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# Structure and support effects on the selective oxidation of dimethyl ether to formaldehyde catalyzed by $MoO_x$ domains

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#### Abstract

The selective oxidation of dimethyl ether (CH<sub>3</sub>OCH<sub>3</sub>) to formaldehyde (HCHO) was carried out on MoO<sub>x</sub> species with a wide range of  $MoO_x$  surface density and structure supported on MgO, Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub>. Raman and X-ray absorption spectroscopies were used to probe the structure of these MoO<sub>x</sub> domains, as they evolved from monomeric species into two-dimensional polymolybdate domains and  $MoO_3$  clusters with increasing  $MoO_x$  surface density. Primary HCHO synthesis rates (per Mo atom) initially increased with increasing  $MoO_x$  surface density (1.5–7 Mo/nm<sup>2</sup>) on all supports, indicating that  $MoO_x$  domain surfaces become more active as two-dimensional monolayers form via oligomerization of monomer species. The incipient formation of MoO<sub>3</sub> clusters at higher surface densities led to inaccessible  $MoO_x$  species and to lower HCHO synthesis rates (per Mo). Areal rates reached constant values as polymolybdate monolayers formed. These areal rates depend on the identity of the support; they were highest on SnO<sub>2</sub>, lowest on Al<sub>2</sub>O<sub>3</sub>, and undetectable on MgO, indicating that the surface properties of polymolybdate structures are strongly influenced by their atomic attachment to a specific support. The catalytic behavior of  $MoO_x$  domains reflects their ability to delocalize electron density during the formation of transition states required for rate-determining C-H bond activation steps within redox cycles involved in HCHO synthesis from dimethyl ether. These conclusions are consistent with the observed parallel increase in the rates of HCHO synthesis and of incipient stoichiometric reduction of  $MoO_x$  domains by H<sub>2</sub> as the domain size increases and as the supports become less insulating and more reducible, and as the energy required for ligandto-metal electronic transitions in the UV-visible spectrum decreases. HCHO selectivities increased with increasing  $MoO_x$  domain size; they were highest on Al<sub>2</sub>O<sub>3</sub> and lowest on SnO<sub>2</sub> supports. The Lewis acidity of the support cations appears to influence HCHO binding energy. HCHO reactions leading to  $CO_x$  and methyl formate via primary and secondary pathways are favored on weaker Lewis acids (with stronger conjugate bases). The Mo-O-support linkages prevalent at low surface densities also favor primary and secondary pathways to  $CO_x$  and methyl formate. When reported on a CH<sub>3</sub>OH-free basis, because of the pathways available for CH<sub>3</sub>OH oxidation to HCHO and for CH<sub>3</sub>OCH<sub>3</sub>–CH<sub>3</sub>OH interconversion, primary HCHO selectivities reached values greater than 95% on Al<sub>2</sub>O<sub>3</sub>-supported polymolybdate monolayers.

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## 1. Introduction

The significant thermodynamic and kinetic hurdles for the direct conversion of  $CH_4$  to alkenes and oxygenates have led to alternate processes involving synthesis gas intermediates [1–5]. For example, oxidative dehydrogenation of methanol is used to manufacture formaldehyde (HCHO) [6,7], which is then used as an intermediate to produce larger molecules containing C–C bonds. Recent advances in the

\* Corresponding author. *E-mail address:* iglesia@cchem.berkeley.edu (E. Iglesia). synthesis of dimethyl ether (DME; CH<sub>3</sub>OCH<sub>3</sub>) from H<sub>2</sub>/CO mixtures [8–10] and the anticipated similarities in the catalytic chemistries of DME and methanol on oxide surfaces [11,12] led us to consider DME as a cheaper and less toxic intermediate for the synthesis of chemicals currently produced from methanol. Previous studies reported low rates and selectivities for the catalytic conversion of DME to HCHO [13–18]. Recently, we have shown that MoO<sub>x</sub> and VO<sub>x</sub> domains supported on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> catalyze this reaction with high primary HCHO selectivities (80–98% HCHO, CH<sub>3</sub>OH-free basis) and high reaction rates at temperatures much lower than in previous studies (500–550 K vs 773 K) [19].

We have also previously reported the effects of the size and structure of  $ZrO_2$ -supported  $MoO_x$  domains on primary and secondary DME reaction rates [20]. Here, we report the significant effects of oxide supports on the structure, reducibility, and catalytic properties of active  $MoO_x$  domains. The acid–base properties [21,22], the surface charge [23,24], and the electronegativity of support cations [25,26] are often implicated in many support effects, but clear evidence of their specific involvement remains elusive. We present here a detailed study of selective DME oxidation reactions on  $MoO_x$  domains supported on MgO, Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> with a wide range of Mo surface density. We also examine the structure of these materials using complementary X-ray diffraction and Raman and X-ray absorption spectroscopic methods. These supports show a wide range of acidbase and redox surface properties. MgO and Al2O3 are unreducible oxides with basic and acid surfaces, respectively. ZrO<sub>2</sub> and SnO<sub>2</sub> are amphoteric oxides with surface sites that can undergo redox cycles and catalyze dehydrogenation steps. These support materials are used here to probe support effects on the structure and redox and catalytic properties of dispersed  $MoO_x$  domains, with specific emphasis on the rational design of selective catalysts for the conversion of dimethyl ether to formaldehyde.

#### 2. Experimental

#### 2.1. Synthesis of catalytic materials

Supported  $MoO_x$  catalysts were prepared by incipient wetness impregnation of ZrO(OH)<sub>2</sub>, SnO<sub>2</sub>, or MgO with aqueous (NH<sub>4</sub>)<sub>2</sub>Mo<sub>2</sub>O<sub>8</sub> (99%, Aldrich) solutions. ZrO(OH)<sub>2</sub> was prepared by hydrolysis of aqueous zirconyl chloride solutions (> 98%, Aldrich) at a pH of  $\sim 10$  using NH<sub>4</sub>OH (14.8 N, Fisher Scientific), followed by filtration of the precipitated powders and drying at 393 K overnight [27]. SnO<sub>2</sub> was prepared by hydrolysis of tin(IV) chloride pentahydrate (98%, Alfa Aesar) at a pH of  $\sim$  7 using NH<sub>4</sub>OH (14.8 N, Fisher Scientific). The precipitates were washed with deionized water until a Ag nitrate test failed to detected Cl ions in the effluent. The resulting solids were treated in flowing dry air (Airgas, zero grade) at 773 K for 3 h. MgO supports were prepared by treating MgO (> 98%, Aldrich) with deionized water at 355-365 K for 4 h, and then treating samples in flowing dry air (Airgas, zero grade) at 773 K for 8 h. Al<sub>2</sub>O<sub>3</sub>-supported MoO<sub>x</sub> catalysts were also prepared by incipient wetness impregnation, using  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> (Degussa, AG) and an aqueous (NH<sub>4</sub>)<sub>6</sub>Mo<sub>7</sub>O<sub>24</sub> (99%, Aldrich) solution at pH 5 [28]. The Mo content was varied by changing the Mo concentration in the impregnating solutions. All samples were dried at 393 K in ambient air after impregnation and then treated in flowing dry air (Airgas, zero grade) at 773 K for 3 h. Bulk MoO<sub>3</sub> was prepared by decomposition of (NH<sub>4</sub>)<sub>2</sub>Mo<sub>2</sub>O<sub>8</sub> (99%, Aldrich) in dry air at 773 K for 3 h. The Mo surface density for all supported samples is reported as  $Mo/nm^2$  per square nanometers based on the Mo content and the BET surface area for each sample.

# 2.2. Structural characterization

Surface areas were measured using N<sub>2</sub> at its normal boiling point (Autosorb-1; Quantachrome) and BET analysis methods. Powder X-ray diffraction patterns were measured at ambient conditions using a Siemens D-5000 diffractometer, Cu-K<sub> $\alpha$ </sub> radiation ( $\lambda = 1.5418$  Å), and a scan rate of 0.07° s<sup>-1</sup>.

Raman spectra were measured using a HoloLab 5000 Raman spectrometer (Kaiser Optical) and a frequencydoubled Nd:YAG laser at a wavelength of 532 nm. Samples were pressed into self-supporting thin wafers, placed on a rotary stage within a quartz cell, and spun at  $\sim 16$  Hz to avoid structural changes caused by laser local heating. The spectra were collected by exposure of the samples to laser for 6 s at a laser power of 50 mW and averaging of 100 spectra. Raman spectra were measured at 298 K for fresh and dehydrated samples. Dehydration was carried out in situ at 773 K for 1 h in flowing 20% O<sub>2</sub>/He.

Mo K-edge X-ray absorption spectra were measured using beamline 4-1 at the Stanford Synchrotron Radiation Laboratory. Samples were diluted with  $Al_2O_3$  for  $MoO_x/Al_2O_3$ samples or with SiO<sub>2</sub> for all other samples to achieve a Mo content of  $\sim 5$  wt% and then pelletized and sieved to retain particles 0.18–0.25 mm in diameter. The particles were placed within a thin quartz capillary tube (1.0-mm od, 0.1mm thickness) supported horizontally within a heated compartment held in the path of the rectangular X-ray beam  $(0.2 \times 6.0 \text{ mm})$  [29]. X-ray absorption spectra were measured in transmission mode with Ar gas in three ion chamber detectors: one chamber was located before the sample to measure the incident X-ray intensity, another chamber was located after the sample and before a Mo reference foil, and the third chamber was placed after the Mo foil. The incident beam energy was selected using a Si (111) crystal monochromator with 5-eV increments in the pre-edge region (19.875-19.975 keV), 0.25-eV increments in the edge region (19.975–20.035 keV), and 0.04  $\text{\AA}^{-1}$  in the fine structure region (20.035-21.024 keV). Energies were calibrated by placing the first inflection point in the Mo foil spectrum at its known absorption energy (19.999 keV). The spectra were analyzed using WinXAS (Version 1.2) [30]. A linear fit to the pre-edge region was subtracted from the entire spectrum, and then the spectrum was normalized with a third order polynomial. After conversion to k-space, the  $k^3$ -weighed fine structure spectrum was Fourier-transformed between 2.0 and 16.0 Å.

#### 2.3. Reducibility of supported $MoO_x$ samples in $H_2$

Reduction rates for  $MoO_x/ZrO_2$  samples with Mo surface densities of 0.5–6.4 Mo/nm<sup>2</sup> and for  $MoO_x/Al_2O_3$  (7.0

Mo/nm<sup>2</sup>) and MoO<sub>x</sub>/MgO (5.8 Mo/nm<sup>2</sup>) in H<sub>2</sub> were measured using a surface area analyzer (Quantachrome) modified with electronic mass flow controllers. The sample temperature was increased from 298 to 1253 K at 0.167 K s<sup>-1</sup> in flowing 20% H<sub>2</sub>/Ar (1.33 cm<sup>3</sup> s<sup>-1</sup>; Matheson UHP, certified mixture). An amount of sample containing 10 mg Mo was placed within a quartz cell. The H<sub>2</sub> concentration in the effluent stream was measured using a thermal conductivity detector after the water formed during reduction was removed using a 13X molecular sieve held at ambient temperature. The thermal conductivity response was calibrated from the reduction of pure CuO powder (99.995%, Aldrich). Initial oxygen removal rates were calculated from these data using kinetic analysis methods previously reported [28,31].

 $MoO_x$  reduction rates in H<sub>2</sub> were also measured for  $MoO_x/SnO_2$  (6.3  $Mo/nm^2$ ),  $MoO_x/ZrO_2$  (6.4  $Mo/nm^2$ ), and  $MoO_x/Al_2O_3$  (7.0  $Mo/nm^2$ ) using in situ X-ray absorption spectroscopy (XANES) (Mo *K*-edge). Near-edge absorption spectra were measured using a Si (111) crystal monochromator with 5-eV increments in the pre-edge region (19.905–19.990 keV), 0.50-eV increments in the edge region (19.990–20.033 keV), and 0.04 Å<sup>-1</sup> in the fine structure region (20.033–20.700 keV). In this approach, the extent of reduction of Mo<sup>6+</sup> to Mo<sup>4+</sup> was subtracted from the XANES spectra analyzed between 19.990 and 20.120 keV using linear superimposition methods [28] and the spectra of bulk MoO<sub>2</sub> and of Mo<sup>6+</sup> in the fresh catalyst samples.

## 2.4. Catalytic dimethyl ether reactions

Dimethyl ether reactions were carried out at 453–553 K in a fixed-bed quartz microreactor using supported MoO<sub>x</sub> samples (0.15–0.30 g) diluted with acid-washed quartz powder ( $\sim 1$  g) to prevent temperature gradients. Samples were treated in flowing 20% O<sub>2</sub>/He (0.67 cm<sup>3</sup> s<sup>-1</sup>) for 1.5 h at 773 K before catalytic measurements. The reactant mixture consisted of 80 kPa DME (99.5%, Praxair), 18 kPa O<sub>2</sub>, and 2 kPa N<sub>2</sub> (2 kPa) (Praxair, certified O<sub>2</sub>/N<sub>2</sub> mixture). Homogeneous dimethyl ether reactions were detected using empty reactors only above 593 K.

The reactor effluent was analyzed by on-line gas chromatography (Hewlett–Packard 6890 GC) using a methyl silicone capillary column (HP-1; 30 m, 0.25 mm, 0.25  $\mu$ m film) and a Porapak Q packed column (80–100 mesh, 1.82 m, 3.18 mm) connected to flame ionization and thermal conductivity detectors, respectively. Methanol, formaldehyde (HCHO), methyl formate (MF), CO, CO<sub>2</sub>, H<sub>2</sub>O, and traces of dimethoxymethane (DMM) were the only products detected.

Reaction rates and product selectivities were measured as a function of dimethyl ether conversion, which was varied by changing the reactant space velocity. Reactant conversions were kept below 10%. DME reaction rates and product selectivities were extrapolated to zero residence time to obtain the corresponding primary rates and selectivities. In view of the available pathways for DME–CH<sub>3</sub>OH interconversion and for CH<sub>3</sub>OH conversion to HCHO, rates and selectivities are reported here, unless noted otherwise, on a methanol-free basis.

## 3. Results and discussion

# 3.1. Structural characterization of supported $MoO_x$ catalysts

Fig. 1a shows Raman spectra for fresh  $MoO_x/SnO_2$  samples with a wide range of Mo surface densities (2.1–11.3 Mo/nm<sup>2</sup>). Raman bands were detected at 636 (s), 777 (w), 864–870 (s), and 953–967 (s) cm<sup>-1</sup> on all samples. The bands at 636 and 777 cm<sup>-1</sup> are assigned to SnO<sub>2</sub> crystallites [24,32,33], while those at 864–870 and 953–967 cm<sup>-1</sup> correspond to bridging Mo–O–Mo and terminal Mo=O stretching modes in oligomeric MoO<sub>x</sub> species, respectively, [24]. At 6.3 Mo/nm<sup>2</sup>, an additional shoulder appeared at 825 cm<sup>-1</sup>, which evolved into a sharp band along with two new bands at 672 and 1001 cm<sup>-1</sup> as the Mo surface density reached 11.3 Mo/nm<sup>2</sup>; these new bands correspond to the incipient formation of crystalline MoO<sub>3</sub> [24] as the surface density increases above that expected for a polymolybdate monolayer (~ 5 Mo/nm<sup>2</sup>) [34].

The band at  $\sim 965 \text{ cm}^{-1}$  for terminal Mo=O stretches shifted to higher frequencies and split into a doublet (986 and  $1003 \text{ cm}^{-1}$ ) (Fig. 1b) after samples were treated in 20% O<sub>2</sub>/He at 773 K; all other bands remained unchanged. The intensity of these two bands (relative to the  $SnO_2$  bands) increased with increasing Mo surface density, in parallel with the observed increase in the intensity of the Mo-O–Mo band at  $\sim 865 \text{ cm}^{-1}$ . Thus, we conclude that these bands correspond to Mo=O bonds in distorted oligomers (986 cm<sup>-1</sup>) and monomers (1003 cm<sup>-1</sup>). Exposure to moisture from ambient air at room temperature led to the coalescence of these two bands into the original feature at  $\sim 965 \text{ cm}^{-1}$ . This reversible process was first reported by Dumesic and co-workers [32], who used it as evidence for the complete dispersion of  $MoO_x$  structures on  $SnO_2$  [32]. The observed shift to higher frequencies on water desorption is associated with a change in Mo coordination and in bond order in terminal Mo=O structures [24].

These results suggest that isolated monomolybdates condense to form oligomeric two-dimensional structures as Mo densities reach ~ 6 Mo/nm<sup>2</sup> on SnO<sub>2</sub> surfaces and then form crystalline MoO<sub>3</sub> at higher coverages. X-ray diffraction measurements showed lines corresponding to MoO<sub>3</sub> only for surface densities above 6.3 Mo/nm<sup>2</sup>. This structural evolution of MoO<sub>x</sub> domains on SnO<sub>2</sub> supports resembles that reported previously on ZrO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> supports [28,35,36].

On  $MoO_x$  domains supported on  $SnO_2$ ,  $ZrO_2$ , and  $Al_2O_3$ , maximum DME conversion rates (per Mo atom) were obtained at Mo surface densities of 6–7 Mo/nm<sup>2</sup>, which resemble those expected for oligomeric two-dimensional struc-



Fig. 1. Raman spectra for  $MoO_x/SnO_2$  catalysts with surface densities of 2.1–11.3 Mo/nm<sup>2</sup> at ambient conditions (a) and after treatment in flowing dry air at 773 K for 1 h (b).

tures (~ 5.0 Mo/nm<sup>2</sup>); in contrast, MoO<sub>x</sub>/MgO samples showed no detectable catalytic activity (as discussed below).

These support effects on the catalytic properties of  $MoO_x$  oligomers were probed first by examining any structural differences detectable in the Raman spectra of samples with different supports but similar surface densities. Figs. 2a and 2b show the Raman spectra for these samples before and after treatment at 773 K in 20% O<sub>2</sub>/He, respectively. Before treatment,  $MoO_x/ZrO_2$  (6.4 Mo/nm<sup>2</sup>) showed two



Fig. 2. Raman spectra for supported  $MoO_x$  catalysts on MgO, Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> with surface densities of 5.8–7 Mo/nm<sup>2</sup> at ambient conditions (a) and after treatment in flowing dry air at 773 K for 1 h (b).

broad bands at ~ 947 and ~ 850 cm<sup>-1</sup> and a shoulder at ~ 920 cm<sup>-1</sup> (Fig. 2a). The removal of water at 773 K shifted the band at ~ 947 to 1003 cm<sup>-1</sup>, without any detectable changes in the other two bands (Fig. 2b). The bands at ~ 850 and 1003 cm<sup>-1</sup> arise from Mo–O–Mo and Mo=O vibrational stretches in two-dimensional polymolybdates [24]. The band at ~ 920 cm<sup>-1</sup> is assigned to symmetric vibrations of O=Mo=O structures in distorted MoO<sub>4</sub> tetrahedral monomers [36] or to Mo–O–Mo stretches in distorted MoO<sub>6</sub> octahedra within two-dimensional oligomers. Raman bands

for MoO<sub>3</sub> were not detected, indicating the absence of threedimensional crystallites.

The two broad bands at  $\sim 874$  and  $920 \text{ cm}^{-1}$  in the fresh  $MoO_x/MgO$  sample (5.8 Mo/nm<sup>2</sup>) (Fig. 2a) are similar to those reported for aquo  $MoO_4^{2-}$  ions, indicating the presence of MoO<sub>4</sub> tetrahedra exposed at MgO surfaces [37]. Water desorption during treatment at 773 K led to Raman bands at 811, 862, 911, and 952  $\text{cm}^{-1}$  (Fig. 2b), and to a spectrum similar to that previously reported for MoO<sub>3</sub>/MgO (5-7 Mo/nm<sup>2</sup>) [37]. The first two bands were assigned to Mo-O-Mo or Mo-O-Mg stretches and the latter two bands to Mo=O symmetric and antisymmetric stretches in polymolybdates [37]. These bands are not associated with crystalline magnesium molybdates, which exhibit sharp bands at ~ 970, ~ 950, and 910 cm<sup>-1</sup> [24,32,37,38]. The structural changes observed on hydration-dehydration are reversible, which precludes the presence of crystalline MgMoO<sub>4</sub> structures reported by others [24,38] and suggests instead a reversible tetrahedral-to-octahedral transformation of surface-exposed  $MoO_x$  species on water desorption [37].

Fresh  $MoO_x/Al_2O_3$  (7.0 Mo/nm<sup>2</sup>) samples showed a broad band at ~ 962 cm<sup>-1</sup>, assigned to two-dimensional oligomers [24,28], and three sharper bands at 673, 825 and 1001 cm<sup>-1</sup>, characteristic of crystalline MoO<sub>3</sub> (Fig. 2a). Treatment at 773 K influenced only the band at ~ 962 cm<sup>-1</sup>, which shifted to 1001 cm<sup>-1</sup> on water desorption. The strong MoO<sub>3</sub> bands in this sample suggest the presence of crystalline MoO<sub>3</sub> structures, but Raman scattering cross sections are 10–10<sup>3</sup> times greater for MoO<sub>3</sub> than for dispersed MoO<sub>x</sub> species [39,40]. Thus, we conclude from the similar intensities for the 962 and the 1001 cm<sup>-1</sup> bands (Fig. 2a) that the sample contains predominately two-dimensional MoO<sub>x</sub> oligomers; this conclusion was confirmed by the absence of strong diffraction lines for MoO<sub>3</sub> and by the lack of features in the X-ray absorption fine structure spectra.

The structure of  $MoO_x$  domains on these supports (at 6–7 Mo/nm<sup>2</sup>) was also probed using radial structure functions obtained from the X-ray absorption fine structure at the Mo *K* edge (Fig. 3). The main feature at ~ 1.3 Å corresponds to first-shell Mo–O distances, while the much weaker features at ~ 3.2 Å correspond to Mo–Mo, Mo–Al, Mo–Sn, or Mo–Zr distances. These radial structure functions differ markedly from those measured for crystalline MoO<sub>3</sub> (Fig. 3), which show the largest feature at Mo–Mo distances of ~ 3.2 Å and two prominent Mo–O bands at ~ 1.3 and 1.6 Å [41]. Thus, supported MoO<sub>x</sub> domains in all samples differ structurally from MoO<sub>3</sub> crystallites, as also concluded from Raman and X-ray diffraction measurements.

# 3.2. Dimethyl ether reaction rates and HCHO selectivities on supported $MoO_x$ catalysts

Fig. 4a shows primary DME reaction rates per Mo atom at 513 K as a function of Mo surface density on various supports. Rates were always higher on  $MoO_x/SnO_2$  than on  $MoO_x/ZrO_2$  and  $MoO_x/Al_2O_3$ .  $MoO_x/MgO$  catalysts did



Fig. 3. X-ray absorption radial structure functions for crystalline MoO<sub>3</sub> and for MoO<sub>x</sub> domains supported on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> after treatment in flowing dry air at 773 K for 1 h.

not give measurable DME conversions at these reaction conditions. On Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> supports, rates increased with increasing Mo surface density and reached maximum values at intermediate surface densities of  $6-7 \text{ Mo/nm}^2$ . For surface densities below those required for maximum rates (per Mo), most Mo atoms in MoO<sub>x</sub> domains are accessible to reactants, as discussed in the previous section. Therefore, the reported rates per Mo represent turnover rates and reflect the intrinsic surface reactivity of available MoO<sub>x</sub> domains.

The observed increase in reaction rates with increasing Mo surface density arises from a higher reactivity of exposed surfaces as the size and dimensionality of  $MoO_x$ domains increase. The size of the dispersed  $MoO_x$  domains increased monotonically with increasing Mo surface density, as recently shown by changes in the electronic properties of supported  $MoO_x$  detected by UV-visible spectroscopy with increasing  $MoO_x$  domain size [28,42]. Monomeric species appear to be much less reactive than two-dimensional oligomers and the reactivity of the latter appears to increase with increasing domain size. Larger and more connected  $MoO_x$  species are able to delocalize the additional electron density imposed by the reduction part of any catalytic redox cycles, such as those likely to be involved in dimethyl ether oxidation to formaldehyde. As Mo surface densities increased further ( $\gtrsim 6.5 \text{ Mo/nm}^2$ ), DME reaction rates (per Mo) decreased, because of the formation of MoO<sub>3</sub> clusters with inaccessible  $MoO_x$  structures.

These effects of surface density and domain size are similar to those observed for oxidative dehydrogenation reactions of alkanes on  $MoO_x$  and  $VO_x$  catalysts [28,42]. As in the case of dehydrogenation reactions, propane oxidation rates increase as electronic transitions between lattice oxy-



Fig. 4. Primary dimethyl ether conversion rates normalized per Mo atom (a) and normalized per catalyst BET surface area (b) as a function of Mo surface density on  $MoO_x$  domains supported on  $Al_2O_3$ ,  $ZrO_2$ ,  $SnO_2$ , and MgO (513 K; 80 kPa DME, 18 kPa  $O_2$ , and 2 kPa  $N_2$ ).

gen and active metal cations (responsible for the UV–visible absorption edge) decrease in energy [42]. These findings differ markedly from those reported for methanol oxidation reactions on supported VO<sub>x</sub> and MoO<sub>x</sub> catalysts, for which reaction rates (per active metal atom) appear to be independent of V or Mo content [43,44], even though redox cycles are involved and the oxide domain size strongly influences the reducibility of the active catalytic domains [7,28].

Areal DME reaction rates were estimated for each supported  $MoO_x$  sample based on their total BET surface areas. Areal rates increased with increasing Mo surface density (Fig. 4b) and reached nearly constant values for surface densities above 6-7 Mo/nm<sup>2</sup> on all supports. These trends suggest that support surfaces become entirely covered with  $MoO_x$  structures and that any  $MoO_x$  species in excess of the amount required to form an active polymolybdate monolayer lead to  $MoO_x$  crystallites, which do not contribute appreciably to the exposed active surface area or to the measured reaction rates. These asymptotic areal rates, however, depend on the identity of the support, as expected from the existing atomic connectivity between MoO<sub>x</sub> layers and oxygen atoms on support surfaces (Table 1). Constant areal rates of about 6, 9, and  $50 \times 10^{-5} \text{ mol}/(\text{m}^2 \text{ h})$  were obtained on  $MoO_x/Al_2O_3$ ,  $MoO_x/ZrO_2$ , and  $MoO_x/SnO_2$ , respectively, as each sample approached polymolybdate monolayer coverages. At low surface densities, the exposed inactive regions on support surfaces lead to low calculated areal rates, which increase as these regions become covered with monomeric molybdates of low intrinsic activity and ultimately with more active  $MoO_x$  oligomers.

Oxidative DME reactions on  $MoO_x$ -based catalysts proceed via primary and secondary pathways depicted in Scheme 1 [20]. These pathways include the direct conversion of DME to CH<sub>3</sub>OH ( $k_0$ ), HCHO ( $k_1$ ), methyl formate (MF)  $(k_2)$ , and CO<sub>x</sub> (CO + CO<sub>2</sub>)  $(k_3)$  in one surface sojourn. They also include secondary reactions of HCHO to form MF ( $k_4$ ) and CO<sub>x</sub> ( $k_5$ ) and of MF to form CO<sub>x</sub> ( $k_6$ ). In each case,  $k_i$  represents the rate constant for reaction *i*. At the low conversions of this study, all primary reactions can be assumed to be pseudo-zero order in DME and O<sub>2</sub>, because their respective concentrations remain nearly constant with changes in reactant residence time or conversion [20]. Secondary reactions are assumed to be pseudo-first order in the concentration of the primary product involved, but any kinetic dependencies on DME and O<sub>2</sub> are neglected for the same reasons proposed for primary reactions. Traces of dimethoxymethane (DMM) were also detected, but are not included in the reaction scheme because of the low selectivities (< 1%). CH<sub>3</sub>OH selectivities were largely unaffected by residence time, suggesting that CH<sub>3</sub>OH is formed directly as a co-product of HCHO formation. Its alternate formation via DME hydration would require H<sub>2</sub>O co-reactants, which are

Table 1

Some results on MoO<sub>x</sub> supported on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> with surface densities of 11.3–12.6 Mo/nm<sup>2</sup> as well as on MgO (5.8 Mo/nm<sup>2</sup>)

Support	Areal rate $(10^{-5} \text{ mol}/(\text{m}^2 \text{ h}))$	Primary HCHO selectivity (%)	$k_1/(k_2 + k_3)$	$k_1/((k_4+k_5)C_{\rm A_0})$
Al <sub>2</sub> O <sub>3</sub>	5.7	98.0	47.0	0.35
ZrO <sub>2</sub>	7.5	81.8	4.5	0.08
SnO <sub>2</sub>	48.0	88.4	7.6	0.10
MgO	0.0	_	-	-



Scheme 1. Primary and secondary reaction pathways for dimethyl ether conversion on  $MoO_x$ -based catalysts

not available at low conversions. The weak effects of residence time on CH<sub>3</sub>OH indicate also that CH<sub>3</sub>OH oxidation is slower than DME oxidation under these reaction conditions.

The rate constants  $k_1$ ,  $k_2$ , and  $k_3$  were calculated from the corresponding pseudo-zero-order primary reaction rates, and  $(k_4 + k_5)C_{A_0}$  ( $C_{A_0}$  = inlet DME concentration) from the observed changes in HCHO selectivity ( $S_B$ ) with  $\nu$  as given by

$$S_{\rm B} = S_{\rm B}^0 \left[ 1 - \frac{1}{2} (k_4 + k_5) C_{\rm A_0} \cdot \nu \right],$$

where  $S_B^0$  is the primary HCHO selectivity,  $C_{A_0}$  is the inlet DME concentration, and  $\nu = \text{g-atom Mo/inlet molar DME}$  rate (g-atom Mo-s/mol DME).  $S_B^0$  is given by  $k_1/(k_1 + k_2 + k_3)$  (on a CH<sub>3</sub>OH-free basis) [20].

Primary HCHO selectivities on  $MoO_x/ZrO_2$  and  $MoO_x/$ SnO2 catalysts increased monotonically with increasing Mo surface density and reached constant values of  $\sim 90\%$  for surface densities higher than  $6-7 \text{ Mo/nm}^2$  (Fig. 5a). Concurrently, primary MF and  $CO_x$  (CO+CO<sub>2</sub>) selectivities decreased to constant values of  $\sim$  7 and 3% (Figs. 5b and 5c). This indicates that exposed SnO<sub>2</sub> and ZrO<sub>2</sub> surfaces or Mo-O–Sn(Zr) structures show higher MF and  $CO_x$  selectivities than Mo-O-Mo structures, as also reported for CH<sub>3</sub>OH oxidation reactions [43,45,46]. Yet, none of these supports led to significant DME conversion rates in the absence of  $MoO_x$ active species. On  $MoO_x/Al_2O_3$ , MF (2–4%) and  $CO_x$ (0-1.2%) selectivities were very low and nearly independent of Mo surface density (Figs. 5b and 5c), while HCHO selectivities increased only slightly from 95.2 to 98.1% as the Mo surface density increased from 1.6 to 11.3 Mo/nm<sup>2</sup> (Fig. 5a). Thus, it appears that Mo-O-Al structures are essentially unreactive in DME conversion reactions to MF and  $CO_x$ . These results show that the coverage of the support surfaces with active  $MoO_x$  domains favor high HCHO selectivities, especially in the case of more reducible ZrO2 and SnO<sub>2</sub> supports.

The redox properties of  $MoO_x$  domains influence DME oxidation rates (as discussed below), while acid–base properties may influence reactions of DME-derived adsorbed in-



Fig. 5. Primary selectivities to HCHO (a), methyl formate (b), and  $CO_x$  (c) as a function of Mo surface density on supported  $MoO_x$  catalysts on  $Al_2O_3$ ,  $ZrO_2$ , and  $SnO_2$  (513 K; 80 kPa DME, 18 kPa  $O_2$ , and 2 kPa  $N_2$ ).

100

termediates, as well as the readsorption and secondary reactions of HCHO. DME oxidation proceeds via methoxy groups formed by DME dissociation on Mo-O sites, followed by a reduction step involving C-H bond activation in methoxy groups (CH<sub>3</sub>O<sup> $\delta$ -</sup>) using lattice oxygen [47]. High primary HCHO selectivities require that adsorbed HCHO species desorb before subsequent reactions. More acidic and less reducible surfaces favor HCHO desorption, while inhibiting the formation of strongly adsorbed dioxymethylene and formate species, which require nucleophilic attack by basic lattice oxygens to form ultimately MF and  $CO_x$ . It appears that exposed ZrO<sub>2</sub> and SnO<sub>2</sub> surfaces or heteroatomic Mo-O-support linkages prevalent at surface densities below 6–7 Mo/nm<sup>2</sup> provide strong binding sites for CH<sub>3</sub>O<sup> $\delta$ -</sup> and HCHO. Similar effects have been proposed for methanol oxidation reactions [45,46]. Infrared and desorption studies of adsorbed methanol detected the presence of strongly adsorbed dioxymethylene and formate species on SnO<sub>2</sub> and  $ZrO_2$  surfaces, which led only to  $CO_x$ , while more weakly adsorbed  $CH_3O^{\delta-}$  species on MoO<sub>x</sub> species formed HCHO [45,46].

Figs. 6a and 6b show  $k_1/(k_2 + k_3)$  and  $k_1/((k_4 + k_5)C_{A_0})$ values, respectively, as a function of Mo surface density for all samples. Values of  $k_1/(k_2 + k_3)$  reflect primary DME conversion rates to HCHO  $(k_1)$  relative to those for MF  $(k_2)$  and CO<sub>x</sub>  $(k_3)$  formation; thus, they provide an alternate measure of primary HCHO selectivities. In contrast,  $k_1/((k_4 + k_5)C_{A_0})$  ratios (at a constant inlet DME concentration,  $C_{A_0}$ ) reflect primary HCHO formation rates  $(k_1)$  relative to the rates for secondary HCHO reactions to form MF  $(k_4)$  and CO<sub>x</sub>  $(k_5)$ ; this kinetic parameter reflects the tendency of a given catalyst to form and convert HCHO. High values of  $k_1/(k_2 + k_3)$  and  $k_1/((k_4 + k_5)C_{A_0})$  lead to higher HCHO selectivities for a given DME conversion; the second parameter becomes increasingly important in determining HCHO yields as DME conversion increases.

Both  $k_1/(k_2+k_3)$  and  $k_1/((k_4+k_5)C_{A_0})$  ratios increased with increasing Mo surface density and reached constant values above 6–7 Mo/nm<sup>2</sup> on MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub>, MoO<sub>x</sub>/ZrO<sub>2</sub>, and  $MoO_x/SnO_2$  (Fig. 6). Thus, the increasing coverage of the support with  $MoO_x$  species and the larger  $MoO_x$ domains formed as surface density increases led to more selective catalytic surfaces and to higher HCHO yields. The higher  $k_1/(k_2+k_3)$  and  $k_1/((k_4+k_5)C_{A_0})$  values on MoO<sub>x</sub> domains supported on Al<sub>2</sub>O<sub>3</sub> relative to those on SnO<sub>2</sub> and  $ZrO_2$  at the constant  $C_{A_0}$  suggest that more reducible supports with weak Lewis acidity favor the conversion of adsorbed HCHO species to MF and  $CO_x$ , as discussed above. Apparently, such supports allow Mo cations in  $MoO_x$  domains to become more electron-rich, thus lowering their affinity for binding electron-donating DME-derived intermediates and HCHO. Similar effects of the Lewis acidity of active oxide cations and of the supports have been reported for secondary reactions of alkenes during oxidative dehydrogenation of alkanes on  $VO_x$ ,  $MoO_x$ , and  $WO_x$ catalysts [48]. In this case, electron-rich alkenes interact

MoO<sub>x</sub>/SnO<sub>2</sub> 10 MoO<sub>x</sub>/ZrO<sub>2</sub> k1/(k2+k3) 1 0.1 0 30 10 20 Mo surface density (Mo/nm<sup>2</sup>) 1 (b) MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub> k<sub>1</sub>/((k<sub>4</sub>+k<sub>5</sub>) C<sub>Ao</sub>) MoO<sub>x</sub>/SnO<sub>2</sub> 0.1 0 С MoO<sub>x</sub>/ZrO<sub>2</sub> 0.01 Á 0.001 0 5 25 10 15 20 30 Mo surface density (Mo/nm<sup>2</sup>)

MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub>

Fig. 6. Rate constant ratios  $k_1/(k_2 + k_3)$  (a) and  $k_1/((k_4 + k_5)C_{A_0})$  ( $C_{A_0}$ ; inlet DME concentration) (b) (presented in a logarithmic scale) as a function of Mo surface density on MoO<sub>x</sub> domains supported on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> (513 K; 80 kPa DME, 18 kPa O<sub>2</sub>, and 2 kPa N<sub>2</sub>).

strongly with more acidic active metal cations, leading to longer surface residence times during their initial formation from alkanes and to a greater tendency for them to readsorb and undergo secondary combustion reactions.

Next, we turn our attention to the underlying basis for the observed support effects on DME reaction rates and HCHO selectivity. DME reaction rates are significantly higher for MoO<sub>x</sub> domains supported on SnO<sub>2</sub> than for those on ZrO<sub>2</sub> or Al<sub>2</sub>O<sub>3</sub>, even when polymolybdate monolayers are prevalent on all supports; in contrast, MoO<sub>x</sub> species on MgO are inactive (Fig. 4, Table 1). Unreducible Al<sub>2</sub>O<sub>3</sub> supports give higher HCHO selectivities than ZrO<sub>2</sub> or SnO<sub>2</sub> (Fig. 5). The  $k_1/(k_2 + k_3)$  and  $k_1/((k_4 + k_5)C_{A_0})$ 

(a)

values decreased from 47 and 0.35 on Al<sub>2</sub>O<sub>3</sub> to 8–9 and 0.08–0.10 on ZrO<sub>2</sub> and SnO<sub>2</sub> at the constant  $C_{A_0}$ , respectively (Fig. 6). Clearly, the chemical properties of a polymolybdate monolayer depend on the identity of the underlying support to which it is atomically connected. In the next section, we examine the effects of support on MoO<sub>x</sub> reduction properties for catalysts containing predominately polymolybdate monolayers (6–7 Mo/nm<sup>2</sup>).

# 3.3. Genesis of support effects in dimethyl ether oxidation on $MoO_x$ domains

For Mo surface densities of  $6-7 \text{ Mo/nm}^2$ , all  $MoO_x$ species on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, SnO<sub>2</sub>, and MgO supports are accessible to reactants; therefore, differences in accessibility cannot account for the observed effects of supports on DME reaction rates. Kinetic and isotopic studies have shown that HCHO formation from DME involves redox cycles and kinetically relevant C-H bond activation steps [47]. The transition states required for such C-H bond activation steps involve the incipient local reduction of  $MoO_x$  domains to form an OH group and adsorbed HCHO species bound to a metal cation. Reactions limited by C-H bond activation on metal oxides, such as oxidative alkane dehydrogenation on  $MoO_x$ and  $VO_x$ , show excellent correlations between catalytic reaction rates and the electronic and redox properties of oxide domains [42]. Thus, we suggest that the strong support effects observed for DME oxidation reactions also reflect differences in the reducibility of  $MoO_x$  domains on the various supports.

The rates of reduction of supported  $MoO_x$  domains were measured using H<sub>2</sub> as the stoichiometric reductant while the sample temperature was increased from 298 to 1253 K. Specifically, the rates of the incipient reduction of  $Mo^{6+}$  to  $Mo^{4+}$  were measured. In-situ X-ray absorption spectra showed that few reduced Mo centers are present during steady-state DME reactions at 513 K [47]. Thus, the initial stages of  $Mo^{6+}$  reduction are the chemical processes most relevant to the redox cycles required for catalytic turnovers, as discussed in detail elsewhere for related oxidation reactions [28,42].

Fig. 7 shows incipient reduction rates in H<sub>2</sub> (at 623 K) as a function of Mo surface density for MoO<sub>x</sub> domains supported on ZrO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, and MgO at surface densities of  $6-7 \text{ Mo/nm}^2$ , as well as for three other ZrO<sub>2</sub>-supported samples with Mo surface densities of 0.5, 3.5, and 5.6 Mo/nm<sup>2</sup> and for a MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub> sample with 3.4 Mo/nm<sup>2</sup>. These incipient H<sub>2</sub> reduction rates increased with increasing Mo surface density for each support. They were higher on MoO<sub>x</sub>/ZrO<sub>2</sub> than on MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub> and MoO<sub>x</sub>/MgO catalysts for a given surface density. For MoO<sub>x</sub>/SnO<sub>2</sub>, however, the extensive concurrent reduction of MoO<sub>x</sub> domains and of the SnO<sub>2</sub> support precludes the use of this method to measure MoO<sub>x</sub> reduction rates.

Fig. 8 shows a parallel increase in the rate of primary DME conversion (513 K) and in the rate of incipient



Fig. 7. Incipient reduction rates in H<sub>2</sub> (at 623 K) for MoO<sub>x</sub> domains supported on ZrO<sub>2</sub> ( $\bullet$ ), Al<sub>2</sub>O<sub>3</sub> ( $\blacktriangle$ ), and MgO ( $\blacksquare$ ) at similar Mo surface densities (5.8–7 Mo/nm<sup>2</sup>), on MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub> (3.4 Mo/nm<sup>2</sup>) ( $\bigstar$ ), and on MoO<sub>x</sub>/ZrO<sub>2</sub> with 0.5, 3.5, and 5.6 Mo/nm<sup>2</sup> ( $\bullet$ ).



Fig. 8. Relationship between primary DME reaction rates (513 K) and initial H<sub>2</sub> reduction rates (623 K) for MoO<sub>x</sub> domains supported on ZrO<sub>2</sub> ( $\bullet$ ), Al<sub>2</sub>O<sub>3</sub> ( $\blacktriangle$ ), and MgO ( $\blacksquare$ ) at similar Mo surface densities (5.8–7 Mo/nm<sup>2</sup>), on MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub> (3.4 Mo/nm<sup>2</sup>) ( $\bigstar$ ), and on MoO<sub>x</sub>/ZrO<sub>2</sub> with 0.5, 3.5, and 5.6 Mo/nm<sup>2</sup> ( $\bullet$ ).

stoichiometric reduction in  $H_2$  (623 K) for the samples shown in Fig. 7. On these samples,  $MoO_x$  species are predominately exposed at surfaces and accessible to both DME and  $H_2$  reactants. Thus, the measured reaction rates (per Mo) reflect the reactivity of exposed surfaces for the two respective reactions. Clearly, the tendency of these domains to undergo reduction with the generation of a small number of oxygen vacancies increases in parallel with their ability to catalyze DME reaction turnovers, which appear to involve



Fig. 9. X-ray absorption spectra near the Mo K edge for  $MoO_x/SnO_2$  (6.3  $Mo/nm^2$ ) after treatment in H<sub>2</sub> (20% H<sub>2</sub>/Ar) at 298, 623, 723, and 823 K and for crystalline MoO<sub>2</sub> at ambient temperature.

the Mars–van Krevelen pathways [49] involving reducible centers and lattice oxygen atoms.

The difficulties caused by the concurrent reduction of  $MoO_x$  and  $SnO_2$  led us to examine the dynamics of reduction in H<sub>2</sub> at 298-823 K using in situ X-ray absorption features near the Mo K edge for  $MoO_x$  domains on SnO<sub>2</sub>, ZrO<sub>2</sub>, and Al<sub>2</sub>O<sub>3</sub> with similar surface densities (6–7  $Mo/nm^2$ ). The concentrations of  $Mo^{6+}$  and  $Mo^{4+}$  were estimated by linear superimposition methods using the starting spectrum and the spectrum of MoO2 as principal components. The Mo K absorption edge arises from the ejection of a 1s photoelectron, while the pre-edge feature at  $\sim$  19.99 keV corresponds to a 1s-to-4d transition that is dipole-forbidden in centrosymmetric structures [29,41]. The Mo<sup>6+</sup> cations in MoO<sub>3</sub> adopt off-center positions within distorted MoO<sub>6</sub> octahedra, which lead to p-d orbital hybridization and to a detectable pre-edge feature in the MoO<sub>3</sub> spectrum. In contrast, MoO<sub>2</sub> has only a very weak pre-edge feature. Recently, Ressler et al. [41] and Chen et al. [28] reported reduction rates for crystalline MoO<sub>3</sub> and supported MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>, respectively, from linear superimposition of Mo K edge spectra.

Fig. 9 shows the Mo K near-edge spectra for  $MoO_x/SnO_2$ (6.3 Mo/nm<sup>2</sup>) during contact with 20% H<sub>2</sub>/Ar at 298, 623, 723, and 823 K, and for crystalline MoO<sub>2</sub> at ambient conditions. The pre-edge feature weakened with increasing temperature and the spectral features approached those typical of MoO<sub>2</sub>. Above 723 K, all spectra are accurately described using only the MoO<sub>2</sub> standard spectrum, without any spectral contributions from either the starting sample or crystalline MoO<sub>3</sub>. Below 723 K, the spectra are accurately described by linear combinations of the spectra for the initial sample and for crystalline MoO<sub>2</sub>.



Fig. 10. MoO<sub>2</sub> fraction measured from linear superimposition of MoO<sub>2</sub> and initial spectra as a function of treatment temperature in 20% H<sub>2</sub>/Ar for MoO<sub>x</sub> domains supported on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> at similar Mo surface density (6.3–7 Mo/nm<sup>2</sup>).

Fig. 10 shows the fraction of the Mo cations present as Mo<sup>4+</sup> as a function of temperature for the three supported samples with surface densities of  $6-7 \text{ Mo/nm}^2$ . The  $Mo^{4+}$  fraction is significantly greater for  $MoO_x/SnO_2$  than for  $MoO_x/ZrO_2$  and  $MoO_x/Al_2O_3$  at all temperatures, indicating that  $MoO_x$  species dispersed on  $SnO_2$  are easier to reduce than on the other two supports. A similar trend was observed for the reactivity of  $MoO_x$  species in these samples for DME reactions (Fig. 4).  $MoO_x$  domains supported on SnO<sub>2</sub> appear to delocalize the electron density that they acquire during stoichiometric reduction processes and also during reduction steps involved in C-H bond activation steps within each catalytic redox cycle. Clearly, the reducibility of the support itself must play a role in these effects, because a shift from refractory MgO and Al<sub>2</sub>O<sub>3</sub> supports to more reducible ZrO2 and SnO2 leads to concomitant changes in the reduction properties of the supported  $MoO_x$ domains.

The observed parallel increases in the reducibility and catalytic reactivity of the active oxides is similar to those observed for other reactions catalyzed by supported  $MoO_x$  and  $VO_x$ , such as the oxidative dehydrogenation of alkanes [28, 31,50] and alcohols [43,44]. These findings are consistent with the involvement of lattice oxygen atoms and of transition states with significant reduction of metal centers in the catalytic cycles required for DME conversion to HCHO [47]. These relationships and mechanistic inferences provide the basis for the catalytic improvements that we have achieved by surface modifications of  $Al_2O_3$  supports, in an effort to increase the reactivity of supported  $MoO_x$  domains, without loss of the selective properties of  $Al_2O_3$ -supported  $MoO_x$  domains for DME conversion to HCHO [51].

## 4. Conclusions

 $MoO_x$  domains on Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, and SnO<sub>2</sub> catalyze the selective oxidation of dimethyl ether to HCHO at 453-553 K. No reaction products, however, were detected on  $MoO_x$  domains supported on MgO at similar surface densities. On all supports, the size and dimensionality of  $MoO_x$ domains increased with increasing  $MoO_x$  surface density. MoO<sub>3</sub> clusters were first detected by Raman spectroscopy at surface densities of  $6-7 \text{ Mo/nm}^2$ , which correspond to those expected for two-dimensional  $MoO_x$  monolayers. Primary HCHO synthesis rates (per Mo atom) were influenced by the Mo surface density on each support. These rates initially increased with increasing  $MoO_x$  surface density on all supports up to 6-7 Mo/nm<sup>2</sup>, indicating that the surface reactivity of  $MoO_x$  domains increases as a two-dimensional monolayer forms via oligomerization of monomolybdate species. Ultimately, the incipient formation of three-dimensional  $MoO_3$  clusters leads to some inaccessible  $MoO_x$  species and lower reaction rates (per Mo). Areal reaction rates reached a constant value as the polymolybdate monolayers form, but these areal rates were highest on SnO<sub>2</sub> supports and lowest on Al<sub>2</sub>O<sub>3</sub>, indicating that the catalytic properties of these polymolybdate layers depend strongly on the chemical identity of the support surfaces to which they are atomically attached. These domain size effects appear to reflect the effectiveness of larger domains to delocalize charge, which is required for kinetically relevant steps in the redox cycles involved in HCHO synthesis. This proposal is consistent with the faster rates of incipient reduction using H<sub>2</sub> with increasing  $MoO_x$  surface density and with changes in support. The domain size and the identity of the support influence charge delocalization properties of  $MoO_x$  domains and thus their catalytic and reduction rates. Reaction rates (per Mo atom) for supported samples with  $6-7 \text{ Mo/nm}^2$  are significantly higher on SnO<sub>2</sub> than on ZrO<sub>2</sub> or Al<sub>2</sub>O<sub>3</sub> supports. Similar trends were observed for stoichiometric reduction rates of these samples using H<sub>2</sub>. The effects of support on primary and secondary HCHO selectivities is consistent with the weaker binding of DME-derived intermediates and of HCHO on Mo<sup>6+</sup> sites with higher Lewis acidity; these sites favor desorption of HCHO precursors, while discouraging HCHO readsorption and secondary reactions to form  $CO_x$ and methyl formate.

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### References

- [1] R. Pitchai, K. Klier, Catal. Rev. Sci. Eng. 28 (1986) 13.
- [2] J.S. Lee, S.T. Oyama, Catal. Rev. Sci. Eng. 30 (1988) 249.
- [3] K. Tabata, Y. Teng, T. Takemoto, E. Suzuki, M.A. Banares, M.A. Pena, J.L. Garcia Fierro, Catal. Rev. Sci. Eng. 44 (2002) 1.
- [4] K. Otsuka, Y. Wang, Appl. Catal. 222 (2001) 145.
- [5] I.C. Bafas, I.E. Constantinou, C.G. Vayenas, Chem. Eng. J. 82 (2001) 109.
- [6] L. Lefferts, J.G. van Ommen, J.R.H. Ross, Appl. Catal. 23 (1986) 385.
- [7] J.M. Tatibouet, Appl. Catal. A 148 (1997) 213.
- [8] J.-L. Li, X.-G. Zhang, T. Inui, Appl. Catal. A 147 (1996) 23.
- [9] T.H. Fleisch, A. Basu, M.J. Gradassi, J.G. Masin, Stud. Surf. Sci. Catal. 107 (1997) 117.
- [10] T. Shikada, Y. Ohno, T. Ogawa, M. Ono, M. Mizuguchi, K. Tomura, K. Fujimoto, Stud. Surf. Sci. Catal. 119 (1998) 515.
- [11] M. Hara, M. Kawamura, J.N. Kondo, K. Domen, K. Maruya, J. Phys. Chem. 100 (1996) 14462.
- [12] F. Ouyang, S. Yao, J. Phys. Chem. 104 (2000) 11253.
- [13] H. Tadenuma, T. Murakami, H. Mitsushima, US Patent 3,655,771.
- [14] R.M. Lewis, R.C. Ryan, L.H. Slaugh, US Patent 4,442,307, 1984.
- [15] R.M. Lewis, R.C. Ryan, L.H. Slaugh, US Patent 4,439,624, 1984.
- [16] R.M. Lewis, R.C. Ryan, L.H. Slaugh, US Patent 4,435,602, 1984.
- [17] G.P. Hagen, M.J. Spangler, US Patent 6,265,528, 2001.
- [18] G.P. Hagen, M.J. Spangler, US Patent 6,350,919, 2002, assigned to BP.
- [19] H. Liu, E. Iglesia, J. Catal. 208 (2002) 1.
- [20] H. Liu, P. Cheung, E. Iglesia, J. Phys. Chem., in press.
- [21] A.N. Desikan, L. Huang, S.T. Oyama, J. Chem. Soc. Faraday Trans. 88 (1993) 3357.
- [22] A. Auroux, D. Sprinceana, A. Gervasini, J. Catal. 195 (2000) 140.
- [23] D.S. Kim, K. Segawa, T. Soeya, I.E. Wachs, J. Catal. 136 (1992) 539.
- [24] G. Mestl, T.K.K. Srinivasan, Catal. Rev. Sci. Eng. 38 (1998) 451.
- [25] M. Niwa, Y. Matsuoka, Y. Murakami, J. Phys. Chem. 91 (1987) 4519.
- [26] X. Gao, E. Wachs, Top. Catal. 18 (2002) 243.
- [27] A. Khodakov, B. Olthof, A.T. Bell, E. Iglesia, J. Catal. 181 (1999) 205.
- [28] K. Chen, S. Xie, A.T. Bell, E. Iglesia, J. Catal. 198 (2001) 232.
- [29] W. Li, G.D. Meitzner, R.W. Borry III, E. Iglesia, J. Catal. 191 (2000) 373.
- [30] T. Ressler, WinXAS 97, Version 1.2, 1998.
- [31] K. Chen, S. Xie, A.T. Bell, E. Iglesia, J. Catal. 195 (2000) 244.
- [32] S.R. Stampel, Y. Chen, J.A. Dumesic, C. Niu, C.G. Hill Jr., J. Catal. 105 (1987) 445.
- [33] B. Amram, H. Pernot, A.M. Govic, M. Gubelmann-Bonneau, Vibr. Spectrosc. 9 (1995) 65.
- [34] Y.C. Xie, Y.Q. Tang, Adv. Catal. 37 (1990) 1.
- [35] K. Chen, S. Xie, E. Iglesia, A.T. Bell, J. Catal. 189 (2000) 421.
- [36] S. Xie, K. Chen, A.T. Bell, E. Iglesia, J. Phys. Chem. 104 (2000) 10059.
- [37] S.-C. Chang, M.A. Leugers, S.R. Bare, J. Phys. Chem. 96 (1992) 10358.
- [38] D.S. Kim, I.E. Wachs, K. Segawa, J. Catal. 146 (1994) 268.
- [39] R. Radhakrishnan, C. Reed, S.T. Oyama, M. Semen, J.N. Kondo, K. Domen, Y. Ohminami, K. Asakura, J. Phys. Chem. 105 (2001) 8519.
- [40] C.C. Williams, J.G. Ekerdt, J.M. Jehng, F.D. Hardcastle, I.E. Wachs, J. Phys. Chem. 95 (1991) 9791.
- [41] T. Ressler, O. Timpe, T. Neisius, J. Find, G. Mestl, M. Dieterle, R. Schlogl, J. Catal. 191 (2000) 75.
- [42] K. Chen, A.T. Bell, E. Iglesia, J. Catal. 209 (2002) 35.
- [43] G. Deo, I.E. Wachs, J. Catal. 146 (1994) 323.
- [44] W. Zhang, A. Desikan, S.T. Oyama, J. Phys. Chem. 99 (1995) 14468.
- [45] M. Niwa, J. Igarashi, Catal. Today 52 (1999) 71.
- [46] N. Narishige, M. Niwa, Catal. Lett. 71 (2001) 63.
- [47] H. Liu, P. Cheung, E. Iglesia, unpublished results.
- [48] K. Chen, A.T. Bell, E. Iglesia, J. Phys. Chem. 104 (2000) 1292.
- [49] P. Mars, D.W. van Krevelen, Chem. Eng. Sci. 3 (1954) 41.
- [50] M.C. Abello, M.F. Gomez, L.E. Cadus, Catal. Lett. 83 (1998) 185.
- [51] H. Liu, P. Cheung, E. Iglesia, Phys. Chem.