of dry air at 550 °C, conditions which would result in extensive dehydroxylation of the zeolite. According to the dehydroxylation model of Kühli and others, cationic Al species are formed as the framework is dealuminated. In fact, NMR results have shown that extensive dealumination can occur at temperatures as low as 500 °C. In the work presented here dehydroxylation and concomitant dealumination were avoided as was shown by the unit cell size of the catalyst after reaction. This is consistent with NMR results of Bösecke et al. which showed that upon deamination at 400 °C under "shallow-bed" conditions only about 3 Al/uc were removed from the framework.

Although the acid strength of different types of zeolites may differ considerably because of changes in bond angle, bond length, etc., it is interesting to compare the specific activity for hexane cracking over a dealuminated Y sample and HZSM-5 having Si/Al = 26. The activities at 350 °C were 11.4 and 8.5 μmol/ (g.min), respectively. Considering the differences in the zeolites the activities are quite similar. For cumene dealkylation the HZSM-5 catalyst was about twice as active as a comparable dealuminated Y, and for methanol dehydration the HZSM-5 was about 4 times as active. Thus, the relative activity depends very much on the probe reaction.

Conclusions

Hexane cracking is essentially a linear function of the lattice Al content over a range of Si/Al ratios from 4.6 to 255. To the extent that hexane cracking is a measure of strong acidity, the acid strength is constant over this range of Si/Al ratios. It is postulated that protons associated with isolated framework aluminum in the 4-rings give rise to the strong acidity which characterizes this material. The presence of aluminum in second-neighbor positions significantly decreases the acidity of a site. This is essentially the Dempsey-Mikovsky-Marshall (MM) model of strong acidity.

Consistent with this model, a normal H–Y zeolite, which has only a small amount of isolated tetrahedral aluminum, is relatively inactive for hexane cracking. Moreover, as described in part 1 the H–Y zeolite is relatively inactive for methanol dehydration, which requires only weak acid sites. This suggests that test reactions to characterize acid sites in zeolites need to be used with caution since a few strong acid sites may dominate the activity, even when a less-demanding reaction is employed.

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Characterization of a New Iron-on-Zeolite Y Fischer–Tropsch Catalyst

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Iron pentacarbonyl adsorbed on dry Na–Y zeolite can be oxidized at subambient temperatures into Fe₂O₃ located in the zeolite supercages (catalyst I). When catalyst I is hydrogen reduced at 357 K most of the iron has agglomerated externally to the zeolite (catalyst II). When the iron carbonyl is thermally decomposed in vacuo at 525 K, an iron phase with a very low degree of dispersion is again obtained (catalyst III). During a Fischer–Tropsch reaction most of the iron is transformed into a Hagg-like carbide phase, located externally to the zeolite. Catalysts II and III rapidly reach steady state and show a Schulz–Flory-type of product distribution, the Hagg carbide being the active phase. Catalyst I slowly moves to steady state and Schulz–Flory behavior. Product selectivity is only found on this catalyst during transient conditions. The physical information on the three catalysts before and after reaction was obtained with transmission electron microscopy and Mössbauer and EXAFS spectroscopies. These techniques supplied consistent and complementary evidence.

Introduction

Iron clusters can be introduced in zeolites via several indirect methods. Hydrogen reduction of Fe(II)-exchanged faujasite-type zeolites is only possible with aluminum-rich samples. Therefore, Fe(II) reduction in Y-type zeolites was attempted with stronger reducing agents. Alternatively, neutral iron complexes such as iron carbonyls can be adsorbed into the pores or cages of a large-pore zeolite and subsequently decomposed. The thermolysis of iron carbonyl adsorbed on zeolites, depending on the

exact decomposition method used, gives rise to discrete iron particle size distributions and dispersions. These systems have been

characterized till now by thermogravimetric, in situ infrared, and magnetic techniques.

For catalyst preparation the carbonyl decomposition has definite advantages over hydrogen reduction of cation-exchanged Fe(II). The activity of Y zeolite loaded with iron carbonyl has been investigated in a Fischer–Tropsch (FT) reaction. The experiments were done in a batch reactor and, although in this way only "average" product distributions are obtained, it was concluded that the zeolite matrix was influencing the product distribution by decreasing suddenly the hydrocarbon growth rates at the carbon numbers 8–10. Similar results have been obtained with Ru clusters on NaY zeolites. It remains therefore vital to decide on the exact influence of the zeolite on the FT selectivity.

In the present work, it was certified with several physical methods (Mössbauer and EXAFS spectroscopies, and transmission electron microscopy) that the FT active phase of the fresh catalyst was confined to the intracrystalline void space of zeolite Y. This catalyst was prepared by the iron carbonyl decomposition method. Reference zeolite-based iron catalysts containing substantial amounts of extra lattice iron particles were prepared as well. The physical characteristics of the steady-state catalysts were compared with fresh materials. This allowed us to draw pertinent conclusions on the way the Y zeolite matrix influences the FT selectivity.

Experimental Section

Materials. A synthetic Na-Y zeolite from Strem Chemicals was treated with an excess of a 0.1 M aqueous solution of NaCl, washed with distilled water till chloride-free, dried in air at 333 K, and stored in an atmosphere of 25% humidity. Iron penta-carbonyl (Fe(CO)₅) from Ventrton with a purity of 99.5% was coldly distilled in the dark and stored over molecular sieve 5A. All gases used, CO, H₂, Ar, He, and O₂, were from L'Air Liquide coldly distilled in the dark and stored over molecular sieve 5A. The catalyst was prepared by the iron carbonyl decomposition method. This allowed us to draw pertinent conclusions on the way the Y zeolite matrix influences the FT selectivity.

Methods. Catalyst Preparation. The moisture-saturated Na-Y zeolite was pelletized, crushed, and sieved. A grain size between 0.25 and 0.5 mm was used for further work. A 300-mg sample of this zeolite was loaded into a stainless steel tubular reactor with a 10-mm internal diameter. This reactor was first mounted in a large glass tube and connected to a grease-free gas handling system. The gas mixture was degassed at 670 K, cooled to 293 K. The catalyst was activated in a high-pressure continuous-flow tubular reactor. The effluent vapors were transferred to the analytical section with a transfer line heated at 420 K. The analytical procedure has been described in detail. It allowed accurate carbon mass balances within ±2%, to be determined. The reactor feed consisted of H₂, CO, and Ar in molar ratios of 55:40:5, respectively. The reaction pressure was 2.0 MPa and the total flow rate 1.8 L/h at the reactor entrance. The catalyst was activated in synthesis gas at 553 K. Product samples were taken periodically till catalyst steady state was reached.

Catalyst Characterization. Electron micrographs were taken with a Jeol JEM 200 CX instrument in the transmission mode (TEM). About 5 mg of sample was ground in an agate mortar and then suspended in 1 mL of ethanol (p.a.). The suspension was dropped on Formvar grids and dried. An electron beam magnification between 500000X and 3000000X was applied. Mössbauer spectra of the 14.4-keV γ-resonance of iron were taken at temperatures between 1.8 and 300 K with a Fe(Rh) source. Samples for Mössbauer spectroscopy were prepared in a quartz reactor under conditions that are similar to those applied in the catalytic reactor. Finally, they were embedded in paraffin so that sample transfer under inert conditions was possible. Thus, Mössbauer spectra were taken of samples representing the original chemical state of the catalysts before reaction.

EXAFS measurements of the K edge of iron were carried out at liquid nitrogen temperature with the spectrometer EXAFS II at HASYLAB (DESY, Hamburg, FRG).

Results and Discussion

Study of the Iron Phases on the Freshly Prepared Samples. Characterization of Highly Dispersed Na-Y/FeOₓ. For more information, please refer to the literature.


Figure 1. Transmission electron micrograph of sample Na-Y/FeOₓ. Dry Na-Y loaded with Fe(CO)₅ as described above was also heated under a vacuum of 1 mPa to 525 K. In this way, part of the adsorbed iron carbonyl is decomposed and another part is desorbed. The sample contains therefore only 6% iron and will be denoted further as Na/YFe(0).

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Results and Discussion

Study of the Iron Phase on the Freshly Prepared Samples. Characterization of Highly Dispersed Na-Y/FeOₓ. For more information, please refer to the literature.
than 50 years it has been known that when oxygen reacts at room temperature with Fe(CO)₅ adsorbed on high surface area supports Fe₂O₃ is formed.²² To avoid overheating during this reaction and consequently agglomeration of the supported species, the dry Na-Y zeolite containing on the average three molecules of the Fe(CO)₅ complex per faujasite supercage¹³,¹⁴ was contacted with oxygen only at 77 K. Gradual oxidation then took place when the sample was allowed to warm to room temperature. Indeed, the cream color of the iron pentacarbonyl–zeolite adduct turned gradually to a light brown. This was the product that has been characterized by TEM, Mössbauer, and EXAFS techniques. An electron micrograph of such a sample, denoted as Na-Y/Fe₂O₃, is shown in Figure 1. Even at the high magnifications applied, no resolved iron metal particles are visible on the zeolite support. This suggests that the Fe₂O₃ particles on Na-Y have an average size smaller than 2 nm. In agreement with the TEM data, an X-ray diffractogram (XRD) of this sample also shows no diffraction line attributable to Fe₂O₃.

From the Mössbauer spectra of sample NaY/FeO₃ shown in Figure 2, the following information can be derived:

i. At low temperatures, a symmetrical six-line pattern exists with a magnetic hyperfine line splitting of 42.8 T and an isomer shift (IS) of 0.3 mm s⁻¹.

ii. At higher temperatures, this pattern disappears, leaving only a symmetrical doublet with an IS of 0.3 mm s⁻¹ and a quadrupole splitting of Vₚ = 1.0 mm s⁻¹. Vₚ was derived by raising the temperature from 1.8 to 300 K. In Table I are collected relevant Mössbauer data on iron-containing systems. It follows that in the Na-Y/FeO₃ sample the presence of the following species can be excluded: iron(0) clusters or particles, iron carbides, Fe(II), and Fe(III) cations at exchangeable positions in the Y zeolite, and unreacted iron pentacarbonyl. This table indicates that the Mössbauer parameters of Fe₂O₃ either supported on Na-Y or on alumina are similar to those of the Na-Y/Fe₂O₃ zeolite. The values of the hyperfine field and the relaxation behavior of the system, however, suggest that very small particles of either α-Fe₂O₃ or γ-Fe₂O₃ are present. Since the value of the hyperfine field of α-Fe₂O₃ decreases with its particle size²³,²⁴ in the size range below 10 nm, a more precise assignment based on Mössbauer data is not possible. Additionally, as a result of the broad spectral width, accurate numerical evaluation of the Mössbauer parameters is difficult. Further evidence for a simple oxide phase dispersed in the zeolitic cages will be presented when the evaluation of the EXAFS Fourier transforms is discussed.

Characterization of Sample Na-Y/Fe(CO)₅. Thermal decomposition under vacuum of Fe(CO)₅ adsorbed on Na-Y zeolite...
shows an enhanced tendency to sinter compared to its rapid thermolysis under inert gas.\textsuperscript{11} The electron micrograph of such a Fe(CO)\textsubscript{5}-Na-Y zeolite adduct subjected to a slow temperature increase under vacuum is shown in Figure 3. It is seen that iron particles with an average size of about 30 nm are located at the outer surface of the zeolite crystals. The Mössbauer spectra on the same sample are given in Figure 4. A typical Fe(0) sextet with a magnetic hyperfine splitting of 33.0 T and a value of IS of 0.0 mm s\textsuperscript{-1} is obtained. These values are also observed for bulk iron (Table I). No relaxation behavior is encountered, since even at room temperature this sextet remains unchanged. This provides evidence for the existence of an almost pure Fe(0) phase with a particle size larger than the zeolite supercages. A small amount of iron(III) oxide, however, seems to be present on this sample since the weak lines have a hyperfine splitting comparable to sample Na-Y/FeO\textsubscript{2} (compare Figures 2 and 4).

The Fourier transform of the EXAFS spectrum of Na-Y/Fe(O) is shown in Figure 5A and compared to that of Na-Y/FeO\textsubscript{2} in Figure 5B. The EXAFS spectrum of sample Na-Y/Fe(O) resembles the pattern of bulk iron and had a nearest-neighbor distance \(d_{\text{nn}}\) of approximately 0.24 nm. For the Na-Y/FeO\textsubscript{2} sample, the main EXAFS peak is found at \(d_{\text{nn}}\) of approximately 0.18 nm. This distance is typical for iron(III) oxides.\textsuperscript{22} The EXAFS data, therefore, are in line with conclusions from TEM and Mössbauer spectroscopy: on Na-Y/Fe(O) large iron particles located outside the zeolite crystals constitute the main iron-containing phase, while on Na-Y/FeO\textsubscript{2} an FeO\textsubscript{2} phase is finely dispersed in the zeolite cages.

**Characterization of Sample Na-Y/FeO\textsubscript{2}-RED.** When sample Na-Y/FeO\textsubscript{2} is hydrogen reduced at relatively high temperatures (575 K) for an extended period of time (50 ks), the sample turns black, large iron particles are observed in TEM, and the Mössbauer spectra typical for sample Na-Y/Fe(O) are found. There is no doubt that for this material the dominating iron phase consists of large iron particles located at the outer surface of the zeolite crystals.

**Characterization of Na-Y/FeO\textsubscript{2} Used as a Fischer–Tropsch Catalyst.** Sample Na-Y/FeO\textsubscript{2} was used as a Fischer–Tropsch (FT) catalyst at 555 K till a steady-state conversion and stable product distribution was obtained. This used catalyst was then characterized with Mössbauer spectroscopy. These spectra recorded at 4.2 and 300 K, respectively, are shown in Figure 6. A striking difference is observed between these spectra and those of the fresh catalyst (Figure 2). Several new absorption lines in the inner range indicate that new iron species are generated during the CO hydrogenation reaction.

The outer lines of this spectrum at 4.2 K (Figure 6A) (at a distance of 8.13 mm s\textsuperscript{-1}) correspond to a magnetic hyperfine field of 49.5 T, which is commonly found for iron(III) oxides (Table I). Since no relaxation model is presently available for this species, which would allow subtraction of the iron oxide fraction, spectral evaluation has to be done by graphic methods using a tentative assignment for other iron species. The result of this procedure is given in Table II. Three additional sextets can be found that fit the set of hyperfine fields obtained for Hagg carbide (FeC\textsubscript{2}) (subsystems A, B, C, and D of Table II and Figure 6A). There is no other iron carbide reported in the literature with hyperfine field values close to 21.9, 18.3, and 11 T (Table I). It is, however,
well-known that magnetic hyperfine fields of iron carbides are subject to change with the nonstoichiometric change in population of iron nearest-neighbor sites by carbon atoms. As a result and because the actual coordination sphere of the iron atoms in the small particles discussed here may have changed, the existence of other carbide phases (e.g., Fe₃C) cannot be completely excluded.

A third iron phase is considered to be partially responsible for the strong lines at -0.4 and +2.4 mm s⁻¹ in the spectrum (Figure 6A). Provided these lines represent a doublet, they are characterized by an isomer shift of 1.0 mm s⁻¹ and a quadrupole splitting, Vzz, of 2.8 mm s⁻¹. The only reasonable assignment of these species seems to be to Fe(II) cations (Table I). In Fe(II) ion-exchanged Y zeolites, high-spin sixfold-coordinated Fe(II) ions located in the hexagonal prisms show IS values between 1.0 and 1.4 mm s⁻¹ and Vzz between 2.29 and 2.44 mm s⁻¹. Fourfold-coordinated Fe(II) has its IS between 0.9 and 1.1 mm s⁻¹ and Vzz between 0.53 and 0.58 mm s⁻¹ (Table I). The subsystem E in the FT catalyst derived from Na₂O₃/Fe₂O₃ represents therefore sixfold-coordinated Fe(II) ions, possibly located in the hexagonal prisms of the zeolite.

The TEM of the used FT catalyst is shown in Figure 7. In contrast to the fresh catalyst on which no extrazeolitic phase could be observed (Figure 1), a great number of extrazeolitic particles are present now with an average diameter of about 50 nm. Since the Mössbauer spectrum of the small Fe₂O₃ particles remains virtually unchanged during reaction, these external large particles have to be associated with the formation of the iron carbide phase. This is further confirmed by the fact that the carbide sextets in the Mössbauer spectra (subsystems B, C, and D in Figure 6) do not relax when the sample temperature is increased to 300 K, but show on the contrary a decreased value for the hyperfine field (Figure 6B). This indicates that larger particles are present.

It follows, therefore, that during a FT reaction a considerable fraction of the small Fe₂O₃ clusters encaged in a Y-type zeolite are transformed into Fe₃C with 2.0 < x < 2.5. These carbide particles have an average diameter of about 50 nm and are located externally to the zeolite crystals. Another fraction of the Fe₂O₃ phase is reduced to sixfold-coordinated Fe(II) ions. A third fraction of the Fe₂O₃ clusters remains virtually unchanged during this reaction. The formation of iron carbides in general has been a well-discussed subject in relation to Fischer–Tropsch catalysis. Even when the reaction is started with completely

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**Table I**

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Fe₂O₃ Conversion</th>
<th>Co. Distribution</th>
<th>α</th>
<th>Carbon Number Range</th>
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<td>Na₂O₃/Fe₂O₃</td>
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<td>60.0</td>
<td>35.0</td>
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**Table II**

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<th>Measuring Temp/K</th>
<th>Hyperfine Splitting/T (±0.0)</th>
<th>Isomer Shift/mm s⁻¹ (±0.01)</th>
<th>Quadrupole Splitting/Vzz/mm s⁻¹ (±0.01)</th>
<th>Relax. Behavior</th>
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<tr>
<td>A</td>
<td>4.2⁰</td>
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<td>B</td>
<td>25.2</td>
<td>20.7</td>
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<td>-</td>
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<tr>
<td>C</td>
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<td>0.2</td>
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<td>-</td>
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<tr>
<td>D</td>
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<td>E</td>
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<td>11.0</td>
<td>0.2</td>
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<td>-</td>
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<tr>
<td>F</td>
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<td>0.8</td>
<td>2.2</td>
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**Table III**

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<th>Time on Stream/h</th>
<th>Conversion</th>
<th>Co. Distribution</th>
<th>α</th>
<th>Carbon Number Range</th>
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<td>Na₂O₃/Fe₂O₃</td>
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<td>C₃-C₁₆</td>
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</tbody>
</table>

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**Figure 7**

Transmission electron micrograph of sample Na₂O₃/Fe₂O₃ used in a FT synthesis for 72 ks.

**Figure 8**

Hydrocarbon distribution over the different carbon numbers (N) during a FT experiment with sample Na₂O₃/Fe₂O₃ after a time on stream of (a) 0.5 h, (b) 15 h, and (c) 120 h, respectively, and at 555 K.
Figure 9. SF plot of the FT product distributions of Figure 8 after (a) 0.5 h, (b) 15 h, and (c) 120 h time on stream.

Figure 10. SF plot of the FT product distribution at steady state obtained at 555 K for (a) Na-Y/FeO$_2$-RED and (b) Na-Y/FeO$_2$ after a time on stream of 60 and 54 h, respectively.

reduced iron, iron carbide phases similar to the presently encountered ones are formed under FT conditions.

Behavior as Fischer-Tropsch Catalysts of the Fe/Na-Y Zeolites. Pertinent catalytic data for the three associations of iron with zeolite Na-Y are given in Figure 8 and in Table III. All catalysts were tested at medium conversion (Table III) so that considerable overheating of the reactor bed which occurs at high conversions is avoided but also that peculiar reaction selectivities which are sometimes associated with low conversions are not obtained.

Under such conditions it is seen (Table III) that the Na-Y/FeO$_2$ catalyst slowly deactivates at increasing times on stream. More significant are the drastic changes in product selectivity associated with it. These changes are given in Figure 8 as distributions per carbon number fraction or in Figure 9 as a so-called Schulz–Flory (SF) plot. SF kinetics obey the following relation:

$$W_N = N(1 - \alpha)^2\alpha^{N-1}$$

for which $W_N$ corresponds to the product weight fraction with carbon number $N$. The chain-growth probability can be calculated from the slope of a log ($W_N/N$) against $N$ plot. FT distributions sometimes deviate from SF behavior. Since several artifacts may cause this deviation, of which non-steady-state behavior and deficiencies in the mass balance are often encountered, the material balance was checked carefully and found to be correct and the catalyst was followed till a stable product distribution was obtained. It is seen in Figure 8 that for longer times on stream, gradually heavier hydrocarbons appear in the product stream. It is also seen in Figure 9 that the growth factor of the hydrocarbon chains derived for the C$_5$-C$_9$ carbon number fraction increases steadily until ultimately a line with two distinct slopes is found. The latter behavior is typical for the presence of two discrete sets of sites.

Relating this behavior to the physical nature of the catalyst investigated earlier, it can be assumed that the sites responsible for the first growth mechanism ($\alpha_1$) are associated with the zeolite-supported FeO$_2$ phase and those giving the $\alpha_2$ growth rate with the extrazolitic large Hägg-type carbide.

A deviation from SF behavior at higher carbon numbers has also been reported for other iron carbonyls physisorbed on Na-Y. However, the present results show that under steady-state conditions a nonnegligible part of the iron has escaped from the zeolite matrix and consequently the typical zeolite-induced selectivity is partially lost. The other FT catalysts, Na-Y/Fe(0) and Na-Y/FeO$_2$-RED at steady state show perfect SF behavior (Figure 10) with very similar hydrocarbon growth rates (Table III). This is the expected behavior when the physical characterization of these catalysts is recalled: indeed, for both systems the major amount of the iron has escaped from the zeolite and agglomerated as large particles outside the crystals. It can be assumed that formation of a Hägg-type carbide phase mixed with Na-Y crystals occurs during the FT reaction.

Conclusions

In the present paper it is shown that when a Na-Y zeolite containing physisorbed iron pentacarbonyl is exposed to oxygen at 77 K and subsequently is heated to room temperature a highly dispersed FeO$_2$ phase is formed in the zeolite supercages. When this sample is hydrogen reduced at 575 K, most of the iron is sintered into large particles located externally to the zeolite. When physisorbed pentacarbonyl is decomposed in vacuo at 525 K, a low dispersed iron phase located mainly outside the zeolite is formed.

The latter two materials behave as classical Fischer–Tropsch catalysts since the distribution of the hydrocarbons formed follows a Schulz–Flory behavior. The selectivity of the FeO$_2$ phase in the Na-Y catalyst changes continuously until a steady state is reached. Such a steady state is characterized by a hydrocarbon distribution that is obtained by two different hydrocarbon growth rates. Physically, the steady-state catalyst only contains a remainder of the original FeO$_2$ phase in the supercages, sixfold-coordinated Fe(II) ions and a new Hägg carbide phase located outside the zeolite crystals. It is therefore obvious to associate the high hydrocarbon growth rate in the FT reaction with the existence of this bulk carbide. Zeolite-induced FT selectivity seems therefore to exist only in transient conditions, unless migration and agglomeration of iron species during reaction is prevented.

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