Production of Fuels and Chemicals from a CO₂/H₂ Mixture

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Abstract: In order to explore co-production alternatives, a once-through process for CO₂ hydrogenation to chemicals and liquid fuels was investigated experimentally. In this approach, two different catalysts were considered; the first was a Cu-based catalyst that hydrogenates CO₂ to methanol and CO and the second a Fisher–Tropsch (FT) Co-based catalyst. The two catalysts were loaded into different reactors and were initially operated separately. The experimental results show that: (1) the Cu catalyst was very active in both the methanol synthesis and reverse-water gas shift (R-WGS) reactions and these two reactions were restricted by thermodynamic equilibrium; this was also supported by an Aspen plus simulation of an (equilibrium) Gibbs reactor. The Aspen simulation results also indicated that the reactor can be operated adiabatically under certain conditions, given that the methanol reaction is exothermic and R-WGS is endothermic. (2) the FT catalyst produced mainly CH₄ and short chain saturated hydrocarbons when the feed was CO₂/H₂. When the two reactors were coupled in series and the presence of CO in the tail gas from the first reactor (loaded with Cu catalyst) significantly improves the FT product selectivity toward higher carbon hydrocarbons in the second reactor compared to the standalone FT reactor with only CO₂/H₂ in the feed.

Keywords: Fischer–Tropsch synthesis; water gas shift reaction; methanol synthesis; CO₂ hydrogenation; Cu-based catalysts; cobalt-based catalysts

1. Introduction

Utilization of CO₂ as a raw material for the synthesis of valuable chemicals and fuels is of interest not only as carbon source but also as a way of mitigating its environmental impact as a greenhouse gas [1–3]. Methanol production is of interest due to its large global market as a bulk chemical product and as a key component in C₁ chemistry, in addition to its low hydrogen requirement during production [4–8]. Methanol synthesis from CO₂ hydrogenation occurs together with the reverse water gas shift (R-WGS) reaction and is subjected to three major limitations [1,9–11]: (1) the thermodynamic equilibrium limits the process to low conversion per pass and, therefore, a large recycle of unconverted gas would be needed to achieve a reasonable overall conversion. (2) A poor methanol selectivity due to the R-WGS reaction. (3) The reaction is also strongly exothermic and thus requires significant cooling duties. These limitations are largely responsible for the high investment and operating cost of methanol synthesis plants.

Various researchers have reported activity data, which clearly shows that CO₂ readily hydrogenated to hydrocarbons over Fischer–Tropsch (FT) catalysts [12–16], which is also another promising way of producing chemicals and fuels from CO₂. However, there is a high selectivity to CH₄ and short chain hydrocarbons, which may not be commercially attractive. Fortunately, results from previous work [15]
show that when even small amounts of CO are added to the feed, the product distribution changes to favor the production of higher hydrocarbons.

CO₂ hydrogenation to methanol and hydrocarbon fuels over their respective appropriate catalysts has been shown to be feasible, but it comes with some important limitations and undesirable by-products. However, in their recent work, Sempuga and Yao [17] have demonstrated the potential benefits of co-production of chemicals by CO₂ hydrogenation; the work looked at a number of CO₂ hydrogenation processes and showed that all of them are potentially inefficient in conserving the chemical potential of hydrogen in their final respective products. Co-production of chemicals seemed to be the only way that can fully conserve the chemical potential of hydrogen and can minimize heat duty requirements, thereby potentially leading to lower capital and operating costs. One of the ways of doing this is to couple methanol synthesis with Fischer–Tropsch synthesis (FTS) in series, where CO₂ and the byproduct, CO, from the R-WGS reaction occurring during methanol synthesis are converted to useful chemicals and transport fuels in downstream FT synthesis sections.

Furthermore, methanol, with its low price and high oxygen content, has been widely considered to be an additive in blended fuels; the addition of methanol could increase the heat of evaporation and reduce soot formation when the fuel is combusted [18,19]. Combining the reactions of CO₂ methanol synthesis, WGS and FT could convert CO₂ and H₂ to both methanol and diesel fuels. As a consequence, these two groups of products could be blended together to produce an alternative fuel with good combustion characteristics and reduced production of particulates.

In order to explore these co-production alternatives, laboratory experiments were conducted where methanol and hydrocarbons were co-produced by using a configuration of reactors in series. CO₂ hydrogenation over a copper-based catalyst (for methanol synthesis and WGS) and a cobalt based catalyst (for FT synthesis) under different reaction conditions were carried out in two plug flow reactors (PFR) which were configured in series. The catalysts activity and selectivity and product distribution were measured, analyzed and compared. Finally, a potentially optimized process for CO₂ hydrogenation to produce methanol and hydrocarbon fuels is suggested based on simulation and experimental results.

2. Experimental Set Up

2.1. Catalyst Preparation

A commercial Cu-based catalyst from Imperial Chemical Industries (ICI), London, UK was used for the methanol synthesis experiments. The test results showed that the catalyst was active for both the R-WGS and methanol synthesis reactions.

A supported Co catalyst (10% Co/90% TiO₂ by mass) was used for the FT CO₂ hydrogenation experiments. The catalyst was prepared by impregnation of TiO₂ (Degussa P25, Degussa Africa Ltd., Midrand, Gaudeng, South Africa) with the cobalt nitrate solution (Co(NO₃)₂·6H₂O, Sigma-Aldrich, St. Louis, MO, USA). The procedure for the catalyst preparation is described in Yao et al. [15].

2.2. Experimental Setup and Procedure

Three cases were investigated by using two plug flow reactors configured as shown in Figure 1: in Case A and Case B the reactors are set up to perform CO₂ hydrogenation separately, one using a Cu-based catalyst (Cu-reactor) and the other using a Co-based catalyst (Co-reactor). In Case C, the Cu-reactor is connected in series to the Co-reactor. The gas exiting the Cu-reactor is fed to the Co-reactor after it is cooled, and the condensate knocked out. The detailed experimental set up and procedure for each case is described below.
Figure 1. Simplified flow schemes of plug flow reactors used in the experiments for Case A, Case B, Case C and modified Case C: (1) plug flow reactor; (2) hot condensable product trap (150 °C); (3) cold condensable product trap (room temperature); (4) online gas chromatograph (GC).

**Case A: CO$_2$ hydrogenation over a Cu-based catalyst**

One gram of the Cu-based catalyst was loaded into the plug flow reactor. The reduction of the catalyst was done for 24 h at atmospheric pressure and at a temperature of 200 °C with H$_2$ (99,999%) at a flowrate of 60 mL/min. Once the reduction was completed, the reactor was allowed to cool down to room temperature. Feed gas, of composition H$_2$: 67.6%, CO$_2$: 22.6% and N$_2$ (as an internal standard for mass balance calculations) 9.8%, was then introduced into the reactor at a flow rate of 60 mL(NTP)/(min·gcat) (NTP stands for Normal Temperature and Pressure). The reactor was run at a pressure of 20 bar (gauge), and at temperatures ranging from 200 °C to 350 °C. The temperature at each set point was kept constant for 72 h during which the tail gas composition was measured frequently to ensure that the system reached steady state. The feed and reaction conditions for CO$_2$ hydrogenation over the Cu-based catalyst are listed in Table 1.

**Table 1.** Reaction and feed conditions for hydrogenation of CO$_2$: (a) a single reactor loaded with a Cu-based catalyst (Case A); (b) a single reactor loaded with a Co-based catalyst (Case B); and (c) the two reactors in series (Case C).

| Items                                           | Case A             | Case B             | Case C             |
|-------------------------------------------------|--------------------|--------------------|--------------------|
| Reactor                                         | One fixed bed      | One fixed bed      | Two fixed bed reactors in series |
| Catalyst weight (g)                             | Cu                 | Co                 | Cu                 |
| Feed: H$_2$/CO$_2$/N$_2$ (%)                    | 67.6%/22.6%/9.8%   | 67.6%/22.6%/9.8%   | 67.6%/22.6%/9.8% |
| Temperature (°C)                                | 200–350            | 200                | 200–350            |
| Flow rate (ml(NTP)/(min·gcat)) $^a$             | 60                 | 60                 | 60                 |
| Pressure (bar gauge)                            | 20                 | 20                 | 20                 |

$^a$ NTP stands for Normal Temperature and Pressure.

**Case B: CO$_2$ hydrogenation over a Co-based catalyst.**

In the same way as in Case A, one gram of the Co/TiO$_2$ catalyst was loaded into a plug flow reactor. The reduction of the catalyst was affected for 24 h at atmospheric pressure and at a temperature of
350 °C with H₂ (99.999%) at a flowrate of 60 mL/min. Once the reduction was completed, the reactor was allowed to cool down to room temperature. Feed gas, of composition H₂: 67.6%, CO₂: 22.6% and N₂ (as an internal standard for mass balance calculations) 9.8%, was then introduced into the reactor at a flow rate of 60 mL (NTP)/(min·gcat). The reactor was run at a pressure of 20 bar (gauge), and at a constant temperature of 200 °C for 72 h. The feed and reaction conditions for CO₂ hydrogenation over the Co-based catalyst are given in Table 1.

Case C: CO₂ hydrogenation in a reactor loaded with Cu-based catalyst in series with a reactor loaded with Co-based catalyst.

The experiments in Case C were conducted with a Cu-reactor and a Co-reactor configured in series. Initially, a trap was placed in between the reactors to condense the methanol and water produced in the Cu-reactor. The tail gas after this condensation trap was introduced into the second reactor. The experimental program for each reactor was exactly the same as when they were run separately, that is: the Cu-catalyst was run at temperatures ranging between 200 °C and 350 °C at constant pressure and flowrate, and the temperature in the reactor was kept constant for 72 h, and the Co-reactor was kept at the same temperature and pressure (200 °C and 20 bar) for all runs. This is summarized in Table 1. For each of these runs, the products and un-reacted materials passed from the exit of the first reactor through the product trap which set at room temperature to condense the water and methanol. The uncondensed gases were fed to the second reactor (see Figure 1 Case C).

With the aim to simplify the process, a modified process named “Modified Case C” was used, in which the condensation trap was removed, so that all the products and unreacted reactants were fed to the Co-reactor (see Figure 1: Modified Case C). The same reaction conditions as those used in Case C were used, and the only difference was that the temperature range was between 200 °C and 260 °C.

2.3. Product Analysis

The tail gas was analyzed every 90 min using an online DANI gas chromatograph (GC) (DANI Analitica, Milan, Lombardy, Italy). The online GC was equipped with four columns and three detectors for the analysis of H₂ (Molecular Sieve 5A column connected to a thermal conductivity detector (TCD)), N₂, CO, CO₂ (Porapack Q packed and molecular sieve 13X columns connected to a TCD) and gas phase hydrocarbon products (PoraPLOT Q-HT column connected to a flame ionization detector (FID). The aqueous from the methanol synthesis reactor was collected in a cold trap (kept at room temperature). Two condensable product traps were used after the FTS reactor: a hot trap (kept at 150 °C) was used to collect the wax products and a cold trap (kept at room temperature) was used for the collection of liquid products. The condensed products (aqueous, wax and oil) from the condensable product traps were collected for further analysis using two off-line GCs.

3. Results

3.1. Case A: CO₂ Hydrogenation Over a Cu-Based Catalyst

The effect of reaction temperature on the hydrogenation of CO₂ over a Cu-based catalyst is shown in Figure 2A. Figure 2A clearly shows that CO₂ conversion increases steadily with increasing reaction temperature. According to the product analysis, the main products for CO₂ hydrogenation were methanol, CO and water. A small amount of CH₄ (less than 1.5%) was detected. The product selectivity as a function of temperature is shown in Figure 2B; it can be seen that CO and CH₄ selectivity increases with temperature. Conversely, the methanol selectivity reduces with increasing temperature. Similar trends are observed when looking at the product formation rates (Figure 2C). The rates of formation of both CO and CH₄ increase steadily with increasing temperature. However, the rate of formation of methanol initially increases, reaches a maximum between 220 °C and 240 °C and then drops as the temperature is further increased.
The experimental data exhibit that two major competing reactions are involved in the CO$_2$ hydrogenation to methanol process; these are the methanol synthesis reaction and the R-WGS reaction. Equilibrium limits what can be achieved in a once-through process and this will determine the maximum amounts of CO$_2$ that can be converted to methanol and CO. Since CO$_2$ hydrogenation to methanol is an exothermic reaction and the R-WGS reaction is endothermic, analyzing the overall heat of reaction is of interest. This will determine the heat load of the reactor and reducing it could potentially reduce operating and capital costs. With the aim of testing how far our experimental results are away from the equilibrium limitation, Aspen Plus $^\text{®}$ V8.6 was used to investigate equilibrium for the methanol synthesis process. In the simulation the following assumptions are made:

- The base method is the Peng-Robinson Equation of State (EOS).
- The RGiibbs reactor was used to calculate equilibrium.
- The feed was: H$_2$ with a flow rate of 3 kmol/h and CO$_2$ with a flow rate of 1 kmol/h
- The outlet was: H$_2$, CO, CO$_2$, H$_2$O and methanol

The simulation results, namely the CO$_2$ conversion, the product selectivity and the heat of reaction as a function of reaction conditions (pressure and temperature) are plotted in Figure 3. It can be seen that: (1) the CO$_2$ conversion for pressures greater than 1 bar, decreases, goes through a minimum and then increases as the temperature is increased from 150 °C to 400 °C. It also shows that the higher the pressure, the higher the total CO$_2$ conversion. (2) Higher temperatures and lower pressures favor the production of CO, while conversely lower temperature and higher pressure favors methanol production. (3) The heat of the reaction is negative at high methanol selectivity and is positive at higher CO selectivity. It can also be seen that the heat of the reaction goes through zero (shown by the red dashed lines) indicating that an adiabatic process is possible at equilibrium. Table 2 summarizes the conditions when both $\Delta$H and $\Delta$G are equal to 0 kW for the methanol synthesis system.

Figure 4 compares the results from the experiments and those predicted by the Aspen simulation. The experimental and simulation results compare well from a temperature above 240 °C, suggesting that equilibrium was reached in the experiments run at this temperature or higher.
Therefore, a thermodynamic model based on equilibrium can be used to predict the behavior of the Cu-reactor at pressures which were not covered in these experiments; from the results (Figure 3), it can be concluded that methanol formation from CO\(_2\) is thermodynamically favored at low temperature and high pressure, whereas CO formation is favored at high temperature. It can also be seen that CO formation decreases with increasing pressure; however, this is due to the increase in methanol selectivity, rather than the increase in pressure, since the R-WGS reaction is not affected by pressure. Daza and Kuhn [20] also mentioned that the R-WGS reaction is equilibrium limited, and its conversion is affected by temperature as it is an endothermic reaction, rather than by pressure due to the stoichiometry of the reaction.

**Figure 3.** The predicted CO\(_2\) conversion, product selectivity and the change in the overall enthalpy of the reaction at equilibrium as a function of reaction conditions for methanol synthesis system. A: at 1 bar; B: at 20 bar; C: at 60 bar; D: at 80 bar.

**Table 2.** Summary of simulation results when both ∆H and ∆G are equal to 0 kW/mol for the methanol synthesis and reverse-water gas shift (R-WGS) system.

| CO\(_2\):H\(_2\) | P (bar) | T (°C) | CO\(_2\) Conversion (%) | Methanol Selectivity (%) | CO Selectivity (%) |
|-----------------|--------|--------|-------------------------|-------------------------|-------------------|
| 3:1             | 20     | 239.5  | 18.4                    | 40.0                    | 60.0              |
| 3:1             | 60     | 294.7  | 26.6                    | 38.3                    | 61.7              |
| 3:1             | 80     | 311.4  | 29.2                    | 37.7                    | 62.3              |

**Figure 4.** Comparison of the predicted equilibrium data and the experimental results for CO\(_2\) hydrogenation over a Cu-based catalyst. Reaction conditions: pressure 20 bar (gauge), flow rate 60 mL (NTP)/(min-gcat) and feed gas H\(_2\)/CO\(_2\)/N\(_2\) = 67.6%/22.6%/9.8%.
3.2. Case B: CO₂ Hydrogenation Over a Co-Based Catalyst

Hydrogenation of CO₂ over a Co-based catalyst was conducted under normal FTS reaction conditions and in this experiment all the conditions were kept constant, at a pressure of 20 bar gauge, temperature of 200 °C and at a feed gas flowrate of 60mL(NTP)/(min·gcat).

The results from the experiments are shown in Table 3. The results show that CO₂ hydrogenation over a Co-based catalyst produces mostly short chain saturated hydrocarbons (paraffins) rich in CH₄ (92.5%). These results correspond to results obtained by many other researchers [12–14]. Table 3 also includes results obtained from previous experiments [15] where hydrogenation was done with a feed containing pure CO (third column of Table 3) and where hydrogenation was done with a mixture of CO and CO₂ (fourth column of Table 3). By looking at the results in Table 3 the following observations can be made:

1. For CO₂ hydrogenation (second column), the product is almost entirely composed of short chain paraffins, rich in CH₄. The analysis results show that CH₄ selectivity is more than 90% and no olefin was detected in the products.
2. FT CO hydrogenation (third column) produces mainly long chain hydrocarbons (including paraffins and olefins) with CH₄ selectivity less than 10%.
3. For a feed which is a mixture of CO and CO₂ (fourth column) with only small quantities of CO (5.9%), the product analysis shows that selectivity of CH₄ is much lower compared to CO₂ hydrogenation only and the products also contain longer chain hydrocarbons in a considerable amount, although still lower compared to CO hydrogenation only.

Table 3. Catalytic performance of a Co-based catalyst: CO, CO₂ and (CO + CO₂) hydrogenation at 200 °C, 20 bar gauge and 60 mL (NTP)/(min·gcat).

| CO₂ Hydrogenation | CO Hydrogenation a | (CO+CO₂) Hydrogenation a |
|-------------------|--------------------|-------------------------|
| CO Conversion (%)| 14.6               | 100.0                   |
| CO₂ Conversion (%)| 24.2               | 13.0                    |
| CH₄ Selectivity (%)| 92.5               | 8.0                     |
| C₂+ Selectivity (%)| 7.5                | 82.0                    |
| C₂−₄ Selectivity (%)| 6.5                | 19.3                    |
| C₅+ Selectivity (%)| 1.0                | 72.1                    |
| O₂/P₂ b          | 0.0                | 0.24                    |
| O₃/P₃ c          | 0.0                | 1.7                     |
| O₄/P₄ d          | 0.0                | 1.1                     |

a: Data come from Ref. [15]. b: Olefin to Paraffin ratio with chain length 2. c: Olefin to Paraffin ratio with chain length 3. d: Olefin to Paraffin ratio with chain length 4.

The above results indicate that even a small amount of CO in the CO₂ rich feed changes the product selectivity significantly. This presents an opportunity for effective conversion of CO₂ by co-production of methanol and R-WGS. In this process, relatively long chain hydrocarbons can be achieved by combining the two catalytic reactors in series: a Cu-reactor for methanol production which also produces CO as by-product, followed by a Co-reactor fed with the tail gas from the Cu-reactor which contains a mixture of unreacted CO₂ and CO formed during methanol formation. The presence of the CO in the tail gas leads to improved selectivity of long chain hydrocarbons in the Co-reactor as shown in Table 3. This combination of reactors in series was tested in the lab as Case C and the results are presented below.

3.3. Case C: CO₂ Hydrogenation Using a Cu-Based Catalyst in Series with a Co-Based Catalyst

Figure 5 shows the comparison of the CO₂ conversion between Case A and Case C as a function of temperature, where temperature is varied in the Cu-reactor only while the Co-reactor temperature is
kept constant. It is clear that total CO$_2$ conversion increases with temperature in both cases and that the conversion in Case C is greater than that of Case A, due to the added reactor in Case C. The comparison of the CO selectivity for both Case A and Case B is shown in Figure 5B. No CO could be detected in the exit from the FT reactor when the temperature of the Cu-reactor was below 250 °C. It is worth noting that the CO selectivity in Case C is much lower than that of Case A, which indicates that the undesirable CO produced in the Cu-reactor was reacted in the Co-reactor (the second reactor in Case C) to valuable hydrocarbon products.

The overall product selectivity and formation rates as a function of temperature of the Cu-reactor are shown in Figure 6. Note that the temperature of Cu-reactor was gradually increased from 200 to 350 °C, while the Co-reactor was maintained at 200 °C. The results in Figure 6A show that CH$_4$ selectivity decreases with increasing operating temperature of the Cu-reactor, while the selectivity of the desired products (methanol + C$_2$-) increases. A similar trend is observed when looking at the reaction rate in Figure 6B where the rate of formation of desired products (methanol + C$_2$-) increases with increasing temperature of the Cu-reactor while the rate of formation of CH$_4$ decreases from $8.6 \times 10^{-5}$ mol/(min·gcat) to $7.4 \times 10^{-5}$ mol/(min·gcat), as the temperature of Cu-reactor increases from 200 to about 250 °C and then remains fairly constant as the temperature increases further.

Figure 6A also shows that the CH$_4$ selectivity in Case C is below 80% and decreases with increasing operating temperature of the Cu-reactor. The CH$_4$ selectivity is lower than that measured in Case B (92.5%); this is due to the presence of CO in the feed gas to the Co-reactor. It is known that the H$_2$/CO ratio in the reactor influences the product selectivity [21,22]: low H$_2$/CO ratio leads to low CH$_4$ selectivity and high long chain hydrocarbon selectivity. Therefore, the selectivity of CH$_4$ in the Co-reactor in Case C decreases as more CO is produced in the Cu-reactor thereby decreasing the H$_2$/CO ratio of the feed gas to the Co-reactor. Since the selectivity and product distribution are linked, it is therefore important to compare the product distribution between Case C and Case B.

Case C was a process in which two reactors were configured in series; a condensation trap, to condense the methanol and water produced from the first reactor (Figure 1: Case C), was placed between the Cu-reactor and the Co-reactor. In order to simplify the process, a modified process named “Modified Case C” was also tested by removing the condensate trap between the methanol (Cu catalyst) and the FT (Co-catalyst) reactors (see Figure 1: Modified Case C), so that all the products as well as the
unreacted feed gas from the Cu-reactor were fed to the Co-reactor. Figure 7 compares the performance of the reactor system for Case C and Modified Case C. Figure 7 shows that a slightly lower CO\textsubscript{2} conversions and higher CH\textsubscript{4} selectivity are obtained for the Modified Case C compared to that of Case C, which indicates that there is a negative effect on the FT reaction when methanol and water produced from the Cu-reactor were introduced to the Co-reactor.

Figure 6. Product selectivity (A) and rate of product formation (B) as a function of reaction temperature in the Cu-reactor during CO\textsubscript{2} hydrogenation in Case C: the first reactor was loaded with a Cu-based catalyst and operated at the following reaction conditions: pressure 20 bar (gauge), flow rate 60 mL(NTP)/(min·g\text{cat}) and feed gas H\textsubscript{2}/CO\textsubscript{2}/N\textsubscript{2} = 67.6%/22.6%/9.8%. The second reactor was loaded with a Co-based catalyst and operated at a reaction temperature of 200 °C.

Figure 7. Comparison of the CO\textsubscript{2} conversion and product selectivity as a function of the temperature of the Cu-reactor in Case C and Modified Case C: (1) Case C: CO\textsubscript{2} hydrogenation in a Cu-reactor and Co-reactor in series; a condensate trap is placed between the two reactors to condense the methanol and water produced from the Cu-reactor. (2) Modified Case C: CO\textsubscript{2} hydrogenation in a Cu-reactor and Co-reactor in series; there is no condensate trap between the two reactors. Reaction conditions: The first reactor was loaded with a Cu-based catalyst and operated under the following conditions: pressure 20 bar (gauge), flow rate 60 mL (NTP)/(min·g\text{cat}) of feed gas H\textsubscript{2}/CO\textsubscript{2}/N\textsubscript{2} = 67.6%/22.6%/9.8%; the second reactor was loaded with a Co-based catalyst and received the gas from the first reactor; the reaction temperature was set at 200 °C. (A) CO\textsubscript{2} conversion; (B) CO selectivity; (C) CH\textsubscript{4} selectivity; and (D) selectivity of (C\textsubscript{2+} + methanol).
There was concern that the Cu-based catalyst might deactivate during the experiments as the reaction temperature was varied over a wide range (200–350 °C) during the experiments in Cases A and C. An experimental run was repeated to test the Cu-based catalyst stability at 260 °C. Figure 8 clearly shows that after 1600 h running, there is no significant deactivation of the catalyst.

![Figure 8](image-url)

Figure 8. Cu-based catalyst stability testing. Reaction conditions: temperature 260 °C, pressure 20 bar (gauge), flow rate 60 mL (NTP)/(min·gcat) and feed gas H₂/CO₂/N₂ = 67.6%/22.6%/9.8%.

The Anderson–Schulz–Flory (ASF) distribution model has been used extensively and has been successful in describing and predicting the product distribution in FTS. If it is assumed that the hydrocarbon chain is formed by step-wise insertion or addition of CH₂ intermediates with constant growth probability (α), then the chain length distribution can be described by the ASF model [23]. If it is further assumed that α is independent of the hydrocarbon chain length, the ASF distribution is described by:

\[
\log\left(\frac{W_n}{n}\right) = n\log(\alpha) + \log\left(\frac{(1-\alpha)^2}{\alpha}\right),
\]

where \(W_n\) is the mass fraction of a hydrocarbon with chain length \(n\). Plotting \(\log\left(\frac{W_n}{n}\right)\) versus \(n\) gives a straight line whose slope (\(\log(\alpha)\)) is used to calculate the value of \(\alpha\). A higher \(\alpha\) value indicates that a heavier hydrocarbon product should be expected. Figure 9 shows the ASF product distributions for Case B and Case C and the results obtained from previous experiments in Yao et al. [15]. The following can be observed:

1. In Case B, with a single Co-reactor (H₂/CO₂ feed gas, Table 1), the results show that CH₄ is the dominant product with a small quantity of short chain hydrocarbons being formed. The results follow a typical ASF distribution with a low \(\alpha\) value of 0.41.
2. In Case C, with a Cu-reactor connected in series to a Co-reactor (H₂/CO₂ feed gas Table 1), two conditions are considered: Case C_1 and Case C_2, which consider the product distribution of the Co-reactor at two temperatures of the Cu-reactor, 300 °C and 350 °C, respectively. Figure 9 shows that the slope of the distribution of Case C_2 is slightly smaller than that of Case C_1 and thus the \(\alpha\) value of Case C_2 is slightly higher than that of Case C_1 (0.81 < 0.84). This is due to the higher CO concentration in the product from the Cu-reactor in Case C_2, due to the higher operating temperature. The CO concentration is higher at high temperatures because, as can be seen in Figure 2B, the CO selectivity in the Cu-reactor increases with temperature. By comparing the \(\alpha\) value in Case B with that of both cases in Case C, the results indicate that a better FT product distribution is obtained by coupling the Co-reactor to the Cu-reactor.
3. A mixture of H₂/CO in the feed (Table 3) [15] produces a considerably higher \(\alpha\) value (0.94) compared to Case B and Case C.

From these results, it can be concluded that the combined reactor in series (Case C) was found to greatly enhance CO₂ conversion and increase the long chain hydrocarbon product selectivity, although it still cannot achieve the same performance as CO hydrogenation.
Figure 9. Fischer–Tropsch synthesis (FTS) product distribution over a Co-based catalyst under the following reaction conditions: pressure 20 bar (gauge), flow rate 60 mL(NTP)/(min·g-cat): (1) Case B: FT CO\textsubscript{2} hydrogenation at a temperature of 200 °C and feed gas H\textsubscript{2}/CO\textsubscript{2}/N\textsubscript{2} = 67.6%/22.6%/9.8%; (2) Case C\textsubscript{1}: dual-reactor in series: the first reactor was loaded with a Cu-based catalyst operated at 300°C and feed gas H\textsubscript{2}/CO\textsubscript{2}/N\textsubscript{2} = 67.6%/22.6%/9.8%. The second reactor was loaded with a Co-based catalyst and operated at 200 °C; (3) Case C\textsubscript{2}: Dual-reactor in series: the first reactor was loaded with a Cu-based catalyst operated at 350 °C and feed gas H\textsubscript{2}/CO\textsubscript{2}/N\textsubscript{2} = 67.6%/22.6%/9.8%. The second reactor was loaded with a Co-based catalyst and operated at temperature of 200 °C; (4) H\textsubscript{2}/CO FTS [15]: FT CO hydrogenation at a temperature of 200 °C and feed gas H\textsubscript{2}/CO\textsubscript{2}/N\textsubscript{2} = 58.8%/30.3%/10.3% over the same Co-based catalyst as the one used in the present work.

4. Discussion and Implications

4.1. Comparison between the Current Work with the Results Reported in the Literature

Figures 5–7 and 9 shows that two reactors in series (a Cu-reactor followed by a Co-reactor) significantly increase the long chain hydrocarbon product selectivity compared to using only a Co-reactor. The catalytic activity and selectivity of this study has been compared to that reported in the literature (see Table 4). Table 4 lists CO\textsubscript{2} hydrogenation results under different reaction conditions over different catalysts, which shows: (1) generally, Cobalt catalysts promote the CO\textsubscript{2} hydrogenation reaction to form methane (>80%); however, He et al. [24] reported a Co\textsubscript{6}/MnO\textsubscript{x} nano-catalyst which could efficiently catalyze CO\textsubscript{2} hydrogenation to form long chain hydrocarbons (53.2% in Table 4); they ran the experiments in batch mode with a liquid solvent; (2) iron catalysts can catalyze the CO\textsubscript{2} hydrogenation and have a low CH\textsubscript{4} selectivity, while one of the drawback is the high selectivity to CO. Table 4 indicates that both Co and Fe catalyst have the potential to catalyze CO\textsubscript{2} hydrogenation to liquid fuels by either combining two catalysts (current work), preparing a bifunctional catalyst [25] or selecting a suitable support [14,26].

In addition, our experimental results (Table 3 and Figures 5–7) suggest a stepwise conversion of CO and CO\textsubscript{2} in the FT reactor. Firstly, most of the CO is converted to hydrocarbons via the FT reaction, then CO\textsubscript{2} is hydrogenated to short chain hydrocarbons, mainly methane. Therefore, further experiments with different ratios of Cu to Co-based catalysts are highly recommended. In addition, more attention needs to be paid to the effect of total pressure on the catalyst performance, as both methanol yield and long chain hydrocarbon selectivity are favored at higher pressure due to the stoichiometry of both the methanation and the FT reaction. Furthermore, a bifunctional catalyst may be another option to convert CO\textsubscript{2} to liquid fuels.
Table 4. Comparison between the present work with the results reported in the literature.

| Reactor | Catalyst | CO₂ Conv (%) | CH₄ Sel (%) | C₅₊ + CH₃OH Sel (%) | CO Sel (%) | Reaction Conditions | Ref |
|---------|----------|--------------|-------------|----------------------|------------|---------------------|-----|
| One reactor | Co/TiO₂ | 24.2 | 92.5 | 1.0 | 0.0 | H₂/CO₂ = 3, 200 °C, 60 ml(NTP)/(min·gcat), 20 bar | Current work |
| Two reactors in series | Cu-200_Co-200 | 23.2 | 76.4 | 17.6 | 0.0 | Reactor one: Cu catalyst; 200–350 °C; 60 ml(NTP)/(min·gcat), 20 bar; reactor two: Co/TiO₂, 200 °C | Current work |
| | Cu-250_Co-200 | 25.6 | 62.9 | 28.6 | 2.2 | | |
| | Cu-300_Co-200 | 30.6 | 50.0 | 33.0 | 6.3 | | |
| | Cu-350_Co-200 | 35.6 | 43.0 | 44.8 | 9.8 | | |
| One reactor | Co/MnOₓ | 15.3 | 46.4 † | 53.2 | 0.4 | H₂/CO₂ = 1, 200 °C, 8 bar, no flow (batch mode), solvent: squalane | [24] |
| | Co/ZnOₓ | / | 80.7 † | 19.2 | 0.1 | | |
| | Coₓ/CoOₓ | / | 89.8 † | 9.8 | 0.0 | | |
| | Coₓ/AlOₓ | / | 94.2 † | 5.7 | 0.0 | | |
| One reactor | Co/MnOₓ/SiO₂/Pt | 18.0 | 95.0 | 1.5 | 33.2 | H₂/CO₂ = 2, 190 °C, 30(ml(NTP)/(min·gcat),10 bar | [14] |
| | Fe/TiO₂ | 11.5 | 33.3 | 4.4 | 35.7 | | |
| | Fe/Al₂O₃ | 22.8 | 38.3 | 7.8 | 11.4 | H₂/CO₂ = 3, 300 °C, 31.6 (ml(NTP)/(min·gcat),10 bar | |
| | Fe/SiO₂ | 6.9 | 23.4 | 0.1 | 71.0 | | |
| | Feₓ/AlOₓ | 30.4 | 7.6 | 23.5 | 40.5 | | |
| One reactor | Fe/Cu/K | 10.8 | 9.1 | 26.9 | 39.3 | H₂/CO₂ = 3, 300 °C, 8 bar, 60ml/(min·gcat), 10-20 bar | [26] |
| | Fe/Cu/Al/K | 11.3 | 8.5 | 27.1 | 45 | | |
| | Fe/Cu/Si/K | 10.2 | 21.1 | 8.8 | 43.4 | | |
| | Fe/Cu/Al/K (2) | 15.6 | 9.9 | 39.4 | 22.8 | | |
| One reactor | Fe/FeOₓ | 14.3 | 40.2 | 1.5 | 33.2 | H₂/CO₂ = 3, 300 °C, 60ml/(min·gcat), 10 bar | [25] |
| | Cu/FeOₓ-6 | 17.3 | 1.8 | 45.3 | 31.7 | | |
| | Cuₓ/Fe₂O₃ | 15.7 | 41.0 | 1.8 | 28.9 | | |

* C₅₊ selectivity (%).
4.2. Multi-Reactor System

A number of studies are available where the direct hydrogenation of CO\(_2\) and CO/CO\(_2\) mixtures to hydrocarbons are reported using traditional FTS catalysts at FT reaction conditions [27]. At present there are two methods of operation favoured for commercial FTS processes [28–30]. The first is the high-temperature (300–350 °C) process, which employs iron-based (Fe-based) catalysts, for the production of gasoline and linear low molecular mass olefins. The second is the low-temperature (180–250 °C) process over either iron or Co catalysts, which produces high molecular weight linear waxes, which in turn can be hydro-cracked to produce diesel of high quality. Iron based catalysts are active for both WGS (R-WGS) and FT reactions [31], and are commercially used for high temperature FT (HTFT) [32]. For HTFT (300–350 °C), the water-gas shift (WGS) reaction is typically sufficiently rapid so that it nearly reaches equilibrium [12]. HTFT synthetic crude (syncrude) typically consists of more highly olefinic intermediate hydrocarbons (predominantly \(\alpha\)-olefins) and is rich in oxygenates, such as alcohols, aldehydes, ketones and acids [33]. Olefins and oxygenates are therefore extracted for the production of chemicals [34]. In principle, iron catalysts would be ideal for use in FTS with CO\(_2\)-containing syngas feeds [14–35]. Currently, the production of light alkanes [36] and alkenes [37,38] from CO\(_2\) is fundamentally a modification of HTFT, where CO\(_2\) is used instead of CO.

CO\(_2\) hydrogenation over an Fe-catalyst is active for both HTFT and WGS reactions. Thus, the introduction of an iron catalyst in the process can help overcome the challenges of co-producing methanol and hydrocarbon fuels when using the Cu and Co catalysts only.

The Fe-catalyst reactor can be introduced as an intermediate between the Cu-reactor and the Co-reactor (Figure 10):

![Figure 10. A new simplified flow scheme for hydrogenation of CO\(_2\) to fuels and chemicals (HC stands for hydrocarbons).](image)

1. In the first reactor, as in Case C, a Cu-based catalyst is used to convert CO\(_2\) to methanol with CO as the byproduct. In this case methanol production is favored by setting reactions conditions to low temperature and high pressure. The aqueous product (a mixture of methanol and H\(_2\)O) is removed from the produced stream, and the tail gas (a mixture of H\(_2\)/CO/CO\(_2\) with a small amount of CH\(_4\)) is introduced into the Fe-catalyst reactor.

2. HTFT occurs over an Fe-based catalyst, where the tail gas from the first reactor (H\(_2\)/CO/CO\(_2\) mixture) is converted to short olefins and oxygenates. Since the feed stream does not contain water, the equilibrium limitation of the R-WGS reaction occurring in the first reactor is eliminated in the second reactor and thus more CO\(_2\) is converted to CO, and because the iron catalyst is WGS active, a new equilibrium can be rapidly reached. Part of the CO formed is consumed in FT reactions.
within the reactor to produce light and medium olefinic hydrocarbons with small amounts of oxygenates, and the remaining CO, goes out in the tail gas (after removal of condensable FT products and water). CO₂ may also react with H₂ to produce FT products on the iron catalyst depending on the amount of CO undergoing FT reaction.

3. The third reactor is a low temperature FT reactor over a Co-based catalyst. This reactor receives a CO rich tail gas mixture (H₂/CO/CO₂) from the second reactor, thus allowing a high selectivity for heavier hydrocarbons relative to lighter ones as discussed previously. Furthermore, if CO is an undesired product, this reactor will reduce the CO to low levels by converting it to FT products.

We therefore see that the introduction of an Fe-based catalyst intermediate reactor may greatly diversify the products from the process. CO₂ conversion to liquid products will be improved and methane selectivity will be reduced.

A possible improvement of the process that would make it more economical, would be to combine Reactor 1 and Reactor 2 into a single reactor. This can be done by mixing the Cu and Fe catalysts at an appropriate ratio, run the reaction at a relatively higher temperature (such as 300 °C) and high pressure, to enable sufficient methanol selectivity and iron catalyst activity. Such a process would enable heat and work integration directly between WGS, methanol and FT synthesis in a single reactor.

4.3. A Proposed “Greener” Process for the Conversion of CO₂ to Valuable Products

Concerns over global warming have led to the search for ways of safe CO₂ disposal; CO₂ hydrogenation could be one of the solutions as it would act as a CO₂ re-use and thereby reduce net CO₂ production [14]. However this requires an environmental friendly source of H₂, which will not produce more CO₂, such as hydro, solar, or nuclear energy [39–41]. If we assume that these sources of hydrogen are available, a greener process can be proposed as shown in Figure 11, in which the methanol synthesis and FTS are combined to convert CO₂ to chemicals and hydrocarbons. The proposed process is much more flexible and controls the yields of methanol and hydrocarbon products by changing the operating conditions. The water produced in each section of the process can be recycled to produce H₂ to be used in the process (via electrolysis using renewable sources of energy suggested above). The diesel fuels and methanol produced from this process could be blended together with different proportions, which will reduce the greenhouse emission with zero aromatic, nitrogen and sulfur components.

![Figure 11. Schematic diagram of the conversion of CO₂ to liquid fuel and chemicals by both methanol synthesis and FTS.](image)

Furthermore CO₂ hydrogenation could also be a way of storing hydrogen in more easily transportable and safer compounds (methanol and hydrocarbons) compared to hydrogen itself, at the same time using CO₂ as green source of carbon compared to conventional sources [14,17,41]. The other advantage for using CO₂ as a source of carbon as opposed to CO, is that CO₂ hydrogenation is less exothermic and therefore the lower heat duty will result in reduced capital and operating costs.

5. Conclusions

Two reactors configured in series were used for the hydrogenation of CO₂ to methanol and hydrocarbon fuels by combining methanol synthesis, R-WGS and subsequently FTS. First, the experiments for methanol
synthesis and hydrocarbon fuels were conducted in separate reactors. The reactors were then coupled in series in the second set of experiments. Experimental results for CO$_2$ hydrogenation over the Cu-catalyst, showed that both methanol synthesis and R-WGS reactions reach equilibrium and the conversion of CO$_2$ per pass is relatively low. Experimental results of CO$_2$ hydrogenation over the Co-catalyst, showed a high CH$_4$ selectivity (92.5%) and also the production of short chain paraffins with a typical low $\alpha$ ASF distribution. It has been previously shown that co-feeding even small amounts of CO with a H$_2$/CO$_2$ feed mixture can significantly change the FT product selectivity [15]. This was effectively achieved in the second set of experiments where the two reactors were coupled in series; the tail gas from the Cu-reactor, containing unreacted feed gas and CO was fed into the Co-reactor. The results showed a significant increase in the long chain hydrocarbon selectivity and an increase in the overall CO$_2$ conversion. The two reactors in series enhanced the CO$_2$ conversion and increased the liquid product selectivity but did not achieve the same performance as the normal FT CO hydrogenation.

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**References**

1. Park, S.-E.; Nam, S.S.; Choi, M.J.; Lee, K.W. Catalytic Reduction of Carbon Dioxide. The Effects of Catalysts and Reductants. *Energy Convers. Manag.* 1995, 36, 573–576. [CrossRef]
2. Yao, L.; Shen, X.; Pan, Y.; Peng, Z. Synergy between Active Sites of Cu-In-Zr-O Catalyst in CO$_2$ Hydrogenation to Methanol. *J. Catal.* 2019, 372, 74–85. [CrossRef]
3. Blumberg, T.; Morosuk, T.; Tsatsaronis, G. CO$_2$-Utilization in the Synthesis of Methanol: Potential Analysis and Exergetic Assessment. *Energy* 2019, 175, 730–744. [CrossRef]
4. Joo, O.-S.; Jung, K.-D.; Moon, I.; Rozovskii, A.Y.; Lin, G.I.; Han, S.-H.; Uhm, S.-J. Carbon Dioxide Hydrogenation to Form Methanol via a Reverse-Water-Gas-Shift Reaction (the CAMERE Process). *Ind. Eng. Chem. Res.* 1999, 38, 1808–1812. [CrossRef]
5. Zhang, Y.; Fei, J.; Yu, Y.; Zheng, X. Study of CO$_2$ Hydrogenation to Methanol over Cu-V/y-Al$_2$O$_3$ Catalyst. *J. Nat. Gas Chem.* 2007, 16, 12–15. [CrossRef]
6. Bahruji, H.; Bowker, M.; Hutchings, G.; Dimitratos, N.; Wells, P.; Gibson, E.; Jones, W.; Brookes, C.; Morgan, D.; Lavey, G. Pd/ZnO Catalysts for Direct CO$_2$ Hydrogenation to Methanol. *J. Catal.* 2016, 343, 133–146. [CrossRef]
7. Li, S.; Guo, L.; Ishihara, T.S.C. Hydrogenation of CO$_2$ to Methanol over Cu/AlCeO Catalyst, *Catal. Today* 2020, 339, 352–361. [CrossRef]
8. Lonis, F.; Tola, V.; Cau, G. Renewable Methanol Production and Use through Reversible Solid Oxide Cells and Recycled CO2 Hydrogenation. *Fuel* 2019, 246, 500–515. [CrossRef]
9. Lange, J.-P. Methanol Synthesis: A Short Review of Technology Improvements. *Catal. Today* 2001, 64, 3–8. [CrossRef]
10. Pérez-Forrestes, M.; Schöneberger, J.C.; Boulamanti, A.; Tzimas, E. Methanol Synthesis Using Captured CO$_2$ as Raw Material: Techno-Economic and Environmental Assessment. *Appl. Energy* 2015, 161, 718–732. [CrossRef]
11. Farsi, M.; Jahanmiri, A. Methanol Production in an Optimized Dual-Membrane Fixed-Bed Reactor. *Chem. Eng. Process. Process Intensif.* 2011, 50, 1177–1185. [CrossRef]
12. Zhang, Y.; Jacobs, G.; Sparks, D.E.; Dry, M.E.; Davis, B.H. CO and CO$_2$ Hydrogenation Study on Supported Cobalt Fischer-Tropsch Synthesis Catalysts. *Catal. Today* 2002, 71, 411–418. [CrossRef]
13. Visconti, C.G.; Lietti, L.; Tronconi, E.; Forzatti, P.; Zennaro, R.; Finocchio, E. Fischer-Tropsch Synthesis on a Co/Al$_2$O$_3$ Catalyst with CO$_2$ Containing Syngas. *Appl. Catal. A Gen.* 2009, 355, 61–68. [CrossRef]
14. Riedel, T.; Claeyts, M.; Schulz, H.; Schaub, G.; Nam, S.-S.; Jun, K.-W.; Choi, M.-J.; Kishan, G.; Lee, K.-W. Comparative Study of Fischer–Tropsch Synthesis with H2/CO and H2/CO2 Syngas Using Fe- and Co-Based Catalysts. *Appl. Catal. A Gen.* 1999, 186, 201–213. [CrossRef]

15. Yao, Y.; Hildebrandt, D.; Glasser, D.; Liu, X. Fischer-Tropsch Synthesis Using H2/CO/CO2 Syngas Mixtures over a Cobalt Catalyst. *Ind. Eng. Chem. Res.* 2010, 49, 11061–11066. [CrossRef]

16. Shafer, W.D.; Jacobs, G.; Graham, U.M.; Hamdeh, H.H.; Davis, B.H. Increased CO2 Hydrogenation to Liquid Products Using Promoted Iron Catalysts. *J. Catal.* 2019, 369, 239–248. [CrossRef]

17. Sempuga, B.C.; Yao, Y. CO2 Hydrogenation from a Process Synthesis Perspective: Setting up Process Targets. *J. CO2 Util.* 2017, 20, 34–42. [CrossRef]

18. Datta, A.; Dutta, S.; Mandal, B.K. Effect of methanol addition to diesel on the performance and emission characteristics of a CI engine. *J. Phys. Conf. Ser.* 2014, 1, 8–13. [CrossRef]

19. Zhu, L.; Cheung, C.S.; Zhang, W.G.; Huang, Z.; Zhu, L.; Cheung, C.S.; Zhang, W.G.; Huang, Z. Influence of methanol—Biodiesel blends on the particulate emissions of a direct injection diesel engine influence of methanol—Biodiesel blends on the particulate emissions of a direct injection diesel engine. *Aerosol Sci. Technol.* 2010, 44, 362–369. [CrossRef]

20. Daza, Y.A.; Kuhn, J.N. CO2 conversion by reverse water gas shift catalysis: Comparison of catalysts, mechanisms and their consequences for CO2 conversion to liquid fuels. *RSC Adv.* 2011, 6, 49675–49691. [CrossRef]

21. De la Osa, A.R.; De Lucas, A.; Romero, A.; Valverde, J.L.; Sánchez, P. Fischer-Tropsch Diesel Production over Calcium-Promoted Co/Alumina Catalyst: Effect of Reaction Conditions. *Fuel* 2011, 90, 1935–1945. [CrossRef]

22. Zhou, W.; Chen, J.G.; Fang, K.G.; Sun, Y.H. The Deactivation of Co/SiO2 Catalyst for Fischer-Tropsch Synthesis at Different Ratios of H2 to CO. *Fuel Process. Technol.* 2006, 87, 609–616. [CrossRef]

23. Tavakoli, A.; Sohrabi, M.; Kargari, A. Application of Anderson—Schulz—Flory (ASF) Equation in the Product Distribution of Slurry Phase FT Synthesis with Nanosized Iron Catalysts. *Chem. Eng. J.* 2008, 136, 358–363. [CrossRef]

24. He, Z.; Cui, M.; Qian, Q.; Zhang, J.; Liu, H.; Han, B. Synthesis of liquid fuel via direct hydrogenation of CO2. *PNAS* 2019, 116, 12654–12659. [CrossRef]

25. Choi, Y.H.; Jang, Y.J.; Park, H.; Kim, W.Y.; Lee, Y.H.; Choi, S.H.; Lee, J.S. Carbon dioxide Fischer-Tropsch synthesis: A new path to carbon-neutral fuels. *Appl. Catal. B* 2017, 202, 605–610. [CrossRef]

26. Prasad, P.S.S.; Bae, J.W.; Jun, K.W.; Lee, K.W. Fischer-Tropsch synthesis by carbon dioxide hydrogenation on Fe-based catalyst. *Catal. Surv. Asia* 2008, 12, 170–183. [CrossRef]

27. Martinelli, M.; Visconti, C.G.; Lietti, L.; Forzatti, P.; Bassano, C.; Deiana, P. CO2 Reactivity on Fe–Zn–Cu–K Fischer–Tropsch Synthesis Catalysts with Different K-Loadings. *Catal. Today* 2014, 228, 77–88. [CrossRef]

28. Dry, M.E. The Fischer-Tropsch Process: 1950–2000. *Catal. Today* 2002, 71, 227–241. [CrossRef]

29. Gorimbo, J.; Muleja, A.; Lu, X.; Yao, Y.; Liu, X.; Hildebrandt, D.; Glasser, D.; Lu Plot and Yao Plot: Models to Analyze Product Distribution of Long-Term Gas-Phase Fischer-Tropsch Synthesis Experimental Data on an Iron Catalyst. *Energy Fuels* 2017, 31, 5682–5690. [CrossRef]

30. Wu, Y.; Feng, J.; Li, W. System Development of Integrated High Temperature and Low Temperature Fischer-Tropsch Synthesis for High Value Chemicals. *Chem. Eng. Res. Des.* 2017, 1–12. [CrossRef]

31. Van der Laan, G.P.; Beenackers, A.A.C.M. Intrinsic Kinetics of the Gas-solid Fischer-Tropsch and Water Gas Shift Reactions over a Precipitated Iron Catalyst. *Appl. Catal. A Gen.* 2000, 193, 39–53. [CrossRef]

32. Steynberg, A.P.; Espinosa, R.L.; Jager, B.; Vosloo, A.C. High Temperature Fischer–Tropsch Synthesis in Commercial Practice. *Appl. Catal. A Gen.* 1999, 186, 41–54. [CrossRef]

33. Huang, Y.; Yi, Q.; Wei, G.; Kang, J.; Li, W.; Feng, J. Energy Use, Greenhouse Gases Emission and Cost Efficiency of an Integrated High- and Low- Temperature Fisher-Tropsch Synthesis Plant From. *Appl. Energy* 2018, 228, 1009–1019. [CrossRef]

34. Leckel, D. Hydroprocessing Euro 4-Type Diesel from High-Temperature Fischer-Tropsch Vacuum Gas Oils. *Energy Fuels* 2009, 23, 38–45. [CrossRef]

35. Dry, M.E. Practical and Theoretical Aspects of the Catalytic Fischer-Tropsch Process. *Appl. Catal. A Gen.* 1996, 138, 319–344. [CrossRef]

36. Saeidi, S.; Amin, N.A.S.; Rahimpour, M.R. Hydrogenation of CO2 to Value-Added Products—A Review and Potential Future Developments. *J. CO2 Util.* 2014, 5, 66–81. [CrossRef]
37. Satthawong, R.; Koizumi, N.; Song, C.; Prasassarakich, P. Light Olefin Synthesis from CO₂ Hydrogenation over K-Promoted Fe-Co Bimetallic Catalysts. *Catal. Today* 2015, 251, 34–40. [CrossRef]

38. Zhang, J.; Lu, S.; Su, X.; Fan, S.; Ma, Q.; Zhao, T. Selective Formation of Light Olefins from CO₂ Hydrogenation over Fe–Zn–K Catalysts. *J. CO₂ Util.* 2015, 12, 95–100. [CrossRef]

39. Hildebrandt, D.; Glasser, D.; Hausberger, B.; Patel, B.; Glasser, B.J. Chemistry. Producing Transportation Fuels with Less Work. *Science* 2009, 323, 1680–1681. [CrossRef]

40. Milani, D.; Khalilpour, R.; Zahedi, G.; Abbas, A. A Model-Based Analysis of CO₂ Utilization in Methanol Synthesis Plant. *J. CO₂ Util.* 2015, 10, 12–22. [CrossRef]

41. Prieto, G. Carbon Dioxide Hydrogenation into Higher Hydrocarbons and Oxygenates: Thermodynamic and Kinetic Bounds and Progress with Heterogeneous and Homogeneous Catalysis. *ChemSusChem* 2017, 10, 1–16. [CrossRef] [PubMed]

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