

Implications of the Molybdenum Coordination Environment in MFI Zeolites on Methane Dehydroaromatisation Performance

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The structure and activity of Mo/Silicalite-1 (MFI, Si/AI = ∞) were compared to Mo/H-ZSM-5 (MFI, Si/AI = 15), a widely studied catalyst for methane dehydroaromatisation (MDA). The anchoring mode of Mo was evaluated by *in situ* X-ray absorption spectroscopy (XAS) and density functional theory (DFT). The results showed that in Mo/Silicalite-1, calcination leads to dispersion of MoO₃ precursor into tetrahedral Mo-oxo species in close proximity to the microporous framework. A

Introduction

Methane dehydroaromatisation (MDA) is a promising route for the valorisation of CH₄ into higher value chemicals as it converts methane directly into aromatics and light hydrocarbons giving H₂ as co-product. Mo-containing ZSM-5 zeolites are among the most studied catalysts for this reaction, with the pore dimensions of the MFI structure being key to provide shape selectivity to benzene (up to 80 %).^[1-6]

It has been traditionally accepted that MDA over Mo/ zeolites occurs *via* a bifunctional mechanism involving two different active sites.^[2,3,7-10] According to this mechanism, molybdenum species constitute the sites responsible for methane activation, forming H₂ as well as C₂H_x and C₃H_x intermediates, which subsequently transform into aromatic

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weaker interaction of the Mo-oxo species with the Silicalite-1 was determined by XAS and DFT. While both catalysts are active for MDA, Mo/Silicalite-1 undergoes rapid deactivation which was attributed to a faster sintering of Mo species leading to the accumulation of carbon deposits on the zeolite outer surface. The results shed light onto the nature of the Mo structure(s) while evidencing the importance of framework AI in stabilising active Mo species under MDA conditions.

products over the zeolite Brønsted acid sites (BAS) associated to framework Al^{3+} . This mechanism was proposed on the basis of several activity studies carried out on Mo-based catalysts prepared using either non zeolitic supports – such as SiO₂ and TiO₂^[11] – or Cs, Ca or Na-exchanged zeolites, with no remaining BAS. While these materials showed low or no selectivity to aromatics,^[12,13] acidic zeolites led to 100–300 times more benzene yield.^[14] Furthermore, Mo₂C and H-ZSM-5 alone were also reported to be poorly active.^[5]

In line with these findings, it has been reported that C₂H₄ conversion on H-ZSM-5 zeolite led to a similar aromatics distribution than that obtained for CH₄ on Mo/H-ZSM-5^[15] whereas Marczewski et al. obtained a direct correlation between the amount of framework Al and aromatic production,^[16] further supporting the idea that the aromatisation step occurs entirely on the Brønsted acid sites of the zeolite. In contrast, titration experiments performed by Tessonier et al. for the quantification of BAS after Mo exchange, revealed that regardless of the amount of BAS left, the catalysts had comparable yield to aromatics.^[17] The authors proposed that the enhanced activity in low Si/Al zeolites was mainly due to a better dispersion of Mo, leading to the formation of highly active isolated Mo species, and that a very few acid sites must be enough to perform the aromatisation of the ethylene intermediates. Interestingly, if the presence of BAS is necessary to produce aromatics from light hydrocarbons as postulated, their absence may allow for a catalyst that is selective to methane coupling products such as ethylene and ethane.

Nonetheless, an alternative monofunctional mechanism has been suggested on the basis that methane aromatisation can occur in the absence of BAS. Although with low conversion and selectivities, aromatisation on MoO₃/SiO₂ has been previously reported.^[18] More recently, Guo *et al.* synthesised a novel Fe@SiO₂ catalyst, wherein isolated iron atoms were embedded



and stabilised in a matrix of amorphous SiO₂.^[19] The authors showed MDA activity and high yields to benzene at 1000 °C. In agreement, catalytic data reported by Kosinov *et al.* for Mocontaining Silicalite-1, the pure siliceous analogue of H-ZSM-5 zeolite, evidenced the formation of aromatics over this catalyst material, leading to the proposal that Brønsted acid sites may not be essential for the aromatisation of methane.^[20] The same group has recently highlighted the involvement of confined carbon in the reaction mechanism instead.^[21,22]

Our previous synchrotron-based operando studies on Mo/H-ZSM-5 evidenced the relevance of Mo speciation in MDA product distribution. X-ray emission showed the selectivity to different hydrocarbon products to be dependent on Mo structure (i.e. partially carburised species were observed during the induction period when light hydrocarbons were produced, whereas fully carburised sites were selective to aromatics).^[23] Xray absorption and diffraction experiments allowed to correlate the Mo species evolution with catalysts deactivation, a major handicap for the commercialisation of MDA route.^[24] This work showed that as a result of complete carburisation, Mo detaches from the zeolite; the subsequent Mo migration from the pores and loss of shape selectivity leads to the formation of carbon deposits on the zeolite outer surface. Accumulation of these deposits block the access of reactants to the active sites resulting in catalyst deactivation.[24]

From the works discussed above it is clear that metal ions such as Mo have a major role in the aromatisation stage than what has been traditionally postulated in the bifunctional mechanism. To date, the extensive research carried out to determine the influence of catalyst support on activity does not include a thorough characterisation to account for differences in metal speciation. Therefore, the aim of this research is to compare the of Mo-containing MFI zeolites in order shed more light onto the role of Mo speciation, framework Al³⁺ and BAS. This is carried out by studying the structure and catalytic activity of Mo/Silicalite-1 with no Al, in comparison with the widely studied Mo/H-ZSM-5 with a silicon to aluminium ratio of 15 and comparable Mo loadings of around 4 wt.%. Notably, the study of a pure siliceous zeolite material containing highly dispersed Mo-oxo species (vide infra) allows us to disentangle the influence of Mo structure and zeolite acidity on the catalytic behaviour largely debated over the past decades. The nature of initial Mo-oxo species is investigated in detail by X-ray absorption spectroscopy (XAS) and density functional theory (DFT), while the physicochemical properties of reacted catalysts are evaluated by characterisation of samples in the first few hours of the MDA reaction.

Results and Discussion

Study of Mo Speciation within Mo-MFI Structures

XAS during in situ Calcination

The Mo K-edge XAS data collected during calcination of the asprepared Mo/Silicalite-1 is shown in Figure 1. The X-ray

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Figure 1. Mo K-edge XAS data collected during *in situ* calcination of Mo/ Silicalite-1 (20% O₂ in He, 700°C, 5°C/min): a) XANES spectra, b) FT-EXAFS (no phase corrected) with vertical lines indicating the peaks assigned to Mo–O and Mo–Mo scattering paths in the MoO₃ precursor.

absorption near edge structure (XANES) spectra (Figure1a) present a pre-edge peak around 20005 eV (dipole-forbidden 1 s \rightarrow 4d electronic transition) and a rising absorption edge at 20015 eV (dipole-allowed 1 s \rightarrow 5p transition). The spectra do not change significantly up to 600 °C showing that the local structure of the MoO₃ precursor (comprising interconnected MoO₆ octahedra) remains fairly stable. Above 600 °C, the pre-edge intensity increases, indicating changes in the symmetry around molybdenum from octahedral to tetrahedral.^[25] In addition, the post-edge region (> 20030 eV) above 600 °C loses its features resulting in a broad peak, suggesting loss of long-range order and dispersion into isolated Mo-oxo species attached to the microporous framework.^[26]

The Fourier transforms of the X-ray absorption fine structure (FT-EXAFS) are shown in Figure 1b. At low temperatures the spectra present features typical of MoO_3 , which contains distorted octahedral MoO_6 units.^[26,27] The scattering from near neighbour O results in two resolved peaks in the FT-EXAFS, with maxima around 1.3 Å (labelled as Mo–O1) and 2.0 Å (Mo–O2). The third peak observed, between 3 and 4 Å, arises



from scattering from neighbouring Mo atoms in agreement with reported Mo-Mo bond distance of c.a. 3.5 Å.^[28] During calcination, a gradual decrease in the peak intensity is observed below 600°C, this decrease is more pronounced for atoms at longer radial distances and can be attributed to an increasing thermal disorder which leads to the damping of the EXAFS oscillations. In line with the XANES results, there are no shifts in the position of the peaks, suggesting that Mo local structure is not significantly altered in this temperature range. Above 600 °C however, sudden changes in the FT-EXAFS indicate the formation of isolated Mo-oxo species attached to the Silicalite-1 framework. The peak at radial distances > 3.0 Å disappear, evidencing loss of Mo-Mo scattering; the peak with maxima around 2.0 Å moves to shorter bond distances, probably corresponding to O atoms bridging to the zeolite, and the peak at lowest radial distances red-shifts, consistent with the formation of short terminal Mo=O double bonds.^[24]

Silicalite-1 contains no framework Al, although the presence of AI as an impurity below the detection limit of ICP-OES cannot be excluded. Unlike Silicalite-1, H-ZSM-5 contains framework Al atoms (Si/Al=15), which provide ion exchange capacity and strong anchoring points for the Mo-oxo species. Characterisation of the acidic properties of the materials here used (NH₃-TPD and FTIR) can be found in Figure S1 in the ESI. In spite of the absence of framework Al, the observed Mo structure evolution into tetrahedral-like Mo-oxo species on Silicalite-1 is analogous to previous reports on acidic Mo/H-ZSM-5.^[23,24,26] For comparison, Figure S2 in the ESI presents the X-ray absorption spectra collected for Mo/H-ZSM-5 during in situ calcination under identical experimental conditions. In agreement with the XAFS data, no MoO₃ crystallites were detected by XRD in the calcined Mo/Silicalite-1 (see Figure S3 in the ESI), indicating that Mo is well dispersed after calcination. Furthermore, a decrease in the micropore volume (14%) was also observed upon calcination (Table 2), supporting the migration of Mo into Silicalite-1 pores. By FTIR a decrease in intensity of the bands corresponding to the silanol groups in Silicalite-1 is seen upon calcination of the physical mixture (Figure S1b), suggesting that metal dispersion may partially occur via interaction of Mo with such groups.

The room temperature FT-EXAFS for calcined Mo/Silicalite-1 and Mo/H-ZSM-5 catalysts are compared in Figure 2. The plot also includes the spectrum of a Fe₂(MoO₄)₃ reference, containing tetrahedral MoO₄ units with an average Mo–O distance of ~ 1.75 Å. Compared to $Fe_2(MoO_4)_3$, Mo/zeolites seem to present a component at lower radial distances, which is consistent with the presence of short Mo=O bonds. $\ensuremath{^{[24]}}$ Interestingly, the two Mo-O distances (i.e. terminal Mo-O and bridging Mo-O bonds to the zeolite framework) in the calcined Mo/Silicalite-1 result in two well resolved peaks, while in contrast, Mo/H-ZSM-5 exhibits a single broad peak. This could be explained by a different degree of interaction between Mo and framework oxygen atoms in both zeolites. Consistent with the absence of framework Al, the interaction of Mo with the pure siliceous Silicalite-1 framework is weaker, resulting in longer Mo-O distances for the bridging oxygen atoms and allowing for better resolution of Mo=O and Mo-O scattering peaks in the FT-EXAFS.

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Figure 2. Mo K-edge FT-EXAFS of Mo/Silicalite-1 and Mo/H-ZSM-5 samples at room temperature after *in situ* calcination together with FT-EXAFS of $Fe_2(MOO_4)_3$ reference with tetrahedral MOO_4 units. Vertical dashed line marks the radial distance for the maxima of the $Fe_2(MOO_4)_3$.

In order to refine the two bond distances, a first shell fit of the EXAFS spectra collected at room temperature after the *in situ* calcination was performed. The best fits were obtained by setting the coordination number 2 for both short and long molybdenum-oxygen bond distances typical of a $(O=)_2Mo$ $(-O-Si)_2$ species. Table 1 presents the parameters refined for Mo/Silicalite-1 and Mo/H-ZSM-5, the experimental and simulated spectra are plotted together in Figure S4 in the ESI. The results suggest similar Mo=O bond distances of 1.69 Å for both samples while the Mo-O distance is considerably longer on the pure siliceous zeolite (i.e. 2.31 Å in Mo/Silicalite-1 vs 1.80 Å in Mo/H-ZSM-5).

Density Functional Theory

In order to verify these observations, DFT calculations were performed, allowing to gain further insight into the interaction of Mo with the Silicalite-1 support. These were concerned with the optimisation of isolated di-coordinated Mo-oxo structures

Table 1. EXAFS fitting parameters of the Mo/Silicalite-1 and Mo/H-ZSM-5 spectra acquired at RT after calcination: So²=0.91, Fit Range: 3 < k < 12, 1 < R < 3. Where CN=coordination number, R=bond length of the Absorber-Scatterer, σ^2 =Mean squared disorder term (sometimes referred to as the Debye Waller factor), E_{σ} =Energy shift, R_{Factor} =A statistic of the fit, which is a way of visualising how the misfit is distributed over the fitting range.

Sample	Schell	CN	R [Å]	σ² [Ų]	Eo	R _{Factor} [%]
Mo/ Silicalite-1	Mo=0	2	1.69	0.0046 (+/-0.0012)	11.93 (+/-2.03)	3.5
	Mo–O	2	2.31 (+/-0.03)	0.0079 (+/-0.0030)		
Mo/ H-ZSM-5	Mo=O	2	1.69	0.0059 (+/-0.0043)	-0.15 (+/-2.61)	3.2
	Mo–O	2	1.80 (+/-0.04)	0.0041 (+/-0.0016)		



with tetrahedral geometry, as suggested by the above X-ray absorption results. The fully geometry optimised structures, shown in Figure 3, comprise pseudo-tetrahedral [MoO2]²⁺ sites attached to the zeolite walls through two Mo-OF bonds to oxygen atoms on oxygens adjacent to the T5 site; they also contain two terminal Mo=O bonds. In the case of Mo/Silicalite-1 (Figure 3a) the bond lengths obtained are 1.73 Å and 2.33 Å for Mo=O and Mo-O_F respectively. These values are comparable to those refined by EXAFS (1.69 Å and 2.31 Å). Note that the Mo–O_F length could be considered too long to be a real bond; nonetheless, the Mulliken charges on the two framework oxygens 'bonded' to Mo are more negative than charges on other framework oxygens, (-0.55 compared to a mean charge of -0.45 for the remaining framework oxygens). This indicates that an electrostatic interaction exists, and that charge transfer occurs between framework oxygens and Mo.

For Mo/H-ZSM-5 (Figure 3b), while the Mo=O bond distance is equivalent to the Mo/Silicalite-1, the Mo–O_F bond obtained is significantly shorter with a value of 1.8 Å. This value is comparable to our EXAFS results in Table 1 as well as previous theoretical calculations on Mo/H-ZSM-5.^[29] The shorter distance compared to Silicalite-1 is presumably due to strong interactions between the Mo-oxo species and the aluminosilicate framework.

By FTIR a range of silanol defect groups were found to exist in the Silicalite-1 sample that may serve as anchoring point of Mo-oxo species. These defects include vicinal, geminal and large number of silanol nests (see Figure S2 for the assignments). Previous DFT studies have shown variations in the stability Mo-oxo species anchored through different silanol



Figure 3. The fully geometry optimised form of the Mo-oxo species bound to Silicalite-1 (a) and H-ZSM-5 (b). The bonding to Silicalite-1 through silanol defects is modelled across the nests (c), vicinal (d) and non-vicinal silanol defects (e). In all structures $[MOQ_2]^{2+}$ contains two terminal Mo=O bonds and is di-coordinated by two short Mo–OF bonds. The framework is shown as stick bonds, Mo is represented in purple, oxygen in red and hydrogen in pink.

defects of an amorphous silica support.^[30] To evaluate the effect of silanol defects on the binding of the initial Mo-oxo to the walls of the zeolite framework, a range of silanol-containing Mo/Silicalite-1 structures were examined. As geminal sites have been reported to be highly unstable due to their rigidity,^[30] only vicinal and non-vicinal sites are considered.

As indicated in the experimental section, the presence of silanol nests were simulated through the removal of a single Si atom from the T5 sites. The four dangling Si–O bonds generated by the introduction of the vacant site were satisfied by the presence of hydroxyls. [MoO2]²⁺ was then di-coordinated across the now vacant T5 site, with the removal of two protons, simulating binding across two terminal silanol defects *via* a condensation reaction. The fully geometry optimised form of this structure is shown in Figure 3c. This structure contains equivalent Mo=O bond lengths of 1.69 Å; Mo–O_F bond lengths however are shorter than those reported for the Mo-oxo species supported on silanol-free Silicalite-1 (in Figure 3a) with one bond slightly longer than the other, Mo–O_F = 1.90 Å and 1.88 Å.

The fully optimised structures of [MoO₂]²⁺ bound across vicinal and non-vicinal silanol groups are shown in Figure 3d and 3e respectively. In these structures, Si has been removed from the T5 and T9 sites and the resulting dangling bonds are satisfied by hydroxyls, simulating the presence of two silanol nests per MFI unit cell. The calculations suggest that bonding to vicinal OHs is more stable than non-vicinal by 52 kJmol^{-1} , contrary to the work by Guesmi et al.^[30] The bond lengths in these structures are Mo=O = 1.70 Å and Mo- $O_F = 1.90$ Å and 1.88 Å for binding across the vicinal site and Mo=O = 1.70 Å and $Mo-O_F = 1.97$ Å and 2.08 Å. Mulliken population analysis suggests that Mo is interacting with both framework oxygens, hence it appears that binding at vicinal sites is preferred as a consequence of stronger, more equal coordination to the framework. The DFT calculations for Mo attached to Silicalite-1 through silanols result in significantly shorter Mo-OF bond lengths (1.88 to 2.08 Å) compared to the ones obtained by EXAFS fitting (2.31 Å). In contrast, the model with no silanols in the structure is closer (Mo– O_F = 2.33 Å) to the fitted values. This suggests that the Mo-oxo species are not bound exclusively across vacancy sites in the Silicalite-1 framework and that interaction and charge transfer occurs between framework oxygen atoms and Mo.

Methane Dehydroaromatisation

Catalytic Activity

To compare the performance of Mo/H-ZSM-5 and Mo/Silicalite-1, MDA reactions were carried out for a period of 10 h, with the outlet gas composition measured by MS. The data corresponding to the main reaction products (i.e. H_2 , C_2H_x and C_6H_6) are presented in Figure 4. Both catalysts exhibit an induction period during the first minutes of reaction where H_2 formation is seen but no hydrocarbon production is detected (data from earlier reaction times are shown in Figures 4a and 4b). This induction period is coincident with the evolution of CO, CO₂ and H₂O (see





Figure 4. MS profiles of the products formed during the methane dehydroaromatisation reaction (50 % CH₄/Ar, 1500 h⁻¹, $700 \degree$ C). First 60 min for Mo/Silicalite-1 (a) and Mo/H-ZSM-5 (b) with vertical lines marking the end of the induction period. 10 h reaction data for Mo/Silicalite-1 (c) and Mo/H-ZSM-5 (d).

Figure S5 for the MS trends of combustion products) and it is accepted that during this stage tetrahedral Mo-oxo species undergo reduction and carburisation leading to the formation of active species for dehydroaromatisation. After the initial induction period, the aromatisation stage was seen to commence for both catalysts evidenced by the evolution of C_6H_6 , together with light hydrocarbons and H_2 . In agreement with Kosinov *et al.*,^[20] the observed formation of C_6H_6 on Mo/ Silicalite-1 suggests that BAS are not required for aromatisation, pointing out to this being also an intrinsic property of molybdenum carbide species.^[5]

A close analysis of the MS data reveals differences in MDA performance of both catalysts. Interestingly, a shorter induction period is observed for Mo/Silicalite-1 (6.0 min) compared with the Mo/H-ZSM-5 (8.5 min) which suggests a faster carburisation of initial Mo-oxo species on the pure Si zeolite. Regarding the aromatisation stage, data over the 10 h of reaction (Figures 4c and 4d) show that C₆H₆ and H₂ gradually decreased for Mo/H-ZSM-5 indicating a gradual catalyst deactivation. For Mo/Silicalite-1 however, C₆H₆ and H₂ were seen to steeply decrease within the first ~3 h of reaction, evidencing a significantly faster deactivation. In addition, the TGA analysis (Table 2) performed for catalysts reacted for different times revealed increased amounts of carbon deposits on Mo/Silicalite-1.

These activity differences can be explained by the degree of interaction of Mo with the zeolite framework as discussed in the

Table 2. N ₂ physisorption	results	and	carbon	content	of	Mo/MFI	catalysts
at different reaction times.							

Sample	Mo [wt. %]	TGA-Carbon 350–600 °C [wt. %]	S _{BET} [m²/g]	V _{micr} [cm ³ /g]
H-ZSM-5	/	/	412	0.15
Mo/H-ZSM-5 calc.	3.80	/	344	0.12
Mo/H-ZSM-5(7 min)	/	0.4	351	0.12
Mo/H-ZSM-5(90 min)	/	3.0	326	0.11
Mo/H-ZSM-5(10 h)	/	6.3	280	0.09
Silicalite-1	/	/	452	0.17
Mo/Silicalite-1 calc.	3.62	/	389	0.14
Mo/Silicalite-1(7 min)	/	0.5	409	0.14
Mo/Silicalite-1(90 min)	/	5.4	333	0.12
Mo/Silicalite-1(10 h)	/	7.4	302	0.11

EXAFS and DFT studies above. In the Mo evolution mechanism proposed from our previous *operando* studies, during the induction period the Mo-oxo species present after calcination are fully carburised; this occurs by the consecutive replacement of terminal and bridging O atoms by C.^[23,24] The weaker interaction of Mo with Silicalite-1 compared with H-ZSM-5 would potentially facilitate the carburisation of bridging O therefore shortening the induction period. Such weaker interaction would also lead to a faster sintering of Mo species and their migration to the outer surface under reaction conditions. In line with the deactivation mechanism previously proposed



by our group,^[24] the shape selectivity to aromatics provided by the MFI pore structure is lost as a consequence of this migration, favouring the formation of large carbonaceous deposits in the zeolite outer surface. Hence, a faster sintering of the active species in Mo/Silicalite-1 would explain the increased formation of carbon deposits in this catalyst - as observed by TGA, and thereby the faster deactivation.

We note that aromatisation over BAS is believed to occur via acid catalysed heterolytic reactions (i.e. oligomerisation, β -scission, hydride transfer, protolysis, alkylation, dealkylation etc),^[9,10] while a radical-mediated reaction mechanism is more plausible for MDA over an acid-free catalyst. Variations in reaction mechanism over Mo/H-ZSM-5 and Mo/Silicalite-1 as a result of the different BAS content could also possibly contribute to the differences in the carbon deposition profile observed.

Post Reaction Characterisation

In order to gain insight into the fast deactivation of Mo/ Silicalite-1, the reacted catalysts (i.e. after 7 min, 90 min and 10 h of reaction) were further characterised.

Table 2 displays the textural properties and the carbon content measured by TGA. Note that the Mo content (determined by ICP for the calcined samples) is similar for both catalysts. As compared to the parent zeolites, the micropore volume of the calcined Mo-containing catalysts was seen to significantly decrease; this decrease is similar for Mo/Silicalite-1 (14% decrease) and Mo/H-ZSM-5 (16% decrease) suggesting a comparable Mo dispersion. As discussed in the XAFS section above, the drop in micropore volume is probably driven by the migration of Mo into the zeolite pores during calcination to form Mo-oxo species. From 7 min to 10 h of reaction, the micropore volume was also seen to decrease as a result of carbon deposition. The increase in the amount of deposited carbon is clearly evidenced by TGA, responsible for the gradual catalyst deactivation observed in Figure 4. The derivatives of the TGA curves at 7, 25 min and 10 h of reaction (Figure S6 in the ESI) also exhibit a gradual increase of peak intensity and the combustion temperature with reaction time, in line with increased carbon deposition as the reaction advances. As shown in Table 2 the amount of carbon deposited on Mo/ Silicalite-1 was always higher than for Mo/H-ZSM-5 consistent with the faster deactivation observed when the pure siliceous zeolite is used as the support.

In order to gain knowledge on the nature of carbon deposits formed during reaction, Raman spectra was acquired for the catalysts reacted for 90 min. Figure 5 shows the first order Raman bands for Mo/Silicalite-1 and Mo/H-ZSM-5, which present two distinct bands typical for carbon compounds. The band around 1360 cm⁻¹ is denoted as D1 (disordered) band and it is ascribed to in-plane breathing vibrations of sp²-bonded carbon (rings). The D1 band is usually attributed to amorphous carbon, carbon nanoparticles or defects in graphitic-type deposits.^[31] The second band is known as the G (graphitic) band, it usually appears at 1580 cm⁻¹ and corresponds to in-

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Figure 5. Raman spectra of Mo/Silicalite-1 and Mo/H-ZSM-5 reacted for 90 min (50 % CH₄/N₂, 700 °C, 1500 h^{-1}).

plane stretching vibrations of pairs of sp² C atoms. This band is observed in graphitic-type carbon as a result of lattice vibrations. $^{\scriptscriptstyle [31-33]}$ In both catalysts these bands are shifted to higher wavenumbers (1611 cm⁻¹). This shift has been reported to be due to a contribution from a second D2 band (1620 cm^{-1}) attributed to edges of graphitic crystallites.[33,34] Thus, the observed shift suggests the presence of very small carbon crystallites with large surface/bulk ratio and therefore a high number of edges.^[33,35] A weak shoulder can be also observed at 1200 cm⁻¹, this Raman shift has been named as a D4 band and it is ascribed to the stretching vibrations of sp²-sp³ C-C bonds in conjugated aliphatic species.^[36] The ratio of D1 and G(D2) bands intensities gives insight regarding the degree of order in the carbon structure,^[31,33] increasing I(D1)/I(G) indicates increasing structural disorder. The higher D1/G value for Mo/Silicalite-1 (0.61) than for Mo/H-ZSM-5 (0.52) indicates that the carbon deposited on the catalyst with no framework Al is more amorphous in nature. Hence, Raman studies confirm the formation of significant amounts of disordered graphitic carbon or small graphite crystallites in the spent samples, with lower degree of crystallinity for Mo/Silicalite-1. From our Raman or TGA data it is difficult to discern between carbon deposits located inside the zeolite channels or on the outer surface. Our previous PXRD studies on Mo/H-ZSM-5 however, suggests that most of the carbon is located in the external surface of the zeolite crystals.^[24]

Microscopy studies (TEM and SEM) were also carried out for reacted catalysts in order to evaluate the Mo sintering process during MDA. High resolution backscattered electron images for Mo/Silicalite-1 reacted for 7 min, 90 min and 10 h are shown in Figure 6 and illustrate the sintering process of Mo species under reaction conditions. The images contain Silicalite-1 crystals of around 200 nm, whereas the observed bright spots are, based on signal intensity, due to molybdenum carbide.^[24] Note that this analysis was done using a low accelerating voltage of





Figure 6. SEM-ESB (Energy Selected Backscattered) electron images (accelerating voltage 1.6 eV) of Mo/Silicalite-1 after 7 min (a), 90 min (b), and 10 h (c) of reaction (50% CH₄/N₂, 700 °C, 1500 h⁻¹).

1.6 eV; hence, the Mo distribution observed corresponds to the outermost surface of the zeolite crystals.

Very few bright spots are present after 7 min of reaction (Figure 6a) suggesting a low degree of molybdenum carbide sintering towards the zeolite surface at this stage of reaction.

a)

The 90 min reacted sample (Figure 6b) presents an increased number of molybdenum carbide particles whereas after 10 h (Figure 6c), the number of these particles increases further such that all zeolite crystals possess sintered molybdenum carbide on the surface. A similar trend is also observed for Mo/H-ZSM-5, in line with previous publications (see Figure S7 in the ESI for equivalent images in Mo/H-ZSM-5).^(8,23)

TEM images were also taken at 90 min of MDA where the steep deactivation of Silicalite-1 is observed in contrast of more stable activity of Mo/H-ZSM-5. As shown in Figure 7a Mo/ Silicalite-1 presents large molybdenum carbide particles of around 15 nm diameter, while Mo/H-ZSM-5 (Figure 7c) exhibits significantly smaller particles (> 5 nm). After 10 h of reaction extensive sintering was observed for both catalysts. As expected, the EDX maps (Figure 7b and 7d) reveal Mo to be associated with C, consistent with the presence of molybdenum carbide particles. An even distribution of Si and O were found



c)

d)

b)



Figure 7. STEM-DF image of the catalysts after 90 min of reaction (50% CH_4/N_2 , 700 °C, 1500 h⁻¹). Mo/Silicalite-1 (a) and Mo/H-ZSM-5 (c); and 10 h reacted STEM-DF with the corresponding EDX elemental maps of Mo/Silicalite-1 (b) and Mo/H-ZSM-5 (d).

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in the zeolite crystals, while the Al distribution in Mo/H-ZSM-5 presents brighter spots probably due to presence of extra framework aluminium or Al molybdate particles. Interestingly, in the C elemental map of Mo/Silicalite-1, which contains a higher amount of C, a more intense carbon signal can be distinguished on the periphery of the Silicalite-1 particles (Figure 7b). This indicates a significant accumulation of carbon deposits on the outer surface of the zeolite.

These results are in agreement with the catalytic activity data and deactivation mechanism discussed in Figure 4. The weak interaction between initial Mo species and Silicalite-1 observed by EXAFS and DFT is a key factor for the enhanced sintering of the Mo active species seen by TEM. The migration of Mo carbides to the outer zeolite surface leads to an increased carbon deposition on the zeolite periphery and thereby, a more rapid deactivation of Mo/Silicalite-1.

Summary and Conclusions

This study compares the properties of Mo/zeolites with MFI structure using zeolites with and without framework AI (H-ZSM-5 and Silicalite-1 respectively). The structural refinement of the initial Mo-oxo species by XAS and DFT, followed by MDA activity studies and characterisation of reacted catalysts bring new insights into the role of framework AI on the catalytic performance.

XAS data collected during *in situ* calcination of Silicalite-1 and MoO₃ physical mixture suggest that upon calcination MoO₃ migrates into zeolite pores, leading to the formation of tetrahedral Mo-oxo species. The structure of these species (i.e. with two terminal Mo=O and two bridging Mo–O groups attached to the zeolite framework) appears to be analogous to the ones previously reported for ion-exchanged Mo/H-ZSM-5, with the BAS acting as anchoring sites for Mo. Longer Mo–O distances to the framework oxygens on Silicalite-1 however allude to a weaker interaction of Mo with this support. DFT calculations are in agreement with the experimental refinement, suggesting that Mo in Mo/Silicalite-1 it is not exclusively bound across silanol defect sites of the Silicalite-1 support and attaches to the zeolite walls through oxygen atoms *via* electrostatic interaction.

The formation of analogous tetrahedral Mo-oxo sites upon calcination catalysts allows us to compare the catalytic activity of both catalysts and to discriminate between the roles of Mo structure, framework AI and BAS on MDA reaction. The activity studies showed benzene formation in both catalysts. In contrast with the traditionally postulated bifunctional mechanism, the aromatisation observed for Mo/Silicalite-1 indicates that the presence of BAS is not essential for the formation of benzene. Therefore, molybdenum carbide species seem to promote aromatisation provided that good initial Mo dispersion into the zeolite pores ensures the shape selectivity to aromatics. These results are in agreement with Kosinov *et al.* who proposed that the conversion of methane to benzene might take place in a monofunctional manner on Mo carbides^[20] in the zeolite micropores, or mediated by carbonaceous species.^[21,37] Such mecha-

nisms open the possibility to optimise active MDA catalysts based on non-acidic zeolites. These supports exhibit enhanced thermal stabilities that represent an advantage for the MDA reaction, which operates > 700 °C and is equilibrium-limited to $\sim 14\%$ at 700 °C, allowing to increase the reaction temperature for improved yields.

While the presence of BAS associated to framework AI may not required for aromatisation to occur, our research suggest that Al plays an important role on the stabilisation of Mo species. The shorter induction period for Mo/Silicalite-1 alludes to a more rapid carburisation of the initial Mo-oxo species. In line, a faster Mo sintering under reaction conditions and severe catalyst deactivation is also seen for this catalyst. These observations can be attributed to the weaker interaction of Mo species with the purely siliceous Silicalite-1 evidenced by EXAFS and DFT. As proposed in our previous studies, such carburisation and sintering comprise the migration of Mo to the zeolite outer surface; this leads to the loss of shape electivity to aromatics provided by the pores and extensive accumulation of bulky carbon deposits in the zeolite outer surface. While variations in the reaction mechanism due to different acidic properties of Mo/H-ZSM-5 and Mo/Sillicalite-1 could also contribute to the different carbon deposition profile observed, our data suggests that the instability of Mo active sites on Silicalite-1 is the main cause of catalyst deactivation. The characterisation carried out for reacted Mo/MFI samples suggests a more pronounced sintering of molybdenum on Silicalite-1. The increased accumulation of carbon deposits is concluded from the N₂ physisorption and TGA; Raman spectra indicate the deposits consist of disordered graphitic carbon, with a lower degree of crystallinity for Mo/Silicalite-1. TEM-EDX maps on 10 h reacted Mo/Silicalite-1 show high C concentration in the zeolite outer surface.

The structure and location of Mo species clearly have a strong impact on catalytic performance. In this regard, framework Al seems to contribute to the stabilisation of Mo species resulting in a delayed catalyst deactivation. In this sense promising engineering solutions are being investigated by several groups to remove carbon deposits (i.e. through catalyst regeneration cycles or by using membrane reactors).[4,15,38-41] Nonetheless, an alternative approach to prevent deactivation would comprise the optimisation of the catalyst formulation, focused on further improving the stability of the active metal species inside the zeolite pores. In this line, follow up investigations should focus on enhancing the interaction of the active metal and the support on MDA catalysts. For instance, with the use of different metals as the active species in MDA (i.e. Fe), by the addition of promotors to the catalyst that could lead to less mobile active species, or by the optimisation of embedding or encapsulating methods to fix the active species and prevent migration and sintering.^[24,42,43]



Experimental Section

Catalyst Preparation

Silicalite-1 (MFI structure, Si/Al = ∞), was prepared by hydrothermal synthesis as described by Lobo *et al.*^[44] and subsequently calcined at 550 °C. The resulting zeolite was then treated with ethylenediamine following the procedure reported by Wang *et al.*^[45] in order to generate silanol groups in the structure by the extraction of framework Si⁴⁺. ZSM-5 (also MFI structure, Si/Al = 15) was purchased from Zeolyst (CBV3024E) in its ammonium form. The proton form was obtained by calcination in static air at 550 °C for 6 h, the heating rate used was 2 °C/min.

Mo/Silicalite-1 and Mo/H-ZSM-5 with metal loadings of ~4 wt.% were prepared by first mixing the zeolite and MoO₃ (Sigma, 99.95%) powders in an agate mortar for 0.5 h. The resulting physical mixtures are referred to as the as-prepared catalysts. The samples were then calcined in air to 700 °C for 0.5 h with a 5 °C/min heating rate.

Catalyst Characterisation

Fourier-transform infrared spectroscopy (FTIR) spectra were recorded in a Nicolet iS10 spectrometer. Samples were pressed into self-supporting wafers with a density of c.a. 10 mg/cm². The wafers were dried prior to the measurements by heating them up to 285°C for 3 h under 70 ml/min He flow. After dehydration, the sample was cooled down to 150°C under dry He before spectra collection. Temperature programmed desorption of ammonia (NH₃-TPD) was performed using an AutoChem II 2920 micromeritics instrument equipped with a moisture trap and a thermoconductivity detector. Samples were first preactivated by flowing pure N_2 and heating up to 550 °C for 30 min (5 °C/min). The reactor was then cooled down to 100°C for ammonia adsorption which was run by flowing 1 % NH₃/N₂ until saturation (~1 h). Next, pure N₂ was flowed for 2 h to remove physisorbed ammonia on the sample. Finally, ammonia desorption was carried out by increasing the temperature up to 1100 °C with a heating rate of 10 °C/min. Powder X-ray diffraction (XRD) patterns were recorded using a Rigaku SmartLab X-Ray Diffractometer fitted with a hemispherical analyser. The measurements were performed using Cu K α radiation source $(\lambda = 1.5406 \text{ Å})$ with a voltage of 40 kV, and a current of 30 mA. The patterns obtained were compared to crystallographic data in the reference library (ICSD database). Elemental analysis was carried out using inductively coupled plasma optical emission spectroscopy (ICP-OES) in a Perkin Elmer Optical Emission Spectrometer Optima 3300 RL. Nitrogen physisorption measurements were performed on a Quadrasorb EVO QDS-30 instrument at 77.3 K. The samples were outgassed at 350 $^{\circ}$ C overnight under vacuum prior to N₂ sorption. The Brunauer-Emmett-Teller equation was used to calculate the specific surface area in the pressure range $p/p_0 = 0.0006 - 0.01$. The micropore volume was calculated from the t-plot curve using the thickness range between 3.5 and 5.4 Å. Thermogravimetric analysis (TGA) was carried out in a TA Q50 instrument. Samples were heated up to 950°C using a rate of 5°C/min under an air flow of 60 mL/ min. Raman spectra were acquired on ULTRA at the Central Laster Facility.[46,47] The measurements were carried out using 400 nm laser to excite the sample (placed in quartz window holders). Toluene impregnated H-ZSM-5 was used for calibration of the detected signals. Scanning electron microscopy (SEM) analysis was performed using a Zeiss ultra 55 Field emission electron microscope. Compositional analysis and low-resolution imaging were carried out with accelerating voltage of 20 kV, 30-60 µm aperture and 7-8 mm working distance. High-resolution images were also taken with an accelerating voltage of 1.6 kV, 20–30 μm aperture

and 2–3 mm working distance. Transmission electron microscopy (TEM) measurements were performed in a JEM 2800 (Scanning) with a voltage of 200 kV and aperture of 70 and 40 μ m. Bright-field imaging mode was performed using a CCD in high magnification, whereas lattice resolution imaging mode was carried out using CCD Dark-field (Z-contrast) imaging in scanning mode using an off-axis annular detector. The secondary electron signal was acquired simultaneously with the other TEM images providing topological information of the sample. Compositional analysis was performed by X-ray emission detection in scanning mode.

X-ray Absorption Spectroscopy during in situ Calcination

XAFS studies on Mo/MFI zeolites were carried out at the B18 beamline at Diamond Light Source^[48] in Harwell, United Kingdom. The storage beam energy was 3 GeV and the ring current 300 mA. Mo K-edge spectra (in the range of 19,797 to 21,000 eV) were collected in transmission mode using ion chamber detectors with a fast scanning Si (111) double crystal monochromator, with a Mo foil placed between It and Iref. X-ray beam dimensions at the sample position were $1 \times 1 \text{ mm}^2$, whereas the acquisition of each spectra took ~60 s. In situ experiments were performed using a setup developed by A. B. Kroner et al.^[49] For the experiments, 40 mg of the as-prepared catalysts (sieve fractions: 0.425-0.150 mm) were placed in a 3 mm diameter quartz capillary. The sample was then heated to 700°C for 30 min using a hot gas blower, under 20% O_2 in He flow (GHSV = 3000 h⁻¹ and heating rate = 5 $^{\circ}$ C/min). After 30 min, the sample was cooled to room temperature under pure He. XAS data was collected during the heating ramp as well when back at room temperature after calcination.

XAS data analysis was performed using the Demeter software package.^[50] The X-ray absorption fine structure (EXAFS) fitting to model spectra was done using an amplitude reduction factor of 0.91, which was obtained by fitting the Mo foil reference to crystallographic data from the ICSD database. K range values used in the fitting were between 3 and 11 Å⁻¹ whereas the R range spanned 1 to 3 Å were used. The Mo K-edge edge position was taken as the energy at half-step height and the Fourier transformed (FT) EXAFS data presented are not phase corrected.

Density Functional Theory

Based on the outcome of the EXAFS studies on Mo/zeolites, $^{\rm [24]}$ only monomeric species were considered in the calculations, which were performed at 0 K.

Firstly, geometry optimisation (all atom positions and cell parameters were allowed to vary) of a single orthorhombic MFI unit cell with Si/Al $= \infty$ (i.e. Silicalite-1) was undertaken. The initial cell was obtained from the IZA zeolite structure database,^[51] and optimised using the periodic DFT code CP2K^[52] at the PBE^[53] level of theory with a DZVP-MOLOPT basis. To give a reasonable starting geometry for monomeric [MoO₄]²⁻, molybdate ([MoO2]²⁺ is known to have a similar O-Mo-O bond angle to [MoO₄]²⁻) was geometry optimised as an isolated cluster using the aperiodic Orca code (PBE functional and SVP basis). Then, the optimised [MoO2]²⁺ moiety was grafted to two framework oxygen atoms (O_F) across the T5 site forming [MoO4]²⁻. Several trial positions were optimised and the results described here relate to the global minimum established in each case. This structure was geometry optimised at the same level of theory as the guest-free Silicalite-1 framework (i.e. PBE and the DZVP-MOLOPT). The interaction of Mo species with different possible types silanols - i.e. nests, vicinal and geminal, was also simulated. For the nests, Si was removed from the T5 sites in



optimised Silicalite-1 structure charge compensating the four dangling Si–O bonds generated with terminating protons. The resultant structure was then geometry optimised at the same level of theory used throughout. Following this, $[MOO_2]^{2+}$ was reintroduced to the same position as before, removing the two protons from the framework oxygens bound to Mo in order to mimic the condensation reaction that is proposed to occur when the initial mobile Mo-oxo species. Vicinal and non-vicinal silanol defects were simulated by removing Si from the T5 and T9 sites in both the vicinal and non-vicinal case. $[MOO_2]^{2+}$ was then grafted across the defects, removing two protons to mimic condensation reactions. The two structures were geometry optimised at the same level of theory used throughout these simulations.

Finally, the Mo/Silicalite-1 models were compared with that of Mo/ ZSM-5 by introducing a single AI to each of the 12 crystallographically distinct framework T-site present in orthorhombic MFI. Brønsted acidic protons were not included in these simulations in order to mimic the liberation of water which is proposed to occur on binding of the initial Mo-oxo species to the walls of the framework. The Mo-oxo species in H-ZSM-5 remained in the same position, grafted across the T5 site.

MDA Catalytic Performance Tests

0.6 g of sieved catalyst (150–425 μ m sieve fractions) was introduced into a tubular quartz reactor (internal diameter = 0.7 mm). The sample was fixed in the isothermal zone of the oven by quartz wool. A total gas flow of 30 mL/min was fed by means of mass flow controllers (GHSV = 1500 h⁻¹).

The as-prepared physical mixtures were first calcined under 20% O₂/He flow by heating up to 700 °C for 30 min using a heating rate of 5 °C/min. After flowing pure Ar for 30 min, methane dehydroar-omatisation reaction was initiated by flowing 50% CH₄/Ar. The reaction products were analysed by an online mass spectrometer (OmniStar GSD 32001). All MS data intensity presented in this manuscript are normalised to the Ar signal and are plotted in logarithmic scale.

To follow the deactivation process, Mo/MFI samples were collected at different reaction times (7 min, 90 min and 10 h) after cooling down the reactor under Ar flow. The resulting samples are denoted by adding the reaction time after the sample code. For example, Mo/Silicalite-1 sample reacted for 90 min is identified as Mo/ Silicalite-1(90 min).

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Conflict of Interest

The authors declare no conflict of interest.

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Coordination in Zeolites: X-ray absorption spectroscopy and density functional theory calculations were performed to investigate the Mostructure on Mo/Silicalite-1 (Si/ $AI = \infty$) and Mo/H-ZSM-5 (Si/AI = 5) catalysts. These studies combined with catalytic activity measurements bring new insight to unravel the roles of Mo speciation, framework AI and Brønsted acid sites on the methane dehydroaromatisation reaction.



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Implications of the Molybdenum Coordination Environment in MFI Zeolites on Methane Dehydroaromatisation Performance