

Quantification of Zinc Atoms in a Surface Alloy on Copper in an Industrial-Type Methanol Synthesis Catalyst**

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Abstract: Methanol has recently attracted renewed interest because of its potential importance as a solar fuel.^[1] Methanol is also an important bulk chemical that is most efficiently formed over the industrial Cu/ZnO/Al₂O₃ catalyst. The identity of the active site and, in particular, the role of ZnO as a promoter for this type of catalyst is still under intense debate.^[2] Structural changes that are strongly dependent on the pretreatment method have now been observed for an industrial-type methanol synthesis catalyst. A combination of chemisorption, reaction, and spectroscopic techniques provides a consistent picture of surface alloying between copper and zinc. This analysis enables a reinterpretation of the methods that have been used for the determination of the Cu surface area and provides an opportunity to independently quantify the specific Cu and Zn areas. This method may also be applied to other systems where metal–support interactions are important, and this work generally addresses the role of the carrier and the nature of the interactions between carrier and metal in heterogeneous catalysts.

It is widely accepted that the presence of ZnO increases the activity of copper-based systems towards methanol synthesis.^[2a,3] Under reducing conditions, the presence of ZnO_x species leads to partially covered Cu particles as a result of strong metal–support interactions (SMSI).^[1a,4] Evidence for surface alloy formation has been obtained from studies of the deposition of metallic zinc on a copper single crystal and by ion-scattering measurements of a Cu/ZnO catalyst with isotopically enriched Cu and Zn.^[5] In a previous study, a change in the copper area was observed subsequent to pretreatments in reducing atmospheres by combining N₂O reactive frontal chromatography (N₂O-RFC) and temperature-programmed desorption of H₂ (H₂-TPD).^[6] Studies of Cu/ZnO model systems have revealed that pretreatments influence the morphology of Cu nanoparticles by wetting/

non-wetting, which was explained by the formation of strongly binding oxygen vacancies at the ZnO surface.^[4a,b,7] Most recently, ambient-pressure X-ray photoelectron spectroscopy (XPS) of Cu/ZnO catalysts with a high Cu content has shown that surface decoration of Cu nanoparticles with ZnO_x species occurs under relatively mild conditions.^[2a,b] It was suggested that the active site consists of Cu steps that are decorated with Zn atoms and stabilized by bulk defects.^[2a,c]

Herein, the nature of the SMSI between ZnO_x and Cu under reducing conditions is investigated for an industrial-type methanol catalyst. A systematic examination using several methods has shown significant dynamical changes in the state of the copper surface as a consequence of an increasing partial pressure of hydrogen. This study elucidates the pitfalls of chemisorption methods for determining the Cu surface area and offer a more accurate determination of the specific Cu surface area together with strong evidence that Zn is present in its metallic state in a Cu/Zn surface alloy. Surface-sensitive methods provide information on the specific copper and zinc areas as a function of the partial pressure of hydrogen during the pretreatment. Furthermore, these methods provide quantitative information on the Zn oxidation state and the coverage of the Cu surfaces.

An industrial-type Cu/ZnO/Al₂O₃ catalyst was subjected to two experimental sequences in a combined HPC (high pressure cell) and UHV (ultra-high vacuum) setup and in a fixed-bed reactor system, respectively (Figure 1). In both

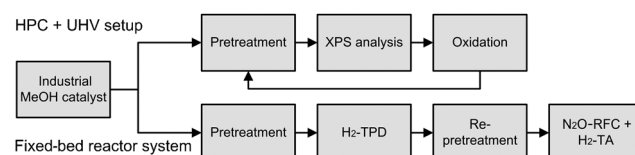


Figure 1. Block diagram of the experimental treatment sequences that were applied in this work. The upper route was used for the combined HPC and UHV system, whereas the bottom sequence was used for a fixed-bed reactor system.

systems, pretreatment of the catalyst consisted of a reduction in a H₂/He gas mixture. The temperature was increased by 2 K min⁻¹ from 300 to 493 K, which was followed by a 16 h dwell time at 493 K where the catalyst was exposed to different partial pressures of hydrogen.

In the combined HPC and UHV system, the Cu/ZnO/Al₂O₃ catalyst and pure ZnO powder (Transpek-Silox Industry) were tablettized and placed in a custom-made stainless-steel sample holder. Before characterization and pretreatment, the catalyst tablet was kept under UHV and heated to

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120 °C for two hours to remove water. After initial XPS, the catalyst was transferred to the HPC and exposed to the pretreatment. The surface was reset by heating the sample to 70 °C in an O₂/Ar atmosphere (5000 ppm) for 16 hours between the pretreatments.

The XPS characterization after the pretreatment was conducted to examine the extent to which metallic Zn is present in the catalyst (Supporting Information, Figure S1). It is not trivial to detect the reduction of ZnO to Zn by XPS. The Zn 2p_{3/2} lines for Zn and ZnO are positioned at 1021.4 eV and 1021.7 eV, respectively.^[8] The change in the valence state from Zn^{II} to Zn⁰ was observed as a more pronounced 3 eV downward shift of the binding energy of the Zn L₃M_{4,5}M_{4,5} Auger peak (Zn Auger; Figure S2b). In Figure 2a, the Zn

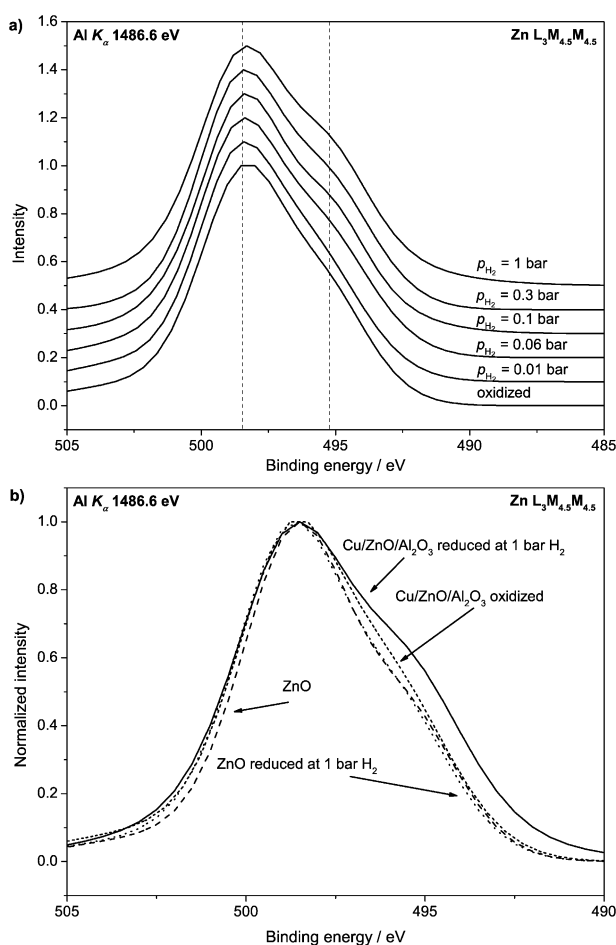


Figure 2. a) The Zn L₃M_{4,5}M_{4,5} Auger peaks for the oxidized and reduced Cu/ZnO/Al₂O₃ catalyst. b) Reference spectra of pure ZnO, reduced ZnO, and the Cu/ZnO/Al₂O₃ catalyst both in its oxidized and reduced form.

Auger peak is shown for the Cu/ZnO/Al₂O₃ catalyst after pretreatment with hydrogen pressures of 0.01–1 bar. In Figure 2b, the Zn Auger lines for the pure ZnO reference and the Cu/ZnO/Al₂O₃ system before and after pretreatment at 1 bar H₂ are displayed. The positions of all Auger peaks were aligned by the C 1s peak, and a Shirley background was subtracted. The Zn Auger peak from Cu/ZnO/Al₂O₃ in

Figure 2b is normalized to the peak height of the ZnO Auger feature. The Zn Auger spectrum shown in Figure 2 consists of a multiplex of many lines,^[9] but only the changes that are due to reduction were considered. In the Zn Auger spectrum of the Cu/ZnO/Al₂O₃ catalyst that was recorded after pretreatment under hydrogen atmosphere (0.01 bar), a shoulder feature appeared at a binding energy that was reduced by 3 eV (Figure 2a). The shoulder feature becomes more pronounced for higher partial pressures of hydrogen. When the pure ZnO tablet was reduced at 1 bar H₂ for 16 hours, no change of the Auger peak was observed (Figure 2b), which is in clear contrast to the results that were obtained with the Cu/ZnO/Al₂O₃ catalyst.

Analysis of the Cu L₃VV Auger peak shows that the Cu nanoparticles are reduced to the metallic state after each pretreatment (Figure S2). The O 1s XPS peak showed no change when the hydrogen partial pressure was varied during the pretreatment (Figure S3).

The Auger peaks in Figure 2 were fitted as a linear combination of the oxidized Auger peak, and the oxidized Auger peak shifted downwards by 3 eV in binding energy. According to the literature, the changes in the shoulder feature of the Zn Auger line could originate from both oxygen vacancies in ZnO thin films and metallic Zn in brass.^[10] The existence of oxygen vacancies was reflected by a shake-up feature of the O 1s peak by a 1.3 eV shift. As the O 1s XPS peak was unaffected by the pretreatment, the Auger shoulder feature may be attributed to metallic Zn. Therefore, a deconvolution of the fit can be used to quantify the amount of metallic Zn and ZnO (Figure S4a). In Figure 3a, the changes in the peak areas of Zn and ZnO are shown for different pretreatment conditions for both the Cu/ZnO/Al₂O₃ catalyst and the pure ZnO reference sample. A gradual enrichment with metallic Zn of up to 10 % was observed for the Cu/ZnO/Al₂O₃ catalyst subsequent to the pretreatment. When the ZnO reference sample was exposed to 1 bar of H₂, metallic Zn was not observed. These measurements demonstrate the importance of copper in industrial-type catalysts for catalyzing the reduction of ZnO.

In the fixed-bed reactor system, two-step series of experiments were applied for each set of pretreatment conditions. First, the Cu area was determined by H₂-TPD (Figure S5) subsequent to the pretreatment. H₂-TPD is known to be nondestructive with respect to the Cu surface area.^[11] Second, a re-pretreatment was followed by N₂O-RFC at 50 °C^[12] (Figure S6) and finally H₂ transient adsorption (H₂-TA) measurement using 1 % H₂/He at room temperature (Figure S7). The re-pretreatment was carried out to ensure that the N₂O-RFC was conducted on an adsorbate-free reduced Cu surface. After each series of experiments, the catalyst sample was replaced.

The Cu areas that were measured for the Cu/ZnO/Al₂O₃ catalyst after pretreatment with different partial pressures of hydrogen are shown in Figure 3b. The Cu area was determined by H₂-TPD, N₂O-RFC, and H₂-TA methods and was measured as the area of the Cu surface per catalyst weight. After pretreatment with 0.01 bar of hydrogen, the N₂O titration method and H₂-TPD essentially gave the same Cu area of approximately 16.5 m² g⁻¹. H₂-TA gave a slightly lower

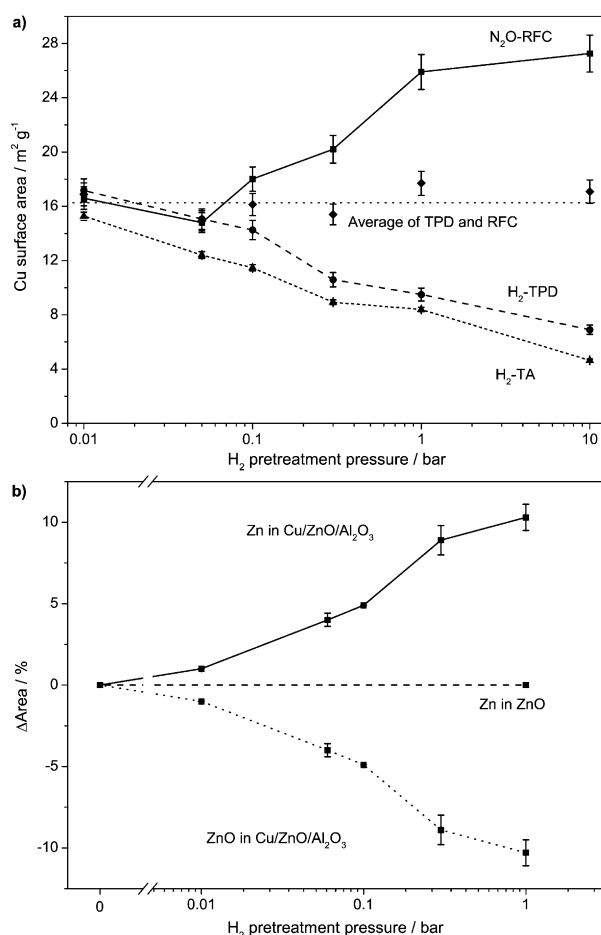


Figure 3. a) The change in the area of the ZnO (----) and Zn (—) Auger peaks for the industrial-type Cu/ZnO/Al₂O₃ catalyst and the change in the area for the ZnO reference sample (---). b) The change in the Cu surface area as determined by N₂O-RFC (■, —), H₂-TPD (●, ---), and H₂-TA (▲,), after pretreatment at different partial pressures of H₂ for 16 hours. For N₂O-RFC and H₂-TA, the error bars are the relative differences between two identical experiments. The error bars for H₂-TPD are 5%, as discussed in the Supporting Information. The average of the Cu areas as determined by H₂-TPD and N₂O-RFC is indicated by ♦. The horizontal dotted line is to guide the eye.

area (−13%). This is ascribed to incomplete reduction of Cu₂O (see the Supporting Information for details). The Cu area determined by N₂O-RFC was observed to increase with the partial pressure of hydrogen in the pretreatment above 0.05 bar. The surface area of the same catalyst, as determined by H₂-TPD and H₂-TA, decreases as a function of the hydrogen partial pressure. Despite the relatively low temperature during the pretreatment, the impact of the hydrogen partial pressure on the surface area as determined by the three different methods varies substantially, that is, up to ±60–65%. The average of the surface area based on the N₂O-RFC and H₂-TPD measurements displayed in Figure 3b (♦) is approximately constant and independent of the pretreatment.

Previous STM studies of Zn deposition on Cu(111) single crystals have shown that a Cu/Zn surface alloy can be formed by the random and homogeneous substitution of Zn atoms

into the Cu(111) surface layer.^[5a,b] The XPS data presented is in agreement with the formation of a Cu/Zn surface alloy. The shoulder feature of the Zn Auger line, which corresponds to metallic Zn, provides strong evidence for the reduction of ZnO to metallic Zn at the interface between the Cu and ZnO particles. The formation of a Cu/Zn surface alloy is also supported by the data that is shown in Figure 3b (see below).

The measurements of the Cu surface area by N₂O-RFC and H₂-TPD deviate to an increasing extent as the partial pressure of H₂ is raised during the pretreatment (Figure 3b). The determination of the Cu area by N₂O-RFC relies on the assumption that only Cu is present in the metallic state, that is, Cu is the only element that is responsible for the oxygen adsorption during N₂O decomposition. However, the oxidation process is not selective in nature. As indicated by XPS, the incorporation of reduced zinc atoms into the surface of the copper nanoparticles will increase the oxygen uptake from N₂O as each Zn atom that has been incorporated into the Cu surface is expected to react with one N₂O molecule, whereas one surface Cu atom only reacts with 0.5 N₂O molecules.

In contrast to the N₂O-RFC measurement, the H₂-TPD peak decreases with an increase in hydrogen partial pressure during the pretreatment. The H₂-TPD method is based on the specific desorption profile that is associated with metallic Cu only. Therefore, the incorporation of reduced Zn at the surface of the Cu particles leads to a decrease in the Cu area that is measured by the H₂-TPD method. Density functional theory (DFT) calculations show that the hydrogen adsorption energy at Zn sites that are incorporated into a Cu(111) surface will decrease by 28 kJ mol⁻¹ compared to that at a clean Cu(111) surface (Table S1). This implies that if Zn forms a surface alloy with Cu, a reduction in the peak area that is obtained by the H₂-TPD measurement can be directly related to a decrease in the Cu surface area and to the incorporation of Zn into the surface of the Cu particles.

The formation of a Cu/Zn surface alloy is supported by a comparison of the XPS data and the Cu areas determined by H₂-TPD. The comparison is based on a simple geometric model and uses the fraction of reduced and oxidized Zn atoms to calculate the loss of Cu surface atoms (Figure S10). In Figure 4a, the specific Cu surface area that was calculated from the XPS data as a function of the pretreatment H₂ pressure is compared to the H₂-TPD data, assuming that the latter only probes the Cu surface atoms. Figure 4a shows a clear correlation between the amount of metallic Zn measured by XPS and the decrease in the Cu surface area that was determined by H₂-TPD, which corroborates the hypothesis that H₂-TPD is an accurate method for determining the specific Cu (Zn-free) surface area.

The Cu surface area derived from the H₂-TA experiment was calculated by assuming that water is formed during the reduction of oxygen that is chemisorbed at the Cu surface (Figure S6).^[13] Despite the fact that the area determined by N₂O-RFC increases with the pretreatment pressure, the amount of H₂ that is consumed in the H₂-TA experiment decreases in the same manner as for the H₂-TPD experiment. This observation is very reasonable in the context of Cu/Zn surface alloy formation. The Zn enrichment in the Cu surface

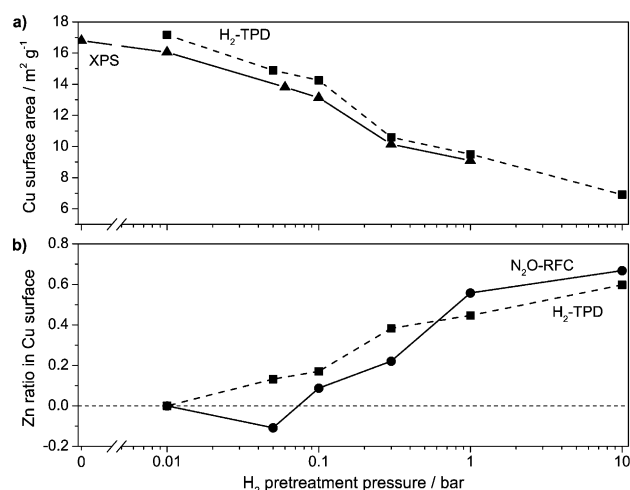


Figure 4. a) Correlation between the Cu surface areas that were determined by H_2 -TPD (■) and that are based on the XPS data (▲) by applying a simple model. b) The ratio of Zn enrichment in the Cu surface was evaluated for both the N_2O -RFC (●) and H_2 -TPD (■) techniques.

layer leads to the formation of ZnO during N_2O -RFC. The less reducible nature of ZnO prohibits H_2 consumption during the H_2 -TA experiment. The H_2 -TA experiment probes the amount of oxygen in the Cu_2O surface layer that is formed during the N_2O -RFC experiment, but not the amount of ZnO in the Cu surface (Figure S8).

The interaction between ZnO and Cu under reducing conditions has often been interpreted as a ZnO_x layer on the Cu surface.^[1a,4] Herein, we provide strong evidence that the identity of this layer is a Cu/Zn alloy. First, the position of the shoulder of the Zn Auger peak agrees well with that for reduced Zn in Cu, and the fact that the O 1s line is unaffected by reduction seems to exclude formation of a ZnO_x layer. Second, the results from the different techniques, that is, XPS, H_2 -TPD, N_2O -RFC, and H_2 -TA, corresponds to those expected for a Cu/Zn surface alloy. This agreement is highly unlikely in the case of a ZnO_x layer on top of the Cu particles. In situ TEM studies have not provided evidence for the formation of a ZnO_x layer on the Cu surfaces under reducing conditions,^[4b,7a] whereas a study that combined in situ TEM and electron energy loss spectroscopy (EELS) provided evidence for Cu/Zn alloy formation in a Cu/ZnO catalyst under CO/H_2 atmosphere.^[14] We therefore conclude that the results presented here provide strong evidence for the formation of a Cu/Zn surface alloy.

Based on this conclusion, the data in Figure 3b enables a quantification of the amount of metallic Zn in the Cu/Zn surface alloy. The relative change in the surface area that was measured after pretreatment at 10 mbar H_2 is interpreted as a change in the Zn/Cu element ratio in the surface of the Cu particles. In Figure 4b, the amount of Zn in the Cu surface was calculated from the H_2 -TPD and the N_2O -RFC experiment, respectively. The Zn enrichment in the Cu surface layer as a function of the pretreatment was seen to be similar even though two independent techniques were applied. The Zn enrichment in the Cu surface layer reached approximately

60% for a pretreatment of 16 hours at 10 bar H_2 . This observation is remarkable taking into account the relatively mild conditions that were applied during the pretreatment. Another interesting observation is that the average surface area measured by N_2O -RFC and H_2 -TPD is approximately independent of the pretreatment hydrogen pressure (see Figure 3b). This indicates that the total geometric surface area of the Cu particles is virtually unchanged during the reduction process, perhaps because wetting of Cu and ZnO particles of a similar size does not significantly change the exposed Cu surface area.

In conclusion, a study that combined XPS, H_2 -TPD, N_2O -RFC, and H_2 -TA measurements has provided clear evidence for an incorporation of metallic Zn atoms into the surface of Cu particles. Merging the results of these measurements gives insight into the debated origin of surface ZnO_x species. XPS revealed an enhancement of the metallic Zn feature in the Zn Auger peak with an increase in the partial pressure of H_2 . The presence of copper particles is found to be essential for the reduction of ZnO . These experimental observations are consistent with the formation of a Cu/Zn surface alloy during pretreatment. The interplay between the surface areas that were evaluated using N_2O -RFC and H_2 -TPD methods can be utilized to quantify the amount of metallic Zn in the Cu surface. The N_2O -RFC method has previously been used several times to determine the Cu surface area.^[15] This study provides a reinterpretation and a more accurate determination of the specific Cu surface, which is based on specific area measurements using four independent and consistent methods. Similar methods may be used for other metal-oxide systems where reduction of the oxide takes place. The presence of Zn atoms in a Cu surface has been shown experimentally to enhance the activity of a Cu surface,^[16] and more recently, this has also been justified theoretically.^[2a] We have now presented evidence that a surface alloy of Zn and Cu can be formed in industrial-type methanol synthesis catalysts under relatively mild conditions. These results support the idea that the presence of Zn atoms in the surface of Cu particles is important for increasing the activity of the Cu surface of a Cu/ZnO-based methanol synthesis catalyst.

Experimental Section

Measurements of the specific Cu areas were carried out in a fixed-bed flow setup. The system consisted of a gas-delivering system, a U-shaped reactor unit, and a gas analyzer, as previously described.^[11,17] Gases of high purity were used: He (99.9999%), H_2 (99.9999%), H_2/He (1% H_2 , 99.9995%), $\text{N}_2\text{O}/\text{He}$ (1% N_2O , 99.9995%). Fast online gas analysis was performed by a calibrated quadrupole mass spectrometer (Balzers GAM 445).

The Cu/ZnO/ Al_2O_3 catalyst that was used for the present experiments was an industrial-type Cu/ZnO/ Al_2O_3 catalyst (60:25:15; mol%) made by a co-precipitation method that is similar to a previously described procedure.^[18] A large sample of the catalyst was aged in dry synthesis gas ($\text{CO}/\text{CO}_2/\text{H}_2 = 5:5:90$) at 250 °C for 72 hours at very low space velocity and passivated in dilute oxygen. Samples of this catalyst were used in the present work. For each experiment, 200 mg of the aged catalyst with a sieve fraction of 150–300 μm were loaded into the U-shaped quartz reactor with an inner diameter of 4 mm. To determine the copper surface area, the procedure for H_2 -TPD experiments that was described by Muhler

et al. was followed.^[11] The copper surface area was equivalently determined by N₂O-RFC as reported by Chinchin et al.^[12]

The XPS measurements were performed in a UHV setup with a base pressure of 10^{−10} Torr. The catalyst could be transferred to an HPC for pretreatment. The pressure in the HPC was controlled by a pressure controller to be 1 bar, and the gas flow of He and H₂ or oxygen was controlled by mass flow controllers. For the XPS measurements, the X-rays were generated from a dual-anode X-ray source from SPECS using Al K_α radiation with a photon energy of 1486.6 eV, whereas the photoelectrons were measured with a Phoibos Hemispherical Energy Analyzer.

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