_2/CO = 2$ |
| | | | | | Diesel | 69.9 | t/h | |
| | | | | | GPL | 1 | t/h | |
| | | | | | Light gas + unre- acted syngas | 35.7 | t/h | |
| MeOH synthesis | De María et al. [113] | Syngas | 11,449.9 | kmol/h | МеОН | 3141.6 | kmol/h | MeOH from syngas, $H_2/CO = 2.94$ |
| | Kiss et al. [123] | Syn- gas + unre- acted prod | 122,003 | kg/h | МеОН | 12,508.7 | kg/h | MeOH from CO ₂ hydrogenation, the reactor is fed with CO ₂ , H ₂ , and recycled unreacted products; MeOH |
| | Perèz-Fortes et al. [121] | CO ₂ | 80.5 | t/h | МеОН | 55.1 | t/h | MeOH from CO_2 hydrogenation, CO2 wt% = 100%, $H_2 \text{ wt\%} = 100\%$ |
| | | H_2 | 11 | t/h | | | | MeOH wt $\% = 99.96\%$ |
| | Van-Dal et al. [117] | CO_2 | 88 | t/h | MeOH | 59.3 | t/h | MeOH from CO ₂ |
| | | H_2 | 12.1 | t/h | | | | hydrogenation |



centralized refineries, making use of the natural gas grid to connect the two environments (farm and refinery). Existing and under-development technologies for the conversion of biomass-to-liquid fuels have been reviewed, and their modelling in the various studies available in the literature analysed, focusing on model assumptions, process conditions, applications, yields, and consumptions.

After a quality-based initial screening, a total of about 57 reviewed modelling papers were selected among the available literature; these included anaerobic digestion, biogas cleaning and upgrading to biomethane, methane reforming to syngas, syngas conversion to hydrocarbons through Fischer–Tropsch synthesis and MeOH synthesis from syngas and CO₂ hydrogenation.

In the reviewed papers, many of the technologies of interest to this paper have not been modelled as stand-alone cases, but are embedded in broader contexts, with more complex flowsheets. For this reason, it was often difficult to extrapolate detailed information on yields and consumptions of the individual processes from their original framework. For example, in the study of Cinti et al. [98], aimed at simulating the production of synthetic green fuels through a system integrating solid oxide electrolyzer and FT synthesis, our analysis did not cover the electrolyzer unit, but was focused only on modelling the Fischer–Tropsch synthesis.

As discussed, the first step of the value chain under investigation foresees the anaerobic digestion process following the BDR model to produce biogas in a sustainable manner. In the literature, no studies that specifically apply this model in Aspen Plus simulation environment have been found (Fig. 5). Therefore, we searched for generalized anaerobic digestion models, which turned out to be not so numerous (i.e. 6); this could be due to the high complexity of the process, which turns out to be particularly difficult to model, especially as regards the biological activity of microorganisms. Nguyen et al. [60] and Scamardella et al. [61] opted for a one-stage model in a single reactor, in which the whole AD phases occurs. In the RSTOIC reactor (used in Nguyen et al. [60]), the reaction kinetics is not considered, whereas in the RCSTR (used in Scamardella et al. [61]) detailed information on the reactions and their kinetics must be input; this means that the first represents a more simplistic modelling approach. However, the two studies ([60] [61]) refer to different AD models (i.e. Buswell equation, ADM1, and comprehensive models), and the papers do not provide information showing a comparison with experimental data. Al-Rubaye et al. [56] and Rajendran et al. [59] adopted a very similar approach, i.e. a two-step model using a RSTOIC reactor for the hydrolytic phase and a RCSTR reactor for the other AD phases, in reference to the ADM1 and comprehensive models. The two studies have been validated against experimental data, resulting in both cases in good agreement.

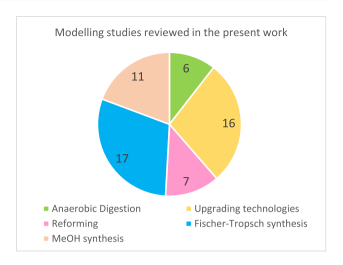


Fig. 5 Number of selected studies reviewed and based on Aspen Plus modelling.

The AD model developed by Serrano Peris [57], which involves two RSTOIC reactors in series, does not take into account the hydrolytic step, and its application is thus limited to post-hydrolyzed wastes. Besides, a higher modelling complexity is given in that of Llanes et al. [62], which integrates ADM1, flow pattern and biofilm characteristics implemented through FORTRAN subroutines; this model is thus not limited to completely mixed flow pattern (as in the case of RCSTR reactors), and results are in agreement with experimental data.

Biogas upgrading to biomethane is a fundamental step in the proposed sustainable value chain, as biomethane must comply with strict Country-specific technical standards in order to be injected into the natural gas grid. Therefore, accurately modelling this step is crucial. In the literature, there is a relevant number of papers (i.e. 16) simulating upgrading processes; these are in large part set in the context of gas sweetening, but nevertheless, in some studies ([8, 34, 39]), the upgrading technologies are employed for syngas upgrading (i.e. CO₂ removal). Many of the upgrading processes modelled in Aspen Plus turned out to be physical absorption (i.e. PWS) and chemical absorption using amine solutions (i.e. MEA, DEA, MDEA). The modelling approach of these technologies is almost the same in every paper: a distillation column for both absorption and stripping processes. The Aspen Plus distillation unit operation chosen in most of the reviewed simulations was the RADFRAC column, which is the most generic column block type. Some models have opted for an equilibriumstage approach for both absorption and stripping column, which assumes that each plate of the column is a theoretical plate (equilibrium plate), and thus, the vapour and the liquid leave any plate at thermodynamic equilibrium [125]. In other simulations, instead, the distillation columns are



rate-based, which means that the mass and energy transfer across the interface are taken into account using rate equation and mass transfer coefficients; therefore, this approach provides a more rigorous modelling of the columns. At the same time, there are studies ([66, 69]) that opted for a combined equilibrium-based and rate-based approach.

Not many models simulating the pressure swing adsorption process in Aspen Plus were found in the literature. As a matter of facts, the PSA process is a dynamic process, and as Aspen Plus is a steady state calculator, no dynamic options are available in the software. For this reason, many authors preferred to model the process in Aspen Adsorption, this being a comprehensive flowsheet simulator more specific for adsorption processes. Finally, only one study (i.e. [61]) modelling the biogas upgrading through membrane separation has been found, consisting in a user-defined model developed in Excel referring to Fick's law with diffusive model assumptions.

About the modellization of the reforming technologies, a discrete number of studies have been found (i.e. 7), most of them assessing the SMR process, which is the most consolidated one in this matter. A good number of papers addressed the process of dry reforming, which is an emerging technology whose reaction kinetics has not been fully described yet. Anyway, there are many studies in which DMR is combined with SMR, as well as POX, while a smaller number of articles assessing the ATR process was found. Most of the reviewed models are aimed at simulating processes for hydrogen production in spite of syngas production, and consequently, in many studies, SMR is combined with WGS. The modelling approaches of the analysed studies are various: the Aspen Plus reactor blocks employed in the simulations are RGIBBS ([8, 81, 83]), REQUIL ([79, 82]), and RPLUG with Langmuir-Hinshelwood-Hougen-Watson (LHHW) kinetics ([16, 80]). The RPLUG reactor type that unlike the RGI-BBS and the REQUIL, encompasses the process reaction kinetics and allows for a complete description of the process under variable conditions and proper reactor sizing; however, a suitable reaction set and the relative kinetic model and thermodynamic data must be provided [126].

A large number of studies (i.e. 17) modelling the Fischer–Tropsch process in Aspen Plus have been found in the literature. As mentioned, the FT process is particularly complex and the full understanding of all mechanisms involved has not yet been reached, though it is not a novel technology. The reference model for most of the reviewed articles is the Anderson-Schulz-Flory (ASF) distribution, a fairly simple model which gives a reasonable description of FT products by representing the synthesis as an addition polymerization reaction with chain growth probability α [127] [91]. Nevertheless, despite the mathematical simplicity of the ASF model, studies have shown that there are

deviations of the FT product composition from the ideal distribution (i.e. higher methane selectivity, lower ethylene selectivity, increasing chain growth probability, and lower olefin-to-paraffin ratio with increasing carbon number) [101, 105, 107, 128]; therefore, correlations accounting for the ASF distribution deviations and dependence of α on process conditions have been developed and are well documented in the literature [89]-[84, 91, 100, 108-110]. In many modelling studies reviewed, the produced hydrocarbons have been assumed to be composed only by paraffins; this assumption is acceptable, as the alkanes are the main product of the FT synthesis. However, there is also a large number of studies considering olefins in the products, while only one model (i.e. Dahl [107]) among those found includes also alcohols. Moreover, it was noticed that in many studies ([95, 97, 98, 105]) the chain growth probability α has a constant value, while others use correlations to relate its values to the process conditions; the most used expression in the reviewed papers is the one derived by Song et al. [89]. Another significant parameter for determining the accuracy of a FT model is the carbon number (n) for the product chain termination; in the modelling studies reviewed, this number was found to be always higher than 20 for the main products (i.e. paraffins). Setting a high n for the hydrocarbon chain termination definitively represents a higher accuracy in describing the Fischer–Tropsch synthesis; in fact, the product slate synthesized ranges from methane up to C-120+[88]. However, as the product selectivity significantly decreases after peaking at around C-7 to C-13 (depending on the process conditions and the catalyst), cutting the chain from n = 20 could be considered satisfactory.

Most of the reviewed Aspen Plus models employ reactor types such as RSTOIC and RYIED, in which the reaction kinetics is not explicitly considered, and the product distribution is imposed. Nevertheless, a small number of modelling studies assessing the FT synthesis reaction kinetics (through external subroutine) were also found (i.e. [83, 101, 103, 107]); these models provide higher flexibility when changing the process conditions.

As concerns the MeOH synthesis from syngas, 5 studies have been found, whose modelling procedures are diverse. Some authors did not consider the reaction kinetics, opting for a RSTOIC reactor ([39]) or for a REQUIL reactor ([34, 112]), which performs chemical and phase equilibrium reactions. At the same time, De Maria et al. [113] and Trop et al. [111] described the reaction kinetics of the catalytic reaction, using, in the former case, a user-defined reactor type developed in Matlab, and a RPLUG reactor in the latter case.

Additionally, opportunities for further pathways have been investigated, including methanol synthesis from captured CO₂, which represents an excellent opportunity from an environmental sustainability point of view. Indeed, CCS and CCU technologies, if associated with bioenergy, can



make the production chain considered not only carbon neutral but also carbon negative. In this perspective, in the proposed bioliquid production chain, the CO₂ separated from biogas through upgrading processes represents again a further feedstock for the MeOH production, and in this way would not be dispersed in the atmosphere. Lots of studies simulating this in Aspen Plus have been found (i.e. 7), meaning that this topic is of great interest to the scientific community. The modelling approaches of the MeOH synthesis from CO₂ adopted in the reviewed papers are very similar to those of in the case of syngas as a feedstock: REQUIL reactor ([112, 114]) and RPLUG reactor with LHHW kinetics ([115, 117, 120, 122, 123]).

As an alternative to MeOH synthesis from syngas, the straight conversion of methane into methanol (DMTM) could be an interesting route, as it gives the possibility to bypass the very energy-intensive step of methane reforming, representing an economical-advantage and environment-friendly option. However, as mentioned, this method still needs to be improved to be suitable for industrial applications. As a matter of fact, no Aspen Plus models simulating this process have been found in the literature.

In view of these considerations, it can be concluded that the simulation system provides all the necessary tools to model the entire sustainable biofuel production chain selected for this work. Moreover, through browsing the suitable technologies (either consolidated or under development) that could be employed, and by analysing and commenting on the diverse options identified in literature, this study could serve as a guideline to assess the feasibility of bio-thermochemical conversion pathways.

The references collected in the literature could be used for further studies aimed at qualitatively and quantitatively assessing multiple aspects related to biofuels production, and besides the possibility to build an overall Aspen Plus simulation model of the proposed sustainable biogas-to-liquid production chain would be an interesting object for future works.

7 Conclusions

Biomass-to-fuel pathways are of interest for many sectors, in particular those relying on liquid fuels also during the short-medium term ecological transition, such as maritime and aviation. The present survey reviewed biogasbiomethane conversion pathways to Fischer–Tropsch liquids and methanol, as an alternative to fossil-based fuels. More specifically, a sustainable biofuel production chain, producing low-ILUC-risk biofuels, has been analysed, and the possibility to model it through a commercial simulation software has been explored. The low-ILUC-risk nature of the value chain is guaranteed by the

implementation of the Biogas Done Right (BDR) model for the production of biomethane in decentralized farms through anaerobic digestion. The produced biomethane constitutes the base to synthesize bio-liquids and biochemicals in centralized conversion plants, making use of the natural gas grid to connect the farm and the refinery environments. The value chain includes thus biogas cleaning and upgrading to biomethane, injection of biomethane into the natural gas grid resulting in downstream extraction, methane reforming to syngas, and syngas conversion to hydrocarbons through Fischer-Tropsch synthesis and to methanol through MeOH synthesis. Several suitable technologies and process solutions have been reviewed and, in addition, opportunities for further pathways to be included have been investigated, in order to integrate the proposed biofuel production chain even more into a perspective of sustainability; these are methanol production from captured CO₂ and direct methane to methanol (DMTM) conversion.

Moreover, a comprehensive literature review of studies modelling the identified conversion pathways using Aspen Plus process simulation software has been carried out, in order to evaluate the maturity of available simulation models. To this aim, 57 publications were selected, which have been deeply analysed from the point of view of plant configuration, modelling approach, and process yield, focusing on understanding the diverse schematizations and assumptions, as well as mass/energy balances. It has been observed that many of the technologies of interest to this article were not modelled as stand-alone studies, but were embedded in broader contexts; thus, it was often difficult to extrapolate detailed information on related yields and consumptions. Anyway, the survey showed that comprehensive and sufficiently complete kinetic data to model well-established technologies (i.e. SMR and MeOH synthesis) are available in the literature, and thus a considerable number of Aspen Plus simulation models addressing them exhaustively was found. Likewise, a large experience in modelling upgrading technologies, such as physical and chemical absorption, has been observed, as well as a remarkable coherence in the modelling approaches. At the same time, the mechanisms involved in the high-complexity processes embedded in the proposed value chain (i.e. AD and FT) still need to be fully understood, but yet studies simulating them satisfactorily and with a good match with experimental data have been found. However, the only technology for which no modelling studies were identified is DMTM, being this process still under investigation.

By investigating a broad spectrum of suitable biomassto-liquid conversion pathways, either consolidated or under development, and by analysing and commenting on the diverse options retrieved in literature studies, this work



could represent a guideline for further studies aimed to analyse qualitatively and quantitatively multiple aspects related to biofuels production.

Moreover, the review revealed that the commercial software chosen for the analysis provides the necessary tools to model the multiple technological solutions that could be employed in the identified sustainable biofuel production chain. Besides, this study contains enough references to build an overall biomass-to-liquid value chain simulation model, making it a solid basis for future works, such as optimizing operational conditions, determining limitations due to process configurations, and possibly exploiting the high potential of process waste streams to further support the sustainability of bioliquids production.

Author contribution Lorenzo Testa (lorenzo.testa@polito.it): data curation, formal analysis, investigation, methodology, visualization, writing — original draft; David Chiaramonti (david.chiaramonti@polito.it): conceptualization, writing and review — review and editing, supervision, methodology; Matteo Prussi (matteo.prussi@polito.it): writing — review and editing, conceptualization, visualization; Samir Bensaid (samir.bensaid@polito.it): writing — review and editing, visualization.

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Declarations

Competing interests The authors declare no competing interests.

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