Comparative TPR and TPD Studies of Cu and Ca Promotion on Fe-Zn- and Fe-Zn-Zr-Based Fischer-Tropsch Catalysts

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Abstract — Comparative TPR and TPD Studies of Cu and Ca Promotion on Fe-Zn- and Fe-Zn-Zr-Based Fischer-Tropsch Catalysts — The present study demonstrates the effect of zirconium promotion on Fe-Zn-based catalysts to boost the active sites of Fischer-Tropsch (FT) catalysts. The catalysts are also promoted by Cu and Ca and the active sites are examined using Temperature-Programmed Reduction (TPR) with H2 and CO and Temperature-Programmed Desorption (TPD) with NH3 and CO2. The results are presented as a comparative study between Fe-Zn- and Fe-Zn-Zr-based catalysts. The results show that addition of Zr to Fe-Zn catalysts increases the availability and dispersion of the precursor to the active sites and promotion with Cu and Ca independently and synergistically enhances reduction of Fe-Zn-Zr-based catalysts. The presence of Ca promotes carburisation, while Cu inhibits carburisation. The impact of the Ca and Cu on the surface acidity/basicity is governed by the nature of the interaction between the phases in the catalysts. The extent of reduction reflects the availability and dispersion of the precursor to the active phase, while the extent of carburisation will impact on the selectivity of the catalysts.
INTRODUCTION

Presently, global energy needs are being met largely by fossil fuels. In particular, due to their high energy density, liquid fuels occupy the central place in the present energy system. Although for socioeconomic, political and environmental reasons, there are motivations for a paradigm shift from fossil fuels to a hydrogen-Based renewable energy-based economy, liquid fuels will still play an important role in the future energy system. Petroleum is the primary resource of liquid fuels; however, there is increasing awareness of the diminishing world crude oil reserves [1, 2]. This has brought about a renaissance of discussions on FT technology in recent times [3-5]. FT technology provides an alternative route to liquid fuels from many carbonaceous resources. Moreover, discovery of more gas resources and regulations to discourage flaring of associated gases in oil wells has been a boost to the current wave of GTL projects [5]. GTL is typically based on the LTFT operation mode.

Catalyst design is a critical aspect of FT technology. Commercial LTFT catalysts are based on iron and cobalt [6]. Neither of the two metals have an absolute advantage over the other. Cobalt catalysts generally have higher intrinsic activity, exhibit lower Water Gas Shift (WGS) activity, and more resistance to moisture and coking-induced deactivation than iron-based catalysts. Hence, there seems to be a preference for cobalt-based catalysts for GTL [7]. However, iron-based catalysts are more resistant to sulphur poisoning, exhibit lower methane selectivity and their product selectivities are less susceptible to variations in process conditions. Moreover, a typical iron-based FT catalyst is about 25 times less expensive than a cobalt-based FT catalyst [8]. In addition, ease of refining of syncrudes to a specified diesel fuel indicates that iron-based LTFT syncrude poses less refining challenges compared with cobalt-based LTFT [9]. Thus, cobalt-based catalysts do not appear to have an absolute advantage over iron-based catalysts. In fact, with rational catalyst design, some of the previous advantages of cobalt over iron as a FT catalyst may cease to exist, while iron retains the lower cost edge that can lead to an enhanced process and FT process economy, and hence the motivation for this study.

The main areas of improvement needed for iron to compete head-to-head with cobalt for LTFT are: lowering of WGS activity/CO2 selectivity, increased activity and improved resistance to coking and deactivation by moisture. In a recent review report, the present authors showed that the key catalytic features that govern the performance of a FT catalyst are: hydrogenation activity, surface acidity and basicity. Hydrogenation activity determines the hydrocarbon formation rate, while surface acidity/basicity influences the product selectivity of the catalyst. State-of-the-art commercial iron-based (Fe-SiO2) catalysts are WGS active, which accounts for their higher CO2 selectivity in LTFT compared with cobalt-based catalysts. Attention has been drawn to the ZnFe2O4 spinel phase and its effect on CO2 selectivity in Fe-Zn-based catalysts [10]. Several reports examine CO [11] and CO2 [12-14] hydrogenation over Fe-Zn-based catalysts and the present authors have arrived at an optimum Zn/Fe ratio of 0.25 to achieve low CO2 selectivity with a minimised compromise of the FTS active site precursor. Although copper and potassium are benchmark promoters in commercial iron-based FT catalysts, potassium has been found to enhance WGS activity in Fe-Zn-based catalysts. To circumvent the undesired boost of WGS activity, we have use calcium in place of potassium. LTFT tests of the unpromoted, Cu- and Ca-promoted catalysts show very low CO2 selectivity. The Cu-promoted Fe-Zn catalyst has a lower surface area, higher hydrocarbon formation rate and virtually zero wax selectivity. The unpromoted Fe-Zn catalyst displays higher wax selectivity than the Ca-promoted catalyst.

It has been argued that the lower FT synthesis rate on Fe-based catalysts compared with cobalt-based catalysts is closely related to lower dispersion of the active components in iron catalysts. Li et al. [15] demonstrated that, in addition to the lower reaction temperature typical of LTFT, iron catalysts of comparable hydrocarbon synthesis rates to their cobalt-based counterparts can be achieved by design strategies that lead to high dispersion of the active component. They have shown that Fe-Zn-K-Cu oxide precursors prepared using a solvent replacement method have higher hydrocarbon synthesis rates (per catalyst mass or volume) compared with those on Co-based catalysts under identical reaction conditions. It has been shown that an increase in dispersion of active sites and higher TOF can be achieved by addition of another promoter (such as Zr, Mn or Cr) to the Fe oxide precursor [16, 17]. Achieving increased dispersion of active sites by introduction and co-precipitation of an additional promoter is more adaptable for large-scale operation than the solvent replacement method.

Building on the low CO2 selectivity of the Fe-Zn-based catalysts, we seek a way to bring their activities on par with that of a typical cobalt-based catalyst. Thus, in this contribution, we explored the introduction of Zr promotion as an easily scalable method to enhance the activity of the Fe-Zn catalysts. We examined the effect of addition of Zr on the FT relevant active sites in the catalysts. Although dispersion of active sites may correlate with textural properties, the results we obtained in the LTFT tests of the
Fe-Zn catalysts suggest their performance is governed by interplay of their active sites rather than their textural properties. The $\text{Fe}_2\text{O}_3 \rightarrow \text{Fe}_3\text{O}_4$ reduction peak in the $\text{H}_2$-TPR profiles of the catalysts is a measure of the hydrogenation activity of the catalysts. Low peak temperature indicates high hydrogenation tendency. The area under the peak $\text{vis-à-vis}$ amount of hydrogen consumed gives a measure of the density of the FTS active precursors. The extent of CO carburisation provides a measure of wax formation, coking tendency and resistance to moisture deactivation. Surface acidity/basicity has an influence on the product distributions of the catalysts. Hence, we focused our study on these characterisation techniques since they provide a direct measure of the active sites.

1 EXPERIMENTAL

1.1 Catalyst Preparation

The catalysts are prepared by the co-precipitation technique. $\text{Na}_2\text{CO}_3$ (2.5 M) solution is added dropwise to a solution (0.5 M) containing the nitrate salts of the metals in the desired ratio to pH $= 8.5 \pm 0.5$ while the temperature is maintained between 70-80°C under mixing with the aid of a motor stirrer. The resulting precipitates are washed with double-distilled water until the filtrates are nitrate-free. The precipitates are dried overnight and calcined at 400°C for 4 hours. The atomic ratio of the constituent metal in the catalysts used in this study is presented in Table 1.

1.2 Temperature-Programmed Reduction (TPR)

TPR and TPD profiles of the catalysts are obtained with ChemiSorb 2720 (Micrometrics, USA) equipped with a TCD detector. The $\text{H}_2$-TPR profiles are obtained by reducing the catalyst samples by a gas mixture of 10% $\text{H}_2$ in Ar with a flow rate of 20 mL/min, and the temperature is increased from ambient to 800°C at a rate of 10°C/min. The CO-TPR are obtained by heating the catalyst samples by a gas mixture of 5.2% CO in He with a flow rate of 20 mL/min and the temperature is increased from ambient to 800°C at a rate of 10°C/min. Hydrogen consumption in the TPR peak area is evaluated by the peak area of the CuO TPR calibration.

1.3 Temperature-Programmed Desorption (TPD)

The TPD is carried out by first reducing a catalyst with $\text{H}_2$ at 400°C for 8 h. The gas flow is then switched to He and the temperature is raised to 450°C at a rate of 10°C/min and kept at 450°C for 30 min and then cooled to ambient temperature to eliminate adsorbed $\text{H}_2$ from the catalyst surface. $\text{NH}_3$ uptake on the catalyst surface takes place when the gas flow is changed to 4% $\text{NH}_3$ in He. The $\text{NH}_3$-TPD profile is obtained by a temperature-programmed desorption under He with a flow rate of 20 mL/min from ambient temperature to 1 000°C at a rate of 10°C/min. $\text{CO}_2$-TPD is obtained by the same procedure using 5.2% $\text{CO}_2$ in He for the $\text{CO}_2$ uptake. The amounts of $\text{NH}_3$ and $\text{CO}_2$ desorbed in the TPD peak area are evaluated by comparing with a calibration graph obtained by measured pulses of $\text{NH}_3$ and $\text{CO}_2$ in a flow of He.

2 RESULTS AND DISCUSSION

2.1 Reduction Behaviour

Fe-Zn oxide precursors are transformed into $\alpha$-$\text{Fe}_2\text{O}_3$ and Zn$\text{Fe}_2\text{O}_4$ phases after calcination at 400°C. The $\alpha$-$\text{Fe}_2\text{O}_3$ phase is recognised to be the precursor to the active sites, carbide phases, during the activation/carburisation step prior to the FT reaction. Figure 1 shows the H-TPR profiles of Fe-Zn oxide at the Zn/Fe ratios 0, 0.1, 0.2 and 0.25. The profile of the zinc-free oxide displays two features, a narrow peak

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<tr>
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<td>Fe-Zn-Zr-Cu</td>
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at 370°C and broad multiple peaks in the range of temperatures 400-900°C. The narrow peaks are assigned to the Fe₂O₃ → Fe₃O₄ transition, while the broad multiple peaks are assigned to Fe₃O₄ → FeO → Fe transitions. The H-TPR profiles of Fe-Zn oxides display similar features to the zinc-free Fe oxide; hence, similar assignments are made for them. The temperature of the peak assigned to the Fe₂O₃ → Fe₃O₄ transition decreases with increasing zinc content. However, the areas under the peaks decrease with increasing zinc content. Part of the iron is used up at the stoichiometric ratio of 2:1 (Zn:Fe) in forming the ZnFe₂O₄ phase. Analyses of the TPR profiles confirm the stoichiometric ratio. The calculated percentage of hydrogen consumption under these peaks corresponds with the experimental value determined by the proportion of the peak to the total hydrogen consumption in each case (Fig. 2). At a Zn/Fe ratio of 0.25, the proportions of the α-Fe₂O₃ and ZnFe₂O₄ phases are expected stoichiometrically. The effects of the promoters are examined on the Fe-Zn (Zn/Fe = 0.25) catalysts.

ZnFe₂O₄, being less reducible than α-Fe₂O₃, is regarded as a textural promoter that increases the surface area and dispersion and prevents the sintering of the α-Fe₂O₃ phase [12]. However, the ZnFe₂O₄ phase has been implicated as an active component which is capable of converting CO₂ into methanol [12-14]. It can therefore be suggested that the ZnFe₂O₄ phase does not only act as a structural promoter, it also plays a key role in the lower CO₂ selectivity by in situ recycling of CO₂ generated by WGS and oxygen removal via a direct CO dissociation mechanism [18]. Phases due to the promoter are virtually absent in X-ray diffractograms of the catalysts. Moreover, the α-Fe₂O₃ phase is the main active phase that is directly connected to hydrocarbon formation; hence, we examine the presence of the promoters which affect the reduction and carburisation of the α-Fe₂O₃ phase in the catalysts.

The readily reducible phases in the catalysts are CuO and α-Fe₂O₃. The promotion effect of Cu on the reduction of α-Fe₂O₃ through the hydrogen spillover effect is well documented. Figure 3 shows the H₂-TPR profiles of the catalysts grouped into Fe-Zn series and Fe-Zn-Zr series. The partition line is used to demarcate the peaks assigned to the Fe₂O₃ → Fe₃O₄ transition. In the two groups, it can be observed that the presence of Ca has no influence on the reducibility of the α-Fe₂O₃ phase. However, Cu promotion enhances the conversion of Fe₂O₃ into Fe₃O₄. In the Fe-Zn series, the peak temperature of the Fe₂O₃ → Fe₃O₄ transition is lowered by 125°C. Co-promotion of Fe-Zn with Cu and Ca further decreases the Fe₂O₃ → Fe₃O₄ reduction temperature by about 40°C. A similar trend of reduction of peak temperature is observed in the Fe-Zn-Zr series. It is noteworthy that the peak temperature of the Fe₂O₃ → Fe₃O₄...
transition in Fe-Zn-Zr-Cu is lower than that of Fe-Zn-Cu. It appears that Cu-CaOx and Cu-ZrO\(_2\) couples in the catalyst display higher efficiency of H\(_2\) cleavage and the hydrogen spillover effect towards the conversion of Fe\(_2\)O\(_3\) into Fe\(_3\)O\(_4\) compared with Cu alone in the catalysts. This suggests that a complementary Lewis basic site is beneficial to a transition metal site (Lewis acidic) for efficient H\(_2\) activation [19].

It is important to draw attention to the right hand side of the partition in Figure 3. There are two reduction peaks in this region of the TPR profiles. In the Fe-Zn series, distinction of the peaks appears more apparent for the Ca-containing catalysts but it becomes less noticeable in the Cu-containing catalysts. The profiles of the Fe-Zn-Zr reveal a similar pattern but here, distinction of the peaks is more noticeable. The presence of Zr and Ca delineates the transition under the peaks and this delineation of the peaks in this region may be attributed to segregation of the phases under the peaks. The segregation appears stronger with Zr than with Ca but it is additive in the presence of Zr and Ca. However, the peak differentiation effect becomes blurred in the Cu-containing catalysts.

A comparison of hydrogen consumptions for the Fe\(_2\)O\(_3\) → Fe\(_3\)O\(_4\) transitions for the catalysts as measured by the area under the peaks is shown in Figure 4. The figure shows that hydrogen consumptions for the Fe\(_2\)O\(_3\) → Fe\(_3\)O\(_4\) transitions in the Fe-Zn series catalysts are very similar. This suggests that the amount of the Fe\(_2\)O\(_3\) phase in the Fe-Zn series catalysts is broadly the same. However, hydrogen consumptions for the Fe\(_2\)O\(_3\) → Fe\(_3\)O\(_4\) transitions in the Fe-Zn-Zr catalyst series present an interesting trend. Beginning from Fe-Zn-Zr, slightly higher hydrogen consumption is observed compared with those of the Fe-Zn series. The most probable reaction at this peak temperature is Fe\(_2\)O\(_3\) → Fe\(_3\)O\(_4\), which suggests that the presence of ZrO\(_2\) in the Fe-Zn oxide precursor decreases the proportion of Fe that is used up in forming the ZnFe\(_2\)O\(_4\) phases. It implies that Zr promotion of Fe-Zn leads to an increased proportion of active site precursors, the Fe\(_2\)O\(_3\) phase, while also serving as a textural component for increased dispersion of the \(\alpha\)-Fe\(_2\)O\(_3\) or ZnFe\(_2\)O\(_4\) phases. This promotion effect is enhanced when Ca is added to Fe-Zn-Zr. Interestingly, Fe-Zn-Zr-Cu displays greater hydrogen consumption than Fe-Zn-Zr-Ca. Here, CuO is also a beneficiary of the dispersion effect of Zr which, in turn, enhances the hydrogen spillover effect. The increased dispersion of the \(\alpha\)-Fe\(_2\)O\(_3\) and ZnFe\(_2\)O\(_4\) phases enhances the hydrogen spillover effect, leading to increased reduction and nucleation of the surface oxides. The promotion effect is further amplified when Fe-Zn-Zr is co-promoted with Cu and Ca. This unprecedented hydrogen consumption in Fe-Zn-Zr-Cu-Ca may be explained as being due to the combined dispersion effect of Zr and Ca and the enhanced hydrogen spillover effect of Cu. The presence of Zr and Ca produces dramatic dispersion of the \(\alpha\)-Fe\(_2\)O\(_3\) and ZnFe\(_2\)O\(_4\) phases, leading to their facile reduction in the presence of Cu. Such synergistic effects of promoters for enhancing the dispersion of active components have not been reported before for iron-based FT catalysts.

### 2.2 Carburisation Behaviour

Figure 5 shows the CO-TPR profiles of the catalysts. The CO-TPR profiles provide information about the carburisation behaviour of the catalysts. The peaks in the CO-TPR profiles are not well resolved, as is the case for the H\(_2\)-TPR profiles. This is because reduction and carburisation occur concurrently during the temperature programme under the flow of CO gas, whereas the H\(_2\)-TPR process is void of this complication. This translates into a challenge in the
interpretation of CO-TPR profiles obtained by a TCD signal. A close examination of Figure 6 shows that the catalysts display similar CO-TPR profile patterns. These are made up of small, broad low temperature peaks below 400°C and larger peaks with shoulders above 400°C. According to Li et al. [11], Fe₂O₃ reduces to Fe₃O₄ in CO at 270°C and subsequent carburisation of Fe₃O₄ occurs above a temperature of 270°C. However, at temperatures above 400°C, carburisation is accompanied by excessive formation of amorphous carbon due to disproportionation of CO (Boudouard reaction). This implies that the peaks at temperatures above 400°C are of little or no relevance in determining catalyst behaviour in actual FT conditions. Thus, further discussion on the CO-TPR profile of the catalysts will be limited to the peaks below 400°C. Since carburisation of the iron oxide phase proceeds without first having been reduced to the metallic state and peak assignment is rather difficult, we looked at CO consumption to estimate the extent of carburisation of the catalysts at temperatures below 400°C. We adopted the reaction:

$$\text{Fe}_2\text{O}_3 + \frac{1}{3} \text{CO} \rightarrow \frac{2}{3} \text{Fe}_3\text{O}_4 + \frac{1}{3} \text{CO}_2$$

as a reference, and the amounts of CO required for this reaction for Fe-Zn and Fe-Zn-Zr series are 2 783 and 2 656 μmol g⁻¹, respectively. CO consumption in excess of the reference amount is taken as CO consumption for carburisation. The CO consumption of the catalysts is presented in Figure 6, which shows that CO consumption in all the catalysts is above the reference amount. This reiterates that the Fe₂O₃ → Fe₃O₄ is facile below 400°C and carburisation of the catalyst can be carried out. Ca promotion of Fe-Zn does not significantly affect CO consumption but addition of Cu to Fe-Zn decreases the CO consumption, indicating that Cu promotion leads to resistance to carburisation.

The Fe-Zn-Zr catalyst series display higher CO consumption than their Fe-Zn counterparts. This may be attributed to the higher amount of α-Fe₂O₃ and the textural promotion effect of Zr in the Fe-Zn-Zr catalyst series. Addition of Ca to Fe-Zn-Zr produces a dramatic increase in CO consumption; this is most probably due to increased dispersion of the α-Fe₂O₃ and ZnFe₂O₄ phases. However, Cu promotion of Fe-Zn-Zr did not significantly affect the carburisation. Here, carburisation resistance of Cu is masked by the presence of Zr on the catalysts. The Cu particles are preferentially distributed close to Zr. This preferential distribution of Cu particles seems relapsed when co-promoted with Ca, and the carburisation resistance effect of Cu re-emerges to account for the lower CO consumption in Fe-Zn-Zr-Cu-Ca compared with Fe-Zn-Zr-Cu and Fe-Zn-Zr. This carburisation behaviour of the catalysts is expected to impact on their performance for FT operations. FT evaluations of the catalysts are ongoing in our laboratory.

2.3 Surface Acidity

Catalyst surface acidity impacts on FT production distribution. High surface acidity promotes acid-catalysed secondary reactions (such as cracking, isomerisation and aromatisation) of primary FT products. It can also aid rapid deactivation of the catalysts due to coke formation. The NH₃-TPD profiles presented in Figure 7 give information on the surface acidity of the two catalysts’ series. For simpler quantitative interpretations, the profiles are partitioned into two temperature ranges to rank

![Figure 6](image6.png)

Figure 6
CO consumption measured by area under the peak in different temperature ranges in the CO-TPR profiles of the catalysts.

![Figure 7](image7.png)

Figure 7
NH₃-TPD profiles of the catalysts.
the surface acid strength in increasing order of the peak temperatures.

The distributions of the surface acid sites of the catalysts are presented in Figure 8. As can be seen from Figure 7, a sharp peak of strong sites emerges on the profile of the Fe-Zn-Ca catalyst (a similar peak is present in the CO₂-TPD profile of the catalyst). A peak connoting the strong acidic site of this nature is quite unexpected for an alkaline cation promotion. The possible phases of Ca in the catalysts are CaO, Ca(OH)₂ or CaCO₃. Ordinarily, precipitation of calcium nitrate with sodium carbonate will produce calcium carbonate and decomposition of CaCO₃ normally takes place at > 800°C, but the peak temperature (580°C) corresponds with the temperature of the reaction [20]:

\[ \text{Ca(OH)}_2(s) \rightarrow \text{CaO}(s) + \text{H}_2\text{O}(g) \]

It thus suggests that Ca is present as the Ca(OH)₂ phase in the Fe-Zn-Ca catalyst. Another plausible explanation is that Ca²⁺ is incorporated into the lattice of α-Fe₂O₃ or ZnFe₂O₄ with strongly bonded hydroxide ions. Cu promotion decreases the acid site densities of Fe-Zn. It appears that Cu titrates the weak acid sites.

Addition of Zr to Fe-Zn brings about site redistribution in favour of weaker acidic sites. Here, Cu promotion of Fe-Zn-Zr leads to an increased proportion of strong acid sites and a slight increase in the total acid sites’ density. However, the presence of Ca in Fe-Zn-Zr brings about almost double the density of the acidic sites. This situation is most probably due to the dispersion effect of Zr discussed earlier and its enhancement when jointly added with Ca to the α-Fe₂O₃ and ZnFe₂O₄ phases. It appears that addition of Ca to Fe-Zn-Zr-Cu produces a similar effect to that observed when Fe-Zn is promoted with Ca. When compared with Fe-Zn-Zr-Cu, strong acid sites seem to increase in Fe-Zn-Zr-Cu-Ca at the expense of weak-strength sites. The acid site distribution suggests that there is a greater tendency for an acid catalysed secondary reaction in the Fe-Zn-Zr than in the Fe-Zn series, while Cu and Ca promotion suppresses surface acidity in Fe-Zn and is again reinforced in Fe-Zn-Zr.

### 2.4 Surface Basicity

The CO₂-TPD profiles of the catalysts are shown in Figure 9. The profile is partitioned into two regions as in the NH₃-TPD profiles, and the densities of the basic sites of the catalysts are presented in Figure 10. The sharp peak at 580°C observed in the NH₃-TPD profile of Fe-Zn-Ca is also present in its CO₂-TPD profile. The implication of this peak has been discussed in Section 2.3. Cu promotion of Fe-Zn slightly increases the proportion of strong basic sites. The basicity of Ca and Cu promotion appears to cancel out in the Cu-Ca-promoted catalyst [21]. The presence of Zr in Fe-Zn does not significantly affect its surface basicity. However, Cu and Ca promotion in the Fe-Zn-Zr series displays a different trend compared with the Fe-Zn series. Creation of strong basic sites is obtained in the Cu-promoted catalyst. Here, Ca promotion imparts formation of strong basic sites and total basic site density is markedly increased. The cancelling-out effect of Cu-Ca co-promotion is also observed in the Fe-Zn-Zr-Cu-Ca catalyst.

Like acidic sites, basic sites can also influence FT product distribution. In our recent review report [10], we commented that the effect of basic sites impacting promoters on FT product selectivity, especially its potential toward producing non-ASF product distribution, has not been rigorously investigated, as is the case for acidic sites. As mentioned before, Cu and K are benchmark promoters in commercial iron (Fe-SiO₂)-

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**Figure 8**
Amount of NH₃ desorbed measured by area under the peak in different temperature ranges in the NH₃-TPD profiles of the catalysts.

**Figure 9**
CO₂-TPD profiles of the catalysts.
based FT catalysts, and most studies on effects of basic sites in FTS have focused on K. The K promoter is usually added to catalysts via impregnation with K₂CO₃ solution followed by calcination at 400-500°C. The impregnated K can present as the K₂CO₃, silicate phase or form a compound/complex with Fe oxides and/or silica. In all these phases, the impregnated K is prone to leaching out of the catalyst in the course of the FT reaction. Hence, the supposed basic sites imparted on the catalyst due to the presence of K may be short-lived and the product distributions obtained may lack the sustained influence of basic sites.

Carbonates, hydroxides and oxides of Group II metals are less susceptible to leaching under FT reaction conditions. They are better candidates for examining the effect of basic sites in FT product selectivity. However, there are very few investigations on this aspect where the Group II cations (nitrate salts) are added to Fe-SiO₂-based catalysts via impregnation [22-24] followed by calcination at 350-450°C for 4 h. There is a tendency for Ca to remain in the nitrate phase since Ca(NO₃)₂ decomposes at temperatures above 550°C even in a reducing atmosphere [25-26]. In this phase, Ca is susceptible to leaching, as described for Tao et al. [27] who introduced Ca into Fe-Mn oxide by co-precipitation. Here, the challenge of Ca(NO₃)₂ decomposition is absent. It also has the advantage of being a one-step method (co-precipitation) instead of a two-step method (co-precipitation followed by impregnation) using K. However, in their study, the authors further added K to the catalyst. This makes it difficult to delineate the promotion effect of Ca from that of K in their catalyst and make a direct comparison with the present study.

CONCLUSIONS

We explored Zr as a promoter for boosting the active sites in Fe-Zn-based catalysts towards developing iron catalysts that can match cobalt-based catalysts in FT activity. For this purpose, we used TPR (H₂ and CO) and TPD (NH₃ and CO₂) to gain insights into FTS active site precursors, hydrogenation tendency and surface acidity/basicity of the catalysts. The results are presented as a comparative study of the effect of Cu and Ca promotion on reduction and carburisation behaviour and surface acidity/basicity of Fe-Zn and Fe-Zn-Zr catalysts. The presence of Zr improved dispersion of the FTS active sites of precursors. Further, Cu and Ca synergistically increased the amount of the FTS active sites of precursors in Fe-Zn-Zr-Cu-Ca-based catalysts. The presence of Ca promotes carburisation but Cu inhibits carburisation. The impact of the Ca and Cu on the surface acidity/basicity is governed by the nature of the interaction between the phases in the catalysts. The extent of reduction reflects the availability and dispersion of the precursor to the active face, while the extent of carburisation will impact on the selectivity of the catalysts. The results from this study suggest that Fe-Zn-Zr-Cu-Ca may compare fairly well with a cobalt catalyst in terms of activity (per gram catalyst) and CO₂ selectivity. When this is confirmed by LTFT tests, the next goal will be to improve the stability of the catalyst.

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REFERENCES


Brockner W., Ehrhardt C., Gijjik M. (2007) Thermal decomposition of nickel nitrate hexahydrate, Ni(NO₃)₂·6H₂O, in comparison to Co(NO₃)₂·6H₂O and Ca(NO₃)₂·4H₂O, *Thermochimica Acta* 456, 64-68.


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