Combination of Cu/ZnO Methanol Synthesis Catalysts and ZSM-5 Zeolites to Produce Oxygenates from CO₂ and H₂

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Abstract
Cu/ZnO methanol catalysts were deposited over several ZSM-5 acid zeolites to directly synthesise oxygenates (methanol and dimethyl ether) from a CO₂/H₂ feed. Catalysts were prepared by two different preparation methodologies: chemical vapour impregnation (CZZ-CVI) and oxalate gel precipitation (CZZ-OG). Chemical vapour impregnation led to Cu/ZnO being deposited on the zeolite surface, whilst oxalate gel precipitation led to the formation of Cu/ZnO agglomerates. For both sets of catalysts a higher concentration of mild and strong acid sites were produced, compared to the parent ZSM-5 zeolites, and CZZ-CVI had a higher concentration of acid sites compared to CZZ-OG. Nevertheless, CZZ-OG shows considerably higher oxygenate productivity, 1322 mmol Kg⁻¹ h⁻¹, compared to 192 mmol Kg⁻¹ h⁻¹ over CZZ-CVI (ZSM-5(50), 250 °C, 20 bar, CO₂/H₂ = 1/3, 30 ml min⁻¹), which could be assigned to a combination of smaller particle size and enhanced methanol mass transfer within the zeolites.

Keywords CO₂ · Cu · ZnO · Dimethyl ether · Hydrogenation · Methanol · ZSM-5

1 Introduction
CO₂ hydrogenation allows us to store hydrogen produced from renewable resources (green hydrogen) in the form of liquid fuels [1]. Thus, green hydrogen [2] can be readily incorporated into the current energy infrastructure, while prominent technologies (e.g. PEM electrolysers) which use hydrogen as an energy vector are well-established. Among the possible CO₂ hydrogenation products [3], CH₃OH is of great interest [4]. CH₃OH can be blended with gasoline or converted into gasoline. For instance, in 1985 to meet the domestic fuel demand in New Zealand two methanol plants were combined with the methanol-to-gasoline process developed by Mobil [5]. Moreover, CH₃OH is also used as an intermediate for the production of numerous important chemicals (e.g. ethylene, propylene) [6]. CH₃OH is currently produced over Cu/ZnO/Al₂O₃ catalysts from syngas (CO + H₂ + CO₂) at 200–300 °C and 80–120 bar [5]. When CO₂ (Eq. 1) is the feed instead of CO (Eq. 2) the CH₃OH yield is constrained by equilibrium. For instance, at 250 °C and 30 bar, CH₃OH yield is limited to ca. 12 C-mol %, while CH₃OH yield increase to ca. 42 C-mol % using a CO/CO₂ = 4 feed (H₂/(CO + CO₂) = 3) [7]. Under laboratory testing conditions (20–50 bar) the CO₂ hydrogenation to CH₃OH is limited to the 200–270 °C range, at lower temperature little CO₂ conversion is observed, while at higher CH₃OH productivity is limited by equilibrium whilst the reverse water gas shift (RWGS) reaction is promoted (Eq. 3) [8].

\[ \text{CO}_2 + 3 \text{H}_2 \rightleftharpoons \text{CH}_3\text{OH} + \text{H}_2\text{O} \quad \Delta H^\circ = -49.5 \text{ kJ mol}^{-1} \] (1)

\[ \text{CO} + 2 \text{H}_2 \rightleftharpoons \text{CH}_3\text{OH} \quad \Delta H^\circ = -90.6 \text{ kJ mol}^{-1} \] (2)

\[ \text{CO}_2 + \text{H}_2 \rightleftharpoons \text{CO} + \text{H}_2\text{O} \quad \Delta H^\circ = 41.2 \text{ kJ mol}^{-1} \] (3)
CH$_3$OH productivity from CO$_2$ can be enhanced by simultaneously dehydrating CH$_3$OH to dimethyl ether (DME) (Eq. 4) by combining a CH$_3$OH synthesis catalyst with a solid acid catalyst (overall reaction in Eq. 5), known as hybrid catalysts. Like CH$_3$OH, DME synthesis from CO$_2$ is favoured at low temperature and high pressure. DME synthesis from CO$_2$ is advantageous compared to CH$_3$OH because of the higher DME equilibrium selectivity. Moreover, DME can be employed as a fuel in diesel engines because of its higher cetane number and lower auto-ignition temperature compared to diesel fuel (55–60 compared to 40–50, and 235 °C compared to 250 °C, respectively), and can be easily liquefied at low pressure, hence it can be used as a liquefied petroleum gas (LPG) substitute [9, 10].

\[
2 \text{CH}_3\text{OH} \rightleftharpoons (\text{CH}_3)_2\text{O} + \text{H}_2\text{O} \quad \Delta H^\theta = -24.0 \text{ kJ mol}^{-1} \\
2 \text{CO}_2 + 6 \text{H}_2 \rightleftharpoons (\text{CH}_3)_2\text{O} + 3 \text{H}_2\text{O} \quad \Delta H^\theta = -123.0 \text{ kJ mol}^{-1}
\]

Based on the actual CH$_3$OH synthesis process (developed by Imperial Chemical Industries), most of the research on the CO$_2$ hydrogenation to CH$_3$OH focus on Cu/ZnO catalysts [11]. As well summarised by Spencer [12], ZnO prevents the sintering of Cu particles by acting as a refractory spatial binder, maintaining small Cu crystallites from agglomerating during reaction [13]. Nevertheless, ZnO not only plays a spatial role in improving the catalyst lifespan, but it is also an activity promoter. As Spencer elucidated: on one hand, under CH$_3$OH synthesis conditions surface CuZn alloys can form, leading to the formation of Cu$^{++}$–O–Zn active sites [14–16]. On the other hand, under laboratory testing conditions (moderate pressure, CO$_2$/H$_2$ = 1/3, high conversion) H$_2$ dissociates on ZnO and spills over onto Cu particles [17]. For the CH$_3$OH dehydration to dimethyl ether (DME) ZSM-5 zeolites are preferred to γ-alumina as the acid catalyst because ZSM-5 zeolites contain Lewis and Brønsted acid sites, their high resistance to water, and their chemical and thermal stability [11]. As for instance, Aguayo et al. [18] compared Cu–ZnO–Al$_2$O$_3$/γ-Al$_2$O$_3$ and Cu–ZnO–Al$_2$O$_3$/NaZSM-5 hybrid catalysts for CH$_3$OH synthesis from the hydrogenation of CO and CO$_2$. It was observed that competitive adsorption between H$_2$O and CH$_3$OH for acid sites on γ-Al$_2$O$_3$, leading to lower CO/CO$_2$ conversion and DME selectivity compared to ZSM-5. Moreover, catalyst deactivation after regeneration was observed for the γ-Al$_2$O$_3$-containing hybrid catalyst, whilst good regeneration was observed for the Cu–ZnO–Al$_2$O$_3$/ZSM-5 hybrid catalyst. Similar conclusions were reported by Naik et al. [19] Bonura and co-workers studied the effect of water on CH$_3$OH dehydration activity over ZSM-5 zeolites. At low temperature H$_2$O competes with acid sites resulting in a decrease in CH$_3$OH conversion, with higher deactivation observed with increasing the zeolite acidity. Whilst, no appreciable decrease in CH$_3$OH dehydration was observed above 180 °C [20]. Readers are referred to reports by Catizzone et al. [21] and Álvarez et al. [11] for a detailed review on the CO$_2$ upgrading to DME.

Frusteri et al. [20] compared the activity of CuZnZr/ZSM-5 hybrid catalysts with CuZnZr supported over ZSM-5 by co-precipitation. Catalysts where the CH$_3$OH synthesis catalyst is supported over the solid acid catalyst are known as integrated catalysts. Slightly higher CO$_2$ conversion and DME productivity was observed over the integrated catalyst, which was attributed to the spatial proximity between the two active functionalities. Due to the benefits of integrated catalysts over hybrid catalysts, the aim of this work is to compare DME productivity between CuZnO/ZSM-5 integrated catalysts prepared via chemical vapour impregnation (CVI) or via oxalate gel precipitation (OG).

## 2 Experimental Section

### 2.1 Materials

All materials employed in this work were purchased from chemical suppliers and used as received without prior purification. ZSM-5 zeolites in the (NH$_4^+$) form were purchased from Alfa Aesar (Si/Al = 23, 50, 80); Zn acetylacetonate (Zn(acac)$_2$, 99%); Cu acetylacetonate (Cu(acac)$_2$, 97%); Cu nitrate hydrate (Cu(NO$_3$)$_2$·3H$_2$O, 99%) and Zn nitrate hydrate (Zn(NO$_3$)$_2$·6H$_2$O, 99%) and ethanol (99.8%) were purchased from Sigma-Aldrich.

### 2.2 Catalysts Synthesis

CuZnO/ZSM-5 catalysts synthesised via Chemical Vapour Impregnation (CZZ-CVI). For the preparation of 3.0 g of CuZnO/ZSM-5 catalyst by CVI (20 wt.%, Cu/Zn mol = 1) 1.24 g of Cu(acac)$_2$, 1.21 g Zn(acac)$_2$ and 2.4 g of activated H-ZSM-5 (static air, 550 °C, 5 °C min$^{-1}$, 4 h) were thoroughly mixed in a glass vial. The homogeneous mixture was then transferred into a Schlenk flask, evacuated (10$^{-3}$ mbar) and heated to 120 °C for 1 h. Samples were then cooled to 30 °C, recovered and annealed in static air (500 °C, 2 °C min$^{-1}$, 16 h).

CuZnO/ZSM-5 catalysts synthesised via Oxalate Gel precipitation (CZZ-OG). The procedure for the preparation of 2.3 g of CuZnO/ZSM-5 catalyst by OG (20 wt.%, Cu/Zn mol = 1) catalysts by OG is as follows: 1.14 g of Cu(NO$_3$)$_2$ and 1.37 g of Zn(NO$_3$)$_2$ were dissolved in 200 ml of ethanol, under vigorous stirring 2.4 g of activated H-ZSM-5 (static air, 550 °C, 5 °C min$^{-1}$, 4 h) were added. 2.16 g of oxalic acid were added to the slurry. The precipitate was aged for 1 h under
stirring, recovered by filtration, dried (static air, 100 °C, 16 h) and annealed (static air, 500 °C, 10 °C min⁻¹, 2 h).

### 2.3 Catalyst Characterisation

Prior to characterisation a portion of the CZZ catalysts were reduced under flowing 5% H₂/Ar (220 °C, 2 °C min⁻¹, 1 h). Powder X-ray diffraction (XRD) patterns were recorded on a (θ-θ) PANalyticalX'pert Pro powder diffractometer fitted with a position sensitive detector using a Cu Kα radiation source (40 keV, 40 mA). Transmission electron microscopy (TEM) images were obtained on a JEOL 2100 (LaB6) instrument fitted with a Gatan Ultrascan 1000xp digital camera operated at 200 kV. Specimens were dry-prepared on 300 mesh copper TEM-grids coated with holey carbon film. Pore size and apparent BET surface area were obtained from N₂ adsorption isotherms at (−196 °C) on a Quantachrome Nova 2200e instrument. Prior to analysis samples were degassed in situ (120 °C, 10 h). Ammonia temperature programmed desorption (NH₃-TPD) were acquired on a Quantachrome ChemBET TPR/TPD equipped with a TCD detector. 150 mg of pre-reduced catalyst were secured with quartz wool. Prior to analysis samples were heated under a He flow (130–140 ml min⁻¹) at 400 °C (10 °C min⁻¹, 1 h). The sample cell was cooled to 30 °C, the flow was then switched to 10% NH₃/Ar. After 15 min the gas was switched back to He, heated to 100 °C (10 °C min⁻¹, 1 h) to remove physiosorbed NH₃, and cooled to 50 °C. NH₃-TPD profiles were then measured (50–700 °C, 10 °C min⁻¹).

### 2.4 Catalyst Performance for Oxygenates Synthesis from CO₂ and H₂

The CO₂ hydrogenation and consecutive CH₃OH dehydrogenation to DME was assessed in a stainless steel fixed-bed continuous flow reactor (50 cm length, 0.5 cm internal diameter). Temperature was controlled through a chromel–alumel thermocouple placed inside the catalyst bed, pressure was controlled using a back pressure regulator. Catalysts were pressed (10 ton, 1 min) into a 1.3 cm die, the die was then crushed, and sieved into (425–600 µm) pellets. 0.5 g of catalyst pellets were placed in the reactor tube without diluent, crushed, and sieved into (425–600 µm) pellets. 0.5 g of catalyst was placed in a (0-0) PANalyticalX'pert Pro powder diffractometer fitted with a position sensitive detector using a Cu Kα radiation source (40 keV, 40 mA). Transmission electron microscopy (TEM) images were obtained on a JEOL 2100 (LaB6) instrument fitted with a Gatan Ultrascan 1000xp digital camera operated at 200 kV. Specimens were dry-prepared on 300 mesh copper TEM-grids coated with holey carbon film. Pore size and apparent BET surface area were obtained from N₂ adsorption isotherms at (−196 °C) on a Quantachrome Nova 2200e instrument. Prior to analysis samples were degassed in situ (120 °C, 10 h). Ammonia temperature programmed desorption (NH₃-TPD) were acquired on a Quantachrome ChemBET TPR/TPD equipped with a TCD detector. 150 mg of pre-reduced catalyst were secured with quartz wool. Prior to analysis samples were heated under a He flow (130–140 ml min⁻¹) at 400 °C (10 °C min⁻¹, 1 h). The sample cell was cooled to 30 °C, the flow was then switched to 10% NH₃/Ar. After 15 min the gas was switched back to He, heated to 100 °C (10 °C min⁻¹, 1 h) to remove physiosorbed NH₃, and cooled to 50 °C. NH₃-TPD profiles were then measured (50–700 °C, 10 °C min⁻¹).

### 3 Results and Discussion

#### 3.1 Catalyst Characterisation

CuZnO/ZSM-5 integrated catalysts (20 wt.% Cu and Zn, Cu/Zn mol = 1) were prepared via oxalate gel precipitation (OG) and chemical vapour impregnation (CVI) over commercially sourced ZSM-5 zeolites (Si/Al = 80, 50 and 23). Precipitation of Cu and Zn nitrate precursors in the presence of carbonates or hydroxides is commonly reported for the preparation of CH₃OH synthesis catalysts [11], however, physicochemical properties strongly depend on preparation parameters such as pH, temperature, precipitation rate, aging and drying [22–24]. Instead, the oxalate gel precipitation method is reported to yield small, well dispersed nanoparticles with high Cu surface area and homogeneous Cu/Zn distribution [25, 26]. This method was employed to prepare CuZnO/ZSM-5 integrated catalysts (CZZ-OG). CVI was also used for the preparation of CuZnO/ZSM-5 integrated catalysts (CZZ-CVI), because it allows the preparation of well dispersed bimetallic catalysts, and can be used over a wide range of supports, avoids the use of chloride metal precursors and avoids the presence of possible impurities from the use of solvents, ligands and reducing/precipitating agents [27–30]. Catalyst characterisation was performed on a portion of the pre-reduced sample (5% H₂/Ar, 220 °C, 2 °C min⁻¹, 1 h).

The porous structure of commercial ZSM-5 (Si/Al = 23, 50 and 80) zeolites after activation (static air, 550 °C, 5 °C min⁻¹, 4 h) was determined by N₂ adsorption. Due to the limitations of using N₂ as adsorbent for the analysis of porous materials (microporous filling due to strong adsorption at low p/p₀, non-uniform monolayer structure leading to different molecular N₂ areas) [31] and that commercially supplied zeolites were employed, values reported in this work were used to qualitatively compare materials. Comparable pore size (ca. 5.5 Å) and apparent BET surface area (ca. 460 m² g⁻¹) were measured for all three zeolites (Table S1). A slight decrease in the pore size was observed for CZZ-OG and CZZ-CVI catalysts, nevertheless, the BET surface area of the integrated catalysts was significantly lower than the parent zeolites (Table 1). Suggesting that Cu/ZnO nanoparticles were mainly at the surface and not within the zeolite pore structure. BET values are representative of the average surface area of the catalyst, for CZZ catalysts reported in this work, ZSM-5 zeolites were loaded with a ca. 9.7 wt.% Cu and ca. 10.3 wt.% Zn. Hence, a decrease in the surface area is expected, and not indicative of the metal distribution within the pore structure.
X-ray powder diffraction (XRD) was employed to determine the crystalline phases present in integrated catalysts after pre-reduction. For comparison, XRD patterns for CZZ(23)-OG, CZZ(23)-CVI and ZSM-5(23) are shown in Fig. 1 (XRD patterns for all integrated catalysts and activated ZSM-5 zeolites can be found in Figure S1 and Figure S2, respectively). The presence of metallic Cu (PDF 01-070-3038) was observed at 43.2°, 50.3° and 73.9°. Reflections not assigned to ZSM-5(23) (MFI morphology PDF 00-89-1421) or to Cu were attributed to the presence of ZnO (PDF 00-036-1451). Sharper Cu and ZnO peaks were observed for integrated catalysts prepared by CVI, compared to catalysts prepared by OG, indicating that the former methodology yielded larger crystallites. Oxalate gel co-precipitation has been reported to yield small and well distributed Cu/ZnO [25, 26], however, differences in the crystallite size between CVI and OG could be also attributed to the difference in annealing treatment. The former was annealed at 500 °C for 16 h to ensure the complete decomposition of the acetylacetone precursors, whilst for the latter the annealing at 500 °C was limited to 2 h to avoid unnecessary crystallite growth. No Cu2O (PDF 01-071-3645) or CuO (PDF 00-048-1548) peaks at 36.4° or 35.6° were observed. The Cu and ZnO crystallite size for CZZ-OG and CZZ-CVI catalysts was estimated using the Scherrer equation (Table 2). Details on how the Scherrer equation was applied to measure Cu and ZnO crystallite sizes can be found in the supporting information (Equation S1–S2). In accordance with N2 adsorption measurements, no significant changes in the pore size were observed between parent zeolites and integrated catalysts, and the measured Cu and ZnO crystallite sizes were considerably larger than the zeolite pores. Hence Cu and ZnO crystallites are physically unable to fit inside the zeolite pores. It was also observed that the oxalate gel precipitation yielded five-fold smaller Cu crystallites compared to CVI.

Morphological differences between CZZ(23)-OG and CZZ(23)-CVI integrated catalysts were analysed by TEM. Figure S3a and Figure S3b show low magnification TEM images for CZZ(23)-OG, it can be observed that most of the Cu and ZnO is not supported at the surface of the ZSM-5(23) zeolite, but instead forms large phase separated aggregates (nm–µm). These large aggregates contradict the Cu crystallite size determined by XRD through the Scherrer equation. However, as shown in Fig. 2a, the large aggregates are made of smaller nanoparticles, in the size range determined by XRD (ca. 15 nm as shown in Fig. 2b).

For CZZ(23)-CVI catalyst two distinct cases are observed. The Cu/ZnO nanoparticles are not well dispersed over every zeolite particle. Zeolite particles either display a complete absence of Cu/ZnO particles (Fig. 3a) or are covered with small and well-dispersed Cu/ZnO nanoparticles (Fig. 3b). Prolonged exposure of the “particle free” zeolite areas under the electron beam led to the formation of Cu/ZnO aggregates on the zeolite surface (Figure S4a to S4c). This behaviour was not observed for the CZZ-OG sample, furthermore, the zeolite showed no morphological change under the same beam conditions. This suggests that the Cu/ZnO is deposited as a film over the surface or as sub-nm particles or clusters inside the zeolite porous framework which are beyond the resolution of the microscope used, and were

| Catalyst   | Cu crystallite size/nm | ZnO crystallite size/nm |
|------------|------------------------|-------------------------|
| CZZ(80)-OG | 15                     | 14                      |
| CZZ(50)-OG | 13                     | 15                      |
| CZZ(23)-OG | 13                     | 15                      |
| CZZ(80)-CVI| 76                     | 21                      |
| CZZ(50)-CVI| 54                     | 22                      |
| CZZ(23)-CVI| 51                     | 38                      |

Table 2 Cu and ZnO crystallite sizes for CZZ-OG and CZZ-CVI catalysts obtained from the Scherrer equation.
formed by beam induced sintering, which could be related to the large Cu and ZnO crystallite sizes observed by XRD analysis.

The CH$_3$OH dehydration to DME is an acid catalysed reaction. Because of the presence of Brønsted and Lewis acid sites zeolites are less prone to deactivation than Al$_2$O$_3$ during CH$_3$OH dehydration. This is explained by the higher affinity of Brønsted acid sites towards CH$_3$OH compared to water [32]. The acidity of zeolites is an important parameter to optimise CH$_3$OH dehydration. Increasing the Al concentration within the structure strengthens the acidity, this leads to improved CH$_3$OH conversion, but usually at the expense of DME selectivity towards the production of other hydrocarbons, and decreases catalyst lifetime due to coke deposition [33]. NH$_3$ is normally employed to characterise the acid properties of porous materials because of its strong basicity and small molecular size. Ammonia temperature programmed desorption (NH$_3$-TPD) was employed to compare acid properties of activated zeolites and Cu–ZnO integrated catalysts. It is worth remembering that NH$_3$-TPD interpretation is challenging because it depends on experimental conditions (e.g. amount of sample, pre-treatment) and several desorption phenomena (e.g. NH$_3$ diffusion within pore structure and re-adsorption) [34, 35]. Often peak deconvolution models are needed to fully interpret NH$_3$-TPD profiles. To add to the NH$_3$-TPD interpretation complexity, one must consider that ZSM-5 zeolites can have 96 different Brønsted acid sites (oxygen bridging a Si and a Al atoms in the framework, although two positions are predominantly stable) [36, 37], which would need to be included into the NH$_3$-TPD peak model. Hence, NH$_3$-TPD measurements in this work are discussed in a qualitative manner. Figure S5 shows NH$_3$-TPD profiles for activated ZSM-5 zeolites. Two distinct peaks were observed; a first desorption peak between 230 and 260 °C associated with weakly bonded NH$_3$, likely to correspond to Lewis acid sites, and a second peak between 440 and 500 °C associated with more strongly bound NH$_3$, likely related to Brønsted acid sites [38]. Although, as observed by the clear shoulder on the first desorption peak for ZSM-5(50), this first peak is made of several desorption contributions. As expected, higher Al content in the zeolite resulted in a higher concentration of acid (higher peak area)
sites and stronger acidity (peak shift towards higher temperature) [20]. NH₃-TPD profiles for CZZ-CVI and CZZ-OG integrated catalysts are shown in Fig. 4a and Fig. 4b, respectively. The NH₃-TPD for integrated catalyst shows that the acid sites of parent zeolites have significantly changed after the addition of Cu and ZnO. Instead of the two distinct peaks observed for ZSM-5 indicating the presence of Lewis and Bronsted acid sites, the integrated catalysts showed multiple NH₃ desorption processes with an extra feature associated with medium acidity (300–380 °C). Moreover, the concentration of strong acid sites increased for synthesised integrated catalysts compared to parent zeolites, as also reported for other CuZnO-zeolite hybrid systems [38, 39]. The acid properties between CZZ(50 and 80)-CVI and CZZ(50 and 80)-OG are fairly comparable. However, a considerably higher concentration of strong acid sites was detected for CZZ(23)-CVI compared to CZZ(23)-OG.

3.2 Catalyst Performance for DME Production

Synthesised CZZ integrated catalysts are bifunctional catalysts, containing Cu-active sites for the CO₂ hydrogenation to CH₃OH and acid sites within the zeolite for the consecutive dehydration of CH₃OH to DME. The catalytic CO₂ hydrogenation (CO₂/H₂/N₂ = 1/3/1, 30 ml min⁻¹) to oxygenates (CH₃OH and DME) was assessed at 230–310 °C and 20 bar. Prior to reaction catalysts were pre-reduced in situ with 5% H₂/Ar (5% H₂/Ar, 220 °C, 2 °C min⁻¹, 1 h). Details on how CO₂ conversion, product selectivity and productivities can be found in the supporting information (Equation S3–S11).

The catalytic trend expected for the CO₂ hydrogenation to CH₃OH with temperature was observed on all synthesised catalysts (Table S2). Increasing the reaction temperature from 230 to 310 °C facilitates CO₂ activation, which translates into higher CO₂ conversion. However, as dictated by thermodynamics, higher temperature favour the endothermic RGWS instead of the exothermic CH₃OH (and DME) synthesis, as observed by the increase in CO selectivity and productivity with temperature [7]. For instance, at 20 bar (CO₂/H₂ = 1/3) the theoretical CH₃OH yield decreases from 10.89 (C mol %) at 230 °C to 1.45 (C mol %) at 300 °C, whilst the CO yield increases from 9.10 to 23.02 (C mol %). When CH₃OH is simultaneously dehydrated a DME yield of 20.26 (C mol %) can be achieved at 230 °C, but it is limited to 1.39 (C mol %) at 300 °C [7].

CZZ-CVI catalysts showed poor CO₂ conversion (ca. 3% at 250 °C) compared to CZZ-OG (ca. 14–20% at 250 °C). This could indicate that mass transfer of CO₂ is limited over CZZ-CVI, which could be attributed to Cu/ZnO blocking the surface of zeolite particles as suggested by TEM. Whilst CZZ-OG catalysts were formed by Cu/ZnO agglomerates not deposited on the zeolite surface, which resulted in an unobstructed zeolite porous structure. Moreover, no hydrocarbons (HC) were observed over CZZ-CVI catalysts, even at 310 °C. The absence of HC on CZZ-CVI catalysts suggested that either CH₃OH cannot diffuse into the ZSM-5 acid sites or that HC formed cannot diffuse out of the zeolite pores. Both scenarios suggest that Cu/ZnO might be
blocking the entrance of ZSM-5 channels. The selectivity
towards oxygenates over CZZ-OG catalysts is higher than
over CZZ-CVI, higher selectivity in addition to higher \( \text{CO}_2 \)
conversion translated in almost a six-fold higher oxygen-
ate productivity over CZZ-OG compared to CZZ-CVI at
230 °C (Fig. 5). As expected from thermodynamics, the
higher oxygenate productivity was observed at lower tem-
perature, while increasing the reaction temperature above
250 °C resulted in a decrease in oxygenate productivity over
CZZ(80 and 50)-OG catalysts. This can be assigned to equi-
librium limitations at high temperature, for both \( \text{CH}_3\text{OH} \) and
DME synthesis from \( \text{CO}_2 \) hydrogenation [7], as well as to
the more favoured side reactions, leading to the formation
of hydrocarbons [32]. Increasing the Si/Al ratio from 50 to
80 favoured \( \text{CH}_3\text{OH} \) dehydration to DME, leading to higher
oxygenates productivity. However, further increasing the Si/
Al ratio, as in CZZ(23)-OG, resulted in the conversion of
oxygenates to undesired by-products.

The formation of undesired hydrocarbons is to be avoided
not only because it decreases the selectivity towards oxygen-
ates and complicates the separation process, but also because
it is normally associated with zeolites deactivation through
coke formation [40]. ZSM-5 zeolites are widely used for
the synthesis of HC from \( \text{CH}_3\text{OH} \) and DME [41–43], in the
so called methanol-to-hydrocarbons process (MTH) [44].
On MTH, methanol absorbs on Brønsted acid sites yielding
surface methoxy species, methoxy species recombine form-
ing \( \text{C}–\text{C} \) bonds (carbene species) with increasing reaction
temperature [45]. Carbene species promote the autocatalytic
formation of hydrocarbons via the dual-cycle mechanism
[46–48]. The Si/Al ratio determines the concentration of
acid sites on zeolites, increasing the Al concentration results
in a higher concentration of Brønsted acid sites, increasing
the chance of forming active carbene species leading to
greater hydrocarbon formation [49]. The effect of increas-
ing the Al content on CZZ-OG catalysts on the productivity
of hydrocarbons is shown in Fig. 6. No HC were detected
over CZZ(80)-OG and CZZ(50)-OG below 270 and 250 °C,
respectively. HC selectivity remained low (< 0.03%) for
CZZ(50 and 80)-OG catalysts at 290 °C, with ethane being
the only by-product detected. Nevertheless, a selectivity of
c.a. 0.5% towards HC was observed over CZZ(23)-OG, even
at 230 °C. Increasing the reaction temperature from 230 to
270 °C resulted in an increase in the HC selectivity to 3.4%.
As shown in Fig. 5, oxygenate productivity for CZZ(23)-
OG does not increase with temperature up to 250 °C, as
observed for CZZ(50 and 80)-OG, but instead decreases
steadily with temperature, indicating the conversion of oxy-
genates to undesired by-products. Over CZZ(23)-OG a range
of by-products up to \( \text{C}_5 \) hydrocarbons were detected. Further
increasing the reaction temperature to 310 °C resulted in a
drop in HC productivity (Table S2), which likely relates to
low \( \text{CH}_3\text{OH} \) availability.

4 Conclusions

The preparation methodology of integrated catalysts has a
great influence on the Cu/ZnO crystallite size and their dis-
tribution over ZSM-5 particles. Thus, influencing the \( \text{CO}_2 \)
hydrogenation activity towards \( \text{CH}_3\text{OH} \) and the concomitant
dehydration to DME.

CVI preparation leads to bigger Cu and ZnO nanoparticles,
and an uneven distribution on CZZ-CVI catalysts: small nanoparticles evenly distributed over the surface of
ZSM-5 were observed, whilst what appeared to be ZSM-5
areas free of metal showed particle agglomeration under
the TEM electron beam. Suggesting that Cu and ZnO were deposited in the outermost zeolite pores or as a film at the surface, leading to poor CO₂ conversion and to possibly mass transfer limitations. In this case, possibly a better route to avoid mass transfer limitations would be to prepare Cu/ ZnO methanol catalysts by CVI and physically mixed them with ZSM-5 zeolites.

OG resulted in Cu/ZnO agglomerates, made of smaller 20 nm aggregates, that did not obstruct the zeolite pores. Which resulted in higher CO₂ conversion and oxygenates productivity compared to CZZ-CVI catalysts. The presence of acid sites is necessary to dehydrate CH₃OH to DME. However, a compromise between acid sites concentration is required to avoid excessive formation of hydrocarbons. The high concentration of acid sites on CZZ(23)-OG led to the conversion of oxygenates to HC, whilst no HC were detected for CZZ(50 and 80)-OG below 250 °C, the temperature with the higher oxygenates productivity (ca. 800 mmol kg⁻¹ h⁻¹ over CZZ(50)-OG).

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Conflict of Interest The authors declare that they have no conflict of interest.

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