Structure and Evolution of Confined Carbon Species during Methane Dehydroaromatization over Mo/ZSM-5

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ABSTRACT: Surface carbon (coke, carbonaceous deposits) is an integral aspect of methane dehydroaromatization catalyzed by Mo/zeolites. We investigated the evolution of surface carbon species from the beginning of the induction period until the complete catalyst deactivation by the pulse reaction technique, TGA, 13C NMR, TEM, and XPS. Isotope labeling was performed to confirm the catalytic role of confined carbon species during MDA. It was found that “hard” and “soft” coke distinction is mainly related to the location of coke species inside the pores and on the external surface, respectively. In addition, MoO3 species act as an active oxidation catalyst, reducing the combustion temperature of a certain fraction of coke. Furthermore, after dissolving the zeolite framework by HF, we found that coke formed during the MDA reaction inside the zeolite pores is essentially a zeolite-templated carbon material. The possibility of preparing zeolite-templated carbons from the most available hydrocarbon feedstock is important for the development of these interesting materials.

KEYWORDS: methane dehydroaromatization, Mo/ZSM-5, hydrocarbon pool, coke, zeolite-templated carbon

INTRODUCTION

Methane dehydroaromatization (MDA) is a promising technology for the direct conversion of methane, the main component of natural gas, to liquid aromatics.1 This reaction has not resulted in a commercial process yet, mainly because the benchmark MDA catalysts such as Mo/ZSM-5 and Mo/MCM-22 quickly deactivate due to coking. Formation of coke over Mo/zeolite catalysts under the MDA conditions cannot be avoided because high temperatures (≥700 °C) are needed for an appreciable conversion of methane to benzene. Under these conditions, polyaromatic hydrocarbons and eventually graphite are more stable than the desired light aromatic products, requiring the use of shape-selective zeolites like ZSM-5 or MCM-22.2,3 Considering the relatively short lifetime of MDA catalysts, several approaches to decrease the coking rate have been investigated. For instance, a small amount of oxygen can be supplied to the reaction to continuously burn the coke species without extensive oxidation of the active Mo phase to Mo(VI). This can be achieved by using either oxygen-permeable ceramic membranes4−6 or short oxygen pulses.7 Complete regeneration of (partially) deactivated MDA catalysts through controlled reaction−regeneration cycling with hydrogen8,9 or oxygen10,11 has also been explored. Despite significant progress in this area, the structure and evolution of the coke species deposited during the MDA reaction remain elusive. Temperature-programmed techniques such as temperature-programmed oxidation (TPO) and thermogravimetric analysis (TGA) are frequently used to characterize the carbonaceous deposits.12 TPO traces of coke formed during the MDA reaction typically contain several features attributed to molybdenum carbide, “soft” or molybdenum-associated coke, and “hard” coke associated with Brønsted acid sites.13−24 There are relatively few spectroscopic investigations of these coke deposits. Wackhuysen et al. performed an extensive XPS study to show that molybdenum carbide, sp2-type aromatic species and hydrogen-poor sp-type species dominate the external surface of spent Mo/ZSM-5.25 Tempelman et al. demonstrated by UV−Raman spectroscopy study that the coke species are polyaromatic hydrocarbons independent of time on stream.26

In a recent publication employing a pulse reaction technique and isotope labeling, we found that polyaromatic carbon species inside the zeolite pores are involved in the formation of benzene during the MDA reaction.27 We established that partially reduced isolated Mo-species catalyze the initial conversion of methane to yet unknown intermediates, which are involved in further reactions with the polyaromatic hydrocarbons eliminating benzene and other aromatics. The involvement of an organocatalytic intermediate resembles the hydrocarbon-pool mechanism for methanol-to-hydrocarbon (MTH) reactions.28 A pronounced difference is that the MDA reaction likely involves radicals, unlike the MTH reaction that proceeds via carbocation chemistry. Figure 1 provides a general scheme of the radical-based hydrocarbon pool mechanism for the MDA reaction.
In this work, we focus on the carbonaceous deposits formed on Mo/ZSM-5 during the MDA reaction at 700 °C using multiple characterization techniques. First, we follow the rapid accumulation of carbon in the micropores of ZSM-5 during the induction period. Isotope labeling with $^{13}$C and D is employed to further underpin the conclusion that confined coke species are actively involved in the MDA reaction. Second, we analyze the evolution of coke species during prolonged MDA reaction by $^{13}$C MAS NMR and DTG and demonstrate the limited ability of TPO-based techniques for studying the chemical speciation of coke. Finally, we characterize the carbon material in spent MDA catalysts after removal of the zeolite by HF etching.

**EXPERIMENTAL SECTION**

**Preparation of Materials.** Supported Mo catalysts were prepared by incipient wetness impregnation of HZSM-5 powder (Si/Al 13, Süd-Chemie, now Clariant) with an aqueous solution of ammonium heptamolybdate tetrahydrate ($\text{NH}_4$)$_6\text{Mo}_7\text{O}_{24} \cdot 4\text{H}_2\text{O}$ (Merck) of appropriate concentration. After impregnation, the catalysts were dried at 110 °C overnight and calcined in air at 550 °C for 6 h with a ramp rate of 2 °C/min. Depending on Mo loading, the catalysts were denoted as 1% Mo (1 wt % of Mo metal loading), 2% Mo, and 5% Mo. Additionally, Mo-containing samples supported on activated carbon (RX-3, Norit), silicalite-1 (prepared according to ref 29), and silica (Grace 643) were prepared by the same method and denoted as Mo/AC (this sample was not calcined at 550 °C), Mo/silicalite-1, and Mo/silica, respectively.

To liberate the carbonaceous deposits from the zeolite, an amount of 1% Mo spent sample was dissolved in an excess of HF (40%) for 2 h. The resulting solid was washed with a copious amount of water on a paper filter and finally dried at 110 °C overnight.

**Catalyst Testing and Characterization.** The prepared catalysts were pelletized, crushed, and sieved to obtain particles with a size of 0.25–0.5 mm. An amount of 0.3 g of sieved catalyst was placed into a tubular quartz reactor (i.d. 4.0 mm, length 250 mm). The catalyst was supported by quartz wool plugs inside the isothermal (700 °C) zone of the reactor. Typically, a 15 mL/min flow of $\text{CH}_4/\text{N}_2$ mixture (95:5) was used, which represents a CH$_4$ WHSV of 2.0 h$^{-1}$. For the isotope-labeling studies, $^{13}$CH$_4$ (99% $^{13}$C, Euriso-Top) and CD$_4$ (99% D, Euriso-Top) were used.

$^{13}$C NMR MAS NMR spectra of coked catalysts were recorded on a 4.7 T Bruker DRX-200 NMR spectrometer operating at 50.3 MHz. The measurements were performed using a 4 mm MAS probe head with a sample rotation rate of 8–10 kHz. Quantitative spectra were recorded using the high-power proton decoupling direct excitation (HPDEC) pulse sequence with an interscan delay of 10 s. The spectra were referenced to the $^{13}$C signal of adamantane at 38.56 ppm. X-ray diffraction (XRD) was performed with a Bruker D2 powder diffractometer system (Cu Kα radiation, scan speed 0.01°/s, 2θ range 5 to 40°). The external surface concentration of Mo species was estimated by X-ray photoelectron spectroscopy (XPS) with a Thermo Scientific K-α spectrometer equipped with a monochromatic Al Kα (1486.6 eV) X-ray source and a 180° double-focusing hemispherical analyzer with a 128-channel detector. Textural properties were analyzed by Ar physisorption carried out at −186 °C using a Micromeritics ASAP-2020 apparatus. Nonlocal density functional theory (NLDFT) was used to determine the pore size distribution of the materials. Prior to the measurements, the samples were outgassed at 400 °C overnight. For the thermogravimetric analysis (TGA) of coke species, a Mettler Toledo TGA/DSC 1 instrument was applied. An amount of ca. 25 mg of spent catalyst was placed in an uncovered alumina crucible, which was heated to 900 °C at a rate of 5 °C/min in a 40 mL/min flow of Ar carrier, MS analysis.

**Figure 1.** Scheme of the hydrocarbon pool MDA mechanism for Mo/ZSM-5. The formation of linear polyaromatic intermediates takes place during the induction stage. Once the hydrocarbon pool is established the formation of light aromatics occurs in parallel with the further growth of polyaromatic species, eventually blocking the zeolite pores.

**Figure 2.** Per-pulse conversion of methane and yields of surface carbon species (estimated by the carbon balance) and benzene during pulsing methane over Mo/ZSM-5 catalysts. Stages of Mo(VI) reduction (activation); extensive formation of surface carbon (induction) and quasi-steady-state benzene production can be distinguished. Additionally Figure S1 shows catalytic performance under flow conditions, and Figure S2 presents the distribution of all detected products during the pulse reaction test. Conditions: 700 °C, 5 mL methane pulses every 200 s, 30 mL/min flow of Ar carrier, MS analysis.
He + 20 mL/min O₂ flow. The morphology of the zeolite and carbon particles was analyzed by scanning electron microscopy (SEM) with a FEI Quanta 200F scanning electron microscope at an accelerating voltage of 3−5 kV using aluminum sample holders. Transmission electron microscopy (TEM) micrographs were obtained with a FEI Tecnai 20 instrument at an electron acceleration voltage of 200 kV.

Modeling of Carbon Species. A structural model of MFI-templated carbon was constructed in Materials Studio 6.0 (Biovia, formerly Accelrys Software, Inc.). The zeolite structure of MFI (silicalite-1) was taken from the internal database. The polyaromatic structure confined inside the silicalite-1 pores was subjected to a geometry optimization using the COMPASS force field at energy and force convergence criteria of 0.1 and 0.5 cal/mol/Å, respectively. The Ewald summation method was employed to account for the electrostatic and van der Waals contributions to the energy. The zeolite framework was removed after this step in order to analyze the pore sizes of the resulting carbon material.

■ RESULTS AND DISCUSSION

Formation of Carbon Species during the Induction Period. We followed the initial stage of the MDA reaction by supplying short pulses of methane to the Mo/ZSM-5 catalyst bed as described before.27

As Figure 2 shows, we can distinguish three reaction stages: (i) activation, involving the reduction of Mo(VI) with CO as main reaction product; (ii) induction, involving formation of a significant amount of surface carbon, which can be linked to an increasing amount of benzene formation; and (iii) quasi-steady-state production of benzene (and other light aromatics). The results show that the formation of surface polyaromatics precedes that of benzene independent of the Mo loading. This sequence of reactions can be explained by the following scheme:

\[
\text{CH}_4 \rightarrow \text{CH}_{x_{\text{surface}}} \rightarrow \text{C}_6\text{H}_6
\]

Further, we used isotope labeling to determine the involvement of surface carbon and proton atoms into the produced aromatic molecules. These experiments were performed by activating and inducing the catalysts with \( ^{13}\text{CH}_4 \) or \( ^{2}\text{CD}_4 \) followed by thorough flushing with Ar to remove any gaseous organic molecules from the system (until complete disappearance of organic MS signals). Finally, pulses of unlabeled CH₄ were supplied to the catalyst bed. Figure 3 shows that the surface carbon and hydrogen atoms are incorporated into the benzene molecules. After activation and induction by \( ^{13}\text{CH}_4 \) pulses, more than 65% of the produced benzene molecules contain at least one \(^{13}\text{C} \) atom during the first \(^{12}\text{CH}_4 \) pulse. The total \(^{13}\text{C} \) concentration in benzene is 20.3%. Upon further \(^{12}\text{CH}_4 \) pulsing, the concentration of \(^{13}\text{C} \) labels in the benzene molecules expectedly decreases. However, even after 10 \(^{12}\text{CH}_4 \) pulses the \(^{13}\text{C} \) concentration is about three times higher than the natural \(^{13}\text{C} \) abundance (1.1%). Although D incorporation was less pronounced than \(^{13}\text{C} \) incorporation, the D content also gradually decreased when more CH₄ was pulsed. Apparent low degree of D incorporation can be explained by a low H/C ratio of confined hydrocarbons and fast H/D exchange of both methane and benzene molecules under applied conditions.
The distribution of benzene isotopologues obtained from the $^{13}$C isotope experiments (Figure 4a) is close to the distribution expected from a random statistical distribution at 20.3% of $^{13}$C. A slight excess of heavy isotopologues $^{13}C_6H_6$, $^{13}C_5^{12}C_1H_6$, and $^{13}C_4^{12}C_2H_6$ and pure $^{12}C_6H_6$ was observed. This observation implies that formed benzene might be in equilibrium with surface carbon species. On the other hand, the concentration of labeled carbon in methane molecules is much lower than that in benzene and naphthalene molecules (Figure 4b). Therefore, the observed isotope exchange is not the result of coke hydrogenation to form labeled methane, which is then further converted to labeled benzene. In other words the following reaction sequence does not significantly contribute to the process:

$$^{13}CH_{\text{surface}} + H_2 \rightarrow ^{13}CH_4$$

$$^{12}CH_4 + ^{13}CH_4 \rightarrow ^{12}C_k^{15}C_{6-k}H_6 + H_2$$

In such a case, the concentration of $^{13}$C label in methane and in aromatic molecules would have to be the same. As the $^{13}$C labels are included selectively into the aromatic molecules, the following overall reaction takes place:

$$^{12}CH_4 + ^{13}CH_{\text{surface}} \rightarrow ^{12}C_k^{13}C_{6-k}H_6 + H_2$$

In this mechanism, mixed benzene isotopologues stem from initial methane activation products (e.g., methyl radicals or ethylene) reacting with the surface species. Previously, we reported that a part of the confined polyaromatic species has a radical character. We also found a correlation between the concentration of organic radicals and the amount of benzene formed during the induction stage. Therefore, we argue that radical pathways likely play an important role in the formation of benzene via surface intermediates.

Pulse reactions and isotope labeling studies showed that the surface carbon formed in the Mo/ZSM-5 pores in addition to the deactivation plays an active intermediate role in the MDA catalytic cycle. The insertion of surface carbon and hydrogen atoms into the benzene molecules likely proceeds via a reaction between the confined polyaromatics and the products of initial methane activation. Under the applied conditions, the contribution of scrambling through reverse hydrogenation of benzene and surface carbon to methane is not significant.

**Evolution of Surface Carbon Species.** The results presented in the previous section are related to the induction period, i.e., the very beginning of the MDA process. In the following sections, we discuss the properties of carbon species formed during the later reaction stages. First, we determined the remaining microporous volume of Mo/ZSM-5 catalysts by Ar physisorption and the total coke content by TG analysis as functions of the time on stream of the MDA reaction. The linear correlation between the amount of coke and the remaining microporous volume (Figure 5) suggests that the carbon is predominantly located on the internal surface. As shown by Okkel et al., the deposition of carbon on the external surface of the zeolite would not result in a significant decrease of microporosity that can be probed by Ar physisorption.

Ar physisorption results demonstrate that carbon species block the pores of 1% Mo and 5% Mo catalysts in a similar manner, independent of the Mo content. In contrast, the results of TG analysis are quite dependent on the Mo loading. Figure 6 shows that increasing the Mo content leads to an increased contribution of “soft” coke at about 450−550 °C. Furthermore, the main combustion feature at 550−600 °C shifts to a lower temperature as the Mo loading increases. In
our previous work we showed that at a Mo loading above 2 wt % a significant fraction of Mo remains as large Mo–oxo clusters on the external surface of fresh catalysts. 11

XPS results (Figure 7) demonstrate that the concentration of Mo on the external surface of 5% Mo catalyst is much higher than on 2% Mo/ZSM-5 and particularly 1%Mo/ZSM-5 catalysts, not only before but also after the MDA reaction. Reduced Mo clusters on the external surface are known to be easily oxidized to MoO$_3$, which can catalyze the combustion of coke and therefore be responsible for the low-temperature DTG feature. Indeed, impregnation of activated carbon (AC) with increasing amounts of ammonium heptamolybdate gradually decreases the temperature necessary to oxidize the carbon from 600 °C (for pristine AC sample) to 450 °C for the 5% Mo/AC sample (Figure 8). Clearly, MoO$_3$ is an efficient combustion catalyst that, if present on the external zeolite surface, can influence the combustion behavior of carbonaceous materials. We argue therefore that the apparent high amount of soft coke on 5% Mo/ZSM-5 is most likely related to the MoO$_3$-catalyzed combustion of carbon species located on/ near the external zeolite surface. Moreover, the heat generated during the combustion of these species results in the shift of the main DTG to lower temperature.

As the majority of carbon species are located in the micropores, XPS cannot be used to characterize these species (Figure S3). To examine the chemical nature and evolution of coke species we performed a $^{13}$C NMR spectroscopy study, applying labeled $^{13}$CH$_4$ for the samples with low carbon content to enhance the quality of spectra. First, we analyzed 5% Mo catalysts exposed to methane for different periods of time and found that the DTG-derived amount of “hard” coke gradually increases with time on stream at the expense of the “soft” coke contribution (Figure 9a). This observation can be explained by a slow diffusion of oxygen and combustion products inside the micropores at increasing carbon content. The NMR spectra of these samples feature only one NMR signal at about 130 ppm (Figure 9b). The intensity of the signal increases with time on stream but the position and relative width remain the same. This signal is assigned to sp$^2$ carbon of aromatic nature. No signals corresponding to sp or sp$^3$ carbon were observed.

A comparison of NMR and DTG data shows that the combustion properties of coke are not related to its structure, as samples with completely different “hard/soft” coke distribution are very similar from the NMR perspective. To underpin this conclusion, we also analyzed spent nonacidic Mo/silicalite-1 and nonmicroporous Mo/SiO$_2$ samples. Spent Mo/SiO$_2$ is characterized by a single low-temperature DTG feature, while Mo/silicalite-1 displays a low- and a high-temperature feature similar to Mo/MFI sample but shifted to lower temperatures (Figure 9c). A stronger contribution of “soft” coke in Mo/silicalite-1 can be explained by the higher external surface concentration of Mo on the SiO$_2$ support that does not contain Bronsted acid sites.32 Despite these differences, $^{13}$C NMR spectra of spent Mo/silicalite-1 and Mo/ZSM-5 are similar (Figure 9d). The $^{13}$C NMR signal in spent Mo/SiO$_2$ is significantly broadened, characteristic of large (pre)graphitic structures, formed on the surface of silica in absence of micropores.33

To summarize, the $^{13}$C NMR and DTG data show that there is no significant chemical difference between the MDA coke burning at low temperature and at high temperature. The low-temperature contribution is related to MoC$_x$ species located on the external surface of spent samples. These species are easily oxidized to MoO$_3$, which is an efficient combustion catalyst. The high-temperature contribution is related to the decreased accessibility of oxygen to the zeolite micropores. The transport of these molecules in the zeolite pores decreases with increasing coke content, resulting in the shift of the combustion features to the higher temperatures and the apparent “hardening” of the coke species.

**Liberation of Carbon Material from the Zeolite.** In order to gain a deeper insight into the properties of carbonaceous deposits formed during MDA, we dissolved a spent (16 h in methane flow at 700 °C) 1% Mo catalyst in HF solution, filtered the remaining solid off, and thoroughly washed it with water. The ash content of the obtained carbon product was below 4% (Figure 10, inset). After liberation of the coke from the zeolite pores, its combustion profile shifted (~100 °C) to a significantly lower temperature (Figure 10). This finding is in line with the conclusion that burning the coke molecules inside the zeolite pores is more difficult due to the presence of coke that blocks diffusion of oxygen.

Electron microscopy analysis reveals that the liberated carbonaceous deposits remain particulate and the size and shape of the primary particles are comparable to the parent carbonaceous deposits...
zeolite crystals (Figure 11). Moreover, since we did not observe any hollowness of the particles by TEM (Figure 11c), it can be concluded that the polyaromatic species (inter)grow throughout the whole zeolite crystal volume.

The analysis of the Ar adsorption isotherm of the liberated carbon material (Figure 12) shows that it possesses a significant microporous volume of 0.07 cm³/g with micropore distribution peaks centered around 5.9, 9.0, and 12.0 Å. The
carbon material also contains mesopores (mesoporous volume of 0.23 cm³/g) and the total BET surface area is 340 m²/g.

The properties of the carbonaceous deposits formed during MDA are similar to those of zeolite-templated carbons (ZTC). These materials can be obtained by high-temperature pyrolysis of organic precursors such as ethylene or furfuryl alcohol inside the zeolite pores, followed by HF dissolution of the zeolite template. The resulting ZTCs are three-dimensional carbon networks displaying varying degrees of order. More generally, ZTCs are porous particulate materials that consist of curved graphene moieties intergrown inside the pores with morphology of particles usually similar to the parent zeolite crystals. ZTCs are promising for numerous applications in catalysis, adsorption, optics, and fuel cells. It should also be mentioned that ZTC materials are known to possess dangling bonds, protected by the zeolite framework. This fact can explain the radical nature of the polyaromatic species confined in ZSM-5, as reported in our previous work.

We modeled the MFI-templated carbon by filling a silicalite-1 model with sp²-hybridized C atoms, placing acenes in the straight channels, connected by phenacene chains via the sinusoidal channels, and pyrene-like moieties in the intersections. The phenacene chains were found to offer the best balance between filling of the sinusoidal channels, the fit and flexibility as compared to other candidate polyaromatics. As a result linear acenes of infinite length lay along the straight channels in the b-direction and slightly twisted picene moieties connect the straight acenes to form a three-dimensional framework (Figure 13). Two types of micropores were found in this model. The first type of pores runs along the b-direction and has dimensions of 8.2 × 12.2 Å. The second type of pores can be viewed along the a-axis and has dimensions of 5.9 × 13.9 Å. These sizes are present in the NLDFT-derived pore size distribution of the liberated carbon material (Figure 12), which confirms the plausibility of the model.

The used model, however, does not account for the structural defects, such as missing linkages and incomplete intergrowth. An important aspect of ZTCs is that their crystallinity depends on the topology of the zeolite template. Highly crystalline ZTC materials with low defect densities and surface areas above 4000 m²/g can be obtained with three-dimensional 12-membered ring (MR) zeolites, such as FAU, BEA, and EMT. Synthesis of ZTC from the 10-MR MFI zeolite results in more disordered materials with no apparent crystalline structure, attributed to steric constraints. Such disorder was also observed in this work, manifested by a high mesoporous volume and moderate surface area. Nevertheless, the NMR, TG, electron microscopy, and Ar adsorption results obtained in this work are with the range of properties previously reported for MFI-templated ZTCs. We expect that the method of direct ZTC synthesis from methane over Mo/zeolite materials can be extended to other zeolite topologies as well. These results show that it is possible to synthesize microporous zeolite-templated carbons directly from methane, which will lead to new possibilities for these interesting materials.

**CONCLUSIONS**

Formation of heavy carbonaceous deposits is an intrinsic property of the MDA reaction. It is highly undesirable as large polyaromatic species block the zeolite pores and lead to the catalyst deactivation. However, these species also play an active intermediate role, as we showed by pulse reaction experiments and isotope labeling. Furthermore, within the sensitivity limit of ¹³C NMR, the structure of coke, formed during MDA, does not depend on time on stream and acidity of the zeolite support. The often reported distinction between “soft” and “hard” coke can be explained solely by the MoO₃-catalyzed combustion of near-surface carbon (“soft” coke feature) and diffusion limitations experienced by molecules of oxygen and combustion products when the zeolite micropores become gradually blocked by the polyaromatic molecules (“hard” coke feature). Finally, the carbon material formed during the MDA after dissolution of the zeolite framework possesses all properties of MFI-templated carbon materials. Conversion of methane over Mo/zeolite catalysts, therefore, offers a direct synthesis route from methane to zeolite-templated carbons.

**ASSOCIATED CONTENT**

Supporting Information

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Figures S1–S3 (PDF)

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Notes

The authors declare no competing financial interest.

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