From Organometallic Zinc and Copper Complexes to Highly Active Colloidal Catalysts for the Conversion of CO₂ to Methanol

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ABSTRACT: A series of zinc oxide and copper(0) colloidal nanocatalysts, produced by a one-pot synthesis, are shown to catalyze the hydrogenation of carbon dioxide to methanol. The catalysts are produced by the reaction between diethyl zinc and bis(carboxylato/phosphinato)copper(II) precursors. The reaction leads to the formation of a precatalyst solution, characterized using various spectroscopic (NMR, UV–vis spectroscopy) and X-ray diffraction/absorption (powder XRD, EXAFS, XANES) techniques. The combined characterization methods indicate that the precatalyst solution contains copper(0) nanoparticles and a mixture of diethyl zinc and an ethyl zinc stearate cluster compound [Et₄Zn₅(stearate)₆]. The catalysts are applied, at 523 K with a 50 bar total pressure of a 3:1 mixture of H₂/CO₂, in the solution phase, quasi-homogeneous, hydrogenation of carbon dioxide, and they show high activities (>55 mmol/gZnOCu/h of methanol). The postreaction catalyst solution is characterized using a range of spectroscopies, X-ray diffraction techniques, and transmission electron microscopy (TEM). These analyses show the formation of a mixture of zinc oxide nanoparticles, of size 2–7 nm and small copper nanoparticles. The catalyst composition can be easily adjusted, and the influence of the relative loadings of ZnO/Cu, the precursor complexes and the total catalyst concentration on the catalytic activity are all investigated. The optimum system, comprising a 55:45 loading of ZnO/Cu, shows equivalent activity to a commercial, activated methanol synthesis catalyst. These findings indicate that using diethyl zinc to reduce copper precursors in situ leads to catalysts with excellent activities for the production of methanol from carbon dioxide.

KEYWORDS: hydrogenation of CO₂, CO₂ reduction, methanol synthesis, colloidal catalysts, Cu-ZnO catalysts, nanoparticles, nanocatalysts, catalysts from organometallic

INTRODUCTION

Using waste CO₂ is a high priority: among the most promising options are the synthesis of polymers and the reduction to alcohols which may serve as fuels.¹ By applying renewable or off-peak power to make hydrogen, for example, via electrolysis of water, it is envisaged that carbon dioxide hydrogenation to methanol could reduce our dependence on fossil fuels.² Methanol is an attractive energy vector and a promising renewable transport fuel, particularly because its use is compatible with existing fuel infrastructures.³ In China, methanol is already being applied, in blends with gasoline, as a fuel.⁴

Methanol is currently produced on a huge scale (>40 million tonnes, globally in 2009), mostly from synthesis gas (CO/H₂), generated by coal gasification or methane reforming, converted using ternary heterogeneous Cu/ZnO/Al₂O₃ catalysts.⁵ There is also a good precedent for these same catalysts to be used for the hydrogenation of carbon dioxide.⁶ They are generally prepared by metal salt coprecipitation, following by calcination, aging, nanostructuring, and reduction cycles.⁷ Controlling the conditions for these steps is essential to maximize catalyst activity. Detailed investigations indicate that maximizing the ZnO/Cu interface is crucial, through controlling the Zn/Cu loading, introducing a suitable support, and ensuring intimate mixing of the phases, on the nanometre scale.⁸,⁹ However, current synthesis methods require the preformation of mineral phases, most commonly zinccian malachite, which fundamentally limit the loading of Zn/Cu (in the case of zinccian...
malachite, to 3:7) and predetermine the products of aging and, hence, the extent and quality of the interface. Furthermore, such coprecipitation routes require forcing conditions to decompose the mineral, thereby limiting the nanomorphology of the catalyst. Recent elegant structural studies of these ternary catalysts have revealed that some surface sites are dynamic. For these reasons, there is an impetus to develop new catalyst syntheses. For example, Tsang and co-workers have mixed (in the solid state) isolated nanoparticles of ZnO and Cu to produce high-activity CO₂ hydrogenation heterogeneous catalysts and have correlated activity to exposure of particular crystal facets. Methanol can also be efficiently produced by slurry phase methods, whereby the active catalyst is dissolved or suspended in a high-boiling solvent (such as squalane). Of particular interest is the use of inorganic/organometallic precursor compounds to produce well-defined, quasi-homogeneous nanoparticles for catalysis. The groups of Fischer and Schüth have pioneered the application of colloidal solutions of nanoparticles (Cu and/or ZnO) as high-activity catalysts for such solution phase syn-gas to methanol processes. Simon et al. reported, as early as 1988 that the reaction between Cu(acac)₂ and Et₂Zn in the presence of 1,3-butadiene in an organic solvent formed an efficient syn-gas catalyst system with a high selectivity for ethanol and methanol formation. Corker and Evans used EXAFS to investigate the precatalyst solution. They proposed it comprised a mixture of colloidal copper and an oxozinc cluster compound (of undefined structure and stoichiometry); however, detailed characterization of the cluster and changes to the system during and after catalysis were not carried out.

In the past 10 years, Fischer and co-workers have pioneered various routes to catalytic nanoparticles, including hydrothermal decomposition, hydrolysis, photochemical decomposition, chemical vapor deposition (CVD), and atomic layer deposition (ALD), using organo-Zn and Cu precursors. In 2005, Schüth and co-workers applied triocetyl aluminum reagents as the reductants and ligands for copper nanoparticles; these systems showed high activities for liquid phase methanol catalysis. Fischer and Mühler have also applied colloidal nanoparticles of ZnO/Cu prepared by hydrothermal decomposition of well-defined inorganic compounds, some of which show excellent activities for the liquid phase synthesis of methanol from syn-gas. In 2006, they reported a high-activity system prepared by the addition of diethyl zinc to a hot (200 °C) solution of a copper(II) alkoxide precursor. This system formed small, unsupported nanoparticles of both Cu and ZnO and showed up to 85% of the activity, in a squalane suspension, of the ternary reference. Other alkyl zinc alkoxides were also shown to undergo thermolysis in the presence of amine surfactants to yield quasi-homogeneous ZnO nanoparticles. Subsequently, Fischer and co-workers reduced Cu/Zn stearate mixtures with hydrogen gas to produce copper nanoparticles, stabilized by zinc stearate.

Our group has focused on CO₂ rather than syn-gas as the carbon source. An alternative zinc oxide synthesis involving the hydrolysis of diethyl zinc, in the presence of substoichiometric quantities of zinc carboxylates/phosphines produces 3–5 nm ZnO nanoparticles which are soluble in various organic solvents/polymer. A high-activity carbon dioxide hydrogenation catalyst system was prepared by combining solutions of ZnO nanoparticles, surface coordinated by di(ocetyl)-phosphininate groups, with copper nanoparticles, surface-coordinated with stearate groups. However, the optimum composition and nature of the precatalyst solution remains under-investigated. Furthermore, uncovering the relative importance of particle sizes, oxidation states of the metals, and interfaces between particles are central to improving catalytic activity. One drawback of this system is the requirement to pre-prepare separate solutions of the nanoparticles. A “one-pot” method to prepare the catalyst would be desirable, as would be the replacement of hydrazine as the reductant in the preparation of copper nanoparticles. This paper presents a “one-pot” approach to catalyst synthesis, applying diethyl zinc as both the reductant to produce copper nanoparticles and the precursor to zinc oxide (by hydrolysis of Zn–C bonds). This method provides highly active carbon dioxide reduction catalysts, and through characterization of the pre- and postcatalyst solutions, the effects of organometallic precursor compound, relative loading, and metal oxidation state can be explored.

RESULTS AND DISCUSSION

Precatalyst Synthesis and Characterization. An in situ, one-pot catalyst preparation method that involved mixing together equimolar quantities of copper(II) bis(stearate) and diethyl zinc in toluene at 298 K (Scheme 1) was discovered. This solution showed several color changes consistent with the reduction of the copper(II) precursor and formation of a red/brown solution and characteristic of Cu(0) nanoparticles. It is proposed that ligand exchange occurs between the copper and zinc complexes, leading to transient formation of copper–alkyl species which are known to undergo rapid decomposition at room temperature to produce copper(0) species. Ligand exchange would also be expected to lead to the formation of some zinc carboxylate species; such species have recently been shown to react with any excess dialkyl zinc to form heteroleptic alkyl zinc carboxylate complexes. The characterization data (vide infra) support this proposed ligand exchange and indicate the reaction proceeds to the formation of a solution containing copper(0) nanoparticles and well-defined zinc organometallic complexes.
The UV–vis spectrum of the red solution confirmed the formation of copper nanoparticles, with a copper surface plasmon absorbance observed at 580 nm (Figure S1).17 The spectrum also clearly indicated the complete consumption of the Cu(II) precursor complex, as shown by the disappearance of its characteristic absorbance at 700 nm.

The speciation of the zinc was revealed by 1H NMR spectroscopy (Figure S2), which showed the complete consumption of the paramagnetic species (e.g., Cu(II)) and the formation of well resolved signals consistent with the presence of diethyl zinc and an ethyl zinc stearate cluster compound, ZnEt2(stearate)6, the structure of which has been reported previously.14b,d Analysis of the peak integrals in the 1H NMR spectrum revealed a 3:2 molar ratio of Et2Zn/[Et2Zn(stearate)]6 (Figure S2). We have also previously established that the hydrolysis of mixtures of diethyl zinc and the zinc stearate cluster, at analogous molar loadings, leads to the efficient formation of 3–5 nm zinc oxide nanoparticles, capped with stearate ligands.14c

The precatalyst solution was also analyzed by XAFS measurements, conducted in toluene solutions, under an inert atmosphere. The copper speciation was studied (Figure 1) by comparing the Cu K-edge X-ray absorption near edge structure (XANES) spectra of the catalyst colloids with appropriate reference compounds, including Cu(stearate)2, CuO, Cu2O, and Cu metal. The precatalyst solution spectrum shows features similar to the Cu foil reference, implying the presence of metallic copper. Moreover, the XANES spectra lacked any features typically associated with oxidic forms of Cu, primarily the peak at 8983.5 eV, indicative of linear Cu2O (1s → 4p), and the peak at 8986.5 eV, which is evidence for CuO (1s → 4p).18a The main edge peaks are less well resolved than those of the copper foil, likely due to the nanoparticle size. Theoretical XANES simulations and experimental measurements of copper clusters have shown related effects, attributed to the absence of higher coordination shells and to a reduced particle size.18

The EXAFS data and calculated fitting parameters are evidence of the formation of copper nanoparticles and are illustrated in Figure S3 and Table S1. These data also indicate that the precatalyst solution contains small copper nanoparticles, again with no evidence for the formation of either of the oxides. The first-shell Cu–Cu coordination number can be used to estimate the particle size, according to the method developed by Beale et al.19 The estimated mean particle size, from the EXAFS data, is 1.2 nm, consistent with the plasmon absorption observed in the UV–vis spectrum.20

The zinc XANES spectrum (Figure 2) of the precatalyst solution shows features in the absorption edge that are consistent with the presence of both bis(stearate) zinc and diethyl zinc. A linear combination fit of the XANES data was calculated using Zn(stearate)2 and diethyl zinc as reference compounds (Figure S4). A reasonable fit was obtained, having an R factor of 0.0025, based on a composition of ~84% diethyl zinc and ~16% Zn(stearate)2. This estimation shows close agreement with the relative ratio of ethyl/stearate groups (~86% of signals are due to zinc bound ethyl groups; ~14% of signals, to zinc bound stearate moieties). It is apparent that the anaerobic techniques were sufficiently robust because the organozinc carboxylate cluster compound (Et4Zn5(stearate)6) was successfully detected, a compound that is known to hydrolyze readily in the presence moisture.14b,d Furthermore, the zinc carbon bond lengths determined by analysis of the integrals for Et2Zn and Et4Zn5(stearate)6 in the 1H NMR spectrum (~86% of signals are due to zinc bound ethyl groups; ~14% of signals, to zinc bound stearate moieties). The large Debye–Waller factor, compared with diethyl zinc, is also indicative of other species having Zn–C or Zn–O bonds, as would be expected from the ethyl zinc stearate cluster compound.

X-ray diffraction measurements were conducted on the precatalyst solution (after removal of all volatiles including diethyl zinc) (Figure S6). The samples are only weakly
Some oxidation peaks (for CuO and Cu₂O) are observed, likely as a result of air exposure during sample preparation and measurement. The average copper particle size, determined for the Cu (111) peak using the Scherrer equation, is 4.8 nm, somewhat larger than the value obtained by EXAFS. However, EXAFS is more sensitive to detection of smaller clusters and particles, and XRD is volume-weighted to larger particles; therefore, a difference in the obtained particle size between the two measurements is to be expected in a polydisperse system. The XRD measurements also showed additional peaks at low angle, which correspond to signals expected for zinc-bound stearate groups, consistent with the presence of the ethyl zinc stearate cluster.

The combined spectroscopic data, therefore, indicate that the red precatalyst solution contains a mixture of 1–5 nm copper nanoparticles, presumably surface-capped with stearate ligands, and a mixture of diethyl zinc and ethyl zinc stearate clusters. Britten and co-workers have previously studied the reaction between diethyl zinc and Cu(II) precursors. They established that diethyl zinc rapidly and irreversibly reduced the Cu(II) species to Cu, which formed as a mirror. This reaction was attributed to rapid ligand exchange and the instability of the species to Cu, which formed as a mirror. This reaction was attributed to rapid ligand exchange and the instability of the species to Cu, which formed as a mirror.

Carbon Dioxide Hydrogenation Catalyst. Catalyst Composition. The precatalyst solution was injected into a degassed solution of squalane in a continuously stirred tank reactor (CSTR), under an atmosphere of N₂ at 298 K. A 3:1 H₂/CO₂ mixture, at 50 bar, was added, and the reactor was heated to 523 K. The reaction was monitored using an in-line GC to detect the formation of methanol. During the first 120 min, the quantity of methanol produced steadily increased (a phenomenon partly resulting from reactor parameters, which introduce a delay in methanol detection by the GC instrument), reaching a steady state after ~2 h and remaining at this level for the duration of each experiment (12 h). Initially, ethane was also detected, but production ceased by 5 h (Figure S7). It is proposed that the ethane evolution results from the reaction of zinc ethyl groups, which are formed by carbon dioxide hydrogenation, and is present in sufficient quantities in the gases/solvents. Indeed, a control experiment in which the precatalyst solution was exposed only to a continuous nitrogen flow in the reactor also resulted in ethane evolution over 5 h, after which time the introduction of the reaction gases (CO₂/H₂) resulted in a catalyst system showing equivalent activity. Regardless of the source of the moisture, the hydrolysis reaction is expected to generate the ZnO nanoparticles, which in combination with the copper nanoparticles comprise the active catalyst (vide infra).

The method was applied to prepare a series of precatalyst solutions, using different molar ratios of diethyl zinc/copper-stearate₂ precursors, while keeping the overall loading of metals constant. The catalytic activities were benchmarked against the total mass of Cu and ZnO present, deduced from the molar quantities of the precursors and assuming complete conversions. The activities obtained using the precatalyst solutions were high and also very reproducible. For example, when a 1:1 molar mixture of diethyl zinc and copper stearate (55:45 weight ratio of ZnO/Cu) was applied, a methanol activity of 60 mmol g⁻¹ h⁻¹ (133 mmol g⁻¹ h⁻¹) was obtained (Table 1, Figure S7). This activity is similar to that obtained using a heterogeneous ternary syn-gas catalyst (preactivated according to the standard protocols) as a benchmark material; however, after 10 h, the best catalyst system showed activity superior to the ternary heterogeneous catalyst control. These findings underscore the stability of the colloidal catalyst because the peak activity does not diminish significantly over the run (12 h), unlike the reference system.

It is clear that activity is strongly dependent on the relative quantities of zinc oxide and copper, at a fixed overall metal concentration (Figure S8). Optimum activities were observed for a 1:1 molar ratio of ZnEt₂/Cu(stearate)₂ precursors, which corresponds to an expected weight ratio of 55:45 ZnO/Cu, assuming complete conversions to the relevant oxide/metal nanoparticles (as indicated by the spectroscopic data). The activity rapidly decreased when either an excess of diethyl zinc (therefore, an increase in the ZnO loading) or an excess of copper stearate was applied. On one hand, sufficient diethyl zinc is required to reduce the copper completely; on the other hand, sufficient stearate (initially coordinated to the copper precursor) is required for nanoparticle stabilization. The colloidal nanoparticle catalysts show good stability over 12 h of the run, with activities remaining at or only slightly lower than the peak value for the duration of the run in the best case. In contrast, the ternary heterogeneous suspension shows a significant loss in activity after 12 h on-stream. Furthermore, in cases where an excess of diethyl zinc is added, significant deactivation is also observed. This is tentatively attributed to a reduction in the stability of the colloidal particles due to the reduced amount of stearate capping ligand available to stabilize the particles.

Table 1. Effect of Varying the Weight Ratio ZnO/Cu on the Catalytic Activity for Carbon Dioxide Hydrogenation to Methanol

<table>
<thead>
<tr>
<th>Entry</th>
<th>Molar ratio precursors: ZnEt₂/Cu(stearate)₂</th>
<th>Expected weight ratio ZnO/Cu</th>
<th>Methanol peak activity after 12 h, mmol g⁻¹ h⁻¹</th>
<th>Methanol activity after 12 h, mmol g⁻¹ h⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1:3</td>
<td>30:70</td>
<td>39</td>
<td>39</td>
</tr>
<tr>
<td>2</td>
<td>1:1</td>
<td>55:45</td>
<td>60</td>
<td>58</td>
</tr>
<tr>
<td>3</td>
<td>2:1</td>
<td>70:30</td>
<td>43</td>
<td>37</td>
</tr>
<tr>
<td>4</td>
<td>7:1</td>
<td>90:10</td>
<td>18</td>
<td>13</td>
</tr>
<tr>
<td>5</td>
<td>Ternary benchmark* (expected weight ratio of ZnO/Cu = 1:2)</td>
<td></td>
<td>60</td>
<td>46</td>
</tr>
</tbody>
</table>

*Alfa Aesar ternary methanol synthesis catalyst (product code: 45776), comprising (by weight) 24.7% ZnO, 63.5% CuO, 11.4% Al₂O₃, and MgO; see ESI for catalyst preactivation details and activity calculations. Error for each measurement is ±4.5% as determined by runs in triplicate. Figures S7 and S8 illustrate the stability of the catalysts showing activity vs time. Reaction conditions: 523 K, 50 bar (3:1, H₂/CO₂), in a solution of squalane/toluene (90:10) at a fixed total volume of 100 mL, a total gas flow of 166 mL min⁻¹, over 16 h. The total catalyst concentration (ZnO + Cu) was constant at 0.20 g dm⁻³.
Catalyst Concentration. The influence of overall nanoparticle concentration in the reactor was investigated at the optimum ratio of ZnO/Cu, 55:45 (1:1, Et₂Zn/Cu(stearate)) (Figure 3 and Table S3). As the nanoparticle concentration increases from 0 to 0.2 gZnOCu/L, the activity also steadily increases, presumably as the probability of an interaction between particles increases, allowing the formation of a catalytically active interface between ZnO and Cu. However, at nanoparticle concentrations above 0.2 gZnOCu/L the activity decreases. At the relatively higher concentrations, aggregation of the nanoparticles appears to increase, presumably reducing the availability of active sites (which must be on the catalyst surface) and thereby reducing activity.22

Influence of the Organometallic Precursor Complexes. Various copper(II) and di(organo)zinc precursor complexes were investigated to establish the generality of the protocol to synthesize the catalyst (Table 2, Figure S10). All the complexes resulted in the formation of effective catalysts for methanol synthesis, with the activity depending weakly on the nature of the precursor. In all cases, the catalysts operated in the same manner, with the evolution of ethane and formation of methanol over the first 5 h, followed by a steady state methanol production that does not significantly decrease over the following 12 h (Figure S10). Substituting diethyl zinc with diphenyl zinc, using bis(stearate) copper(II) as the copper source reduced the catalytic activity by ~25%. This reduction is tentatively attributed to differences in the reactivity of diphenyl zinc, both toward the Cu(II) precursor and toward water during the formation of zinc oxide in the reactor.

The influence of the copper precursor on catalytic activity, using diethyl zinc as the zinc oxide source, was more subtle. Using bis(laurate) or bis(octanoate) copper(II) precursors resulted in slightly more active catalyst than using bis(stearate) copper(II). It is likely that this effect relates to the balance between optimum solubility of the nanoparticles and steric hindrance and protection of the surface active sites. Thus, the optimum C-chain length for these carboxylate capped nanoparticles appears to be 8 (octanoate).

We have previously reported a catalyst system prepared by mixing solutions of preformed nanoparticles of ZnO/Cu.15 In that system, it was observed that increasing the reductive stability of the zinc oxide nanoparticle capping ligand by substituting carboxylate for di(octyl)phosphinate resulted in increased catalytic activity.15 In this study, changing the copper precursor to di(octyl)phosphinate has a much lesser influence on activity, resulting in only a small increase. The Et₂Zn/Cu(di(octyl)phosphinate)₂ precursor was also tested at the 0.2 gZnOCu/L catalyst concentration (optimum for stearate), resulting in a peak activity of 60 mmol gZnOCu⁻¹ h⁻¹, which is close to the value for the Cu(stearate)₂ precursor (Figure S11).

Characterization. The optimum precatalyst solution was formed by the in situ reduction of copper stearate/phosphinate complexes with diethyl zinc. TEM analysis showed small nanoparticles (5–10 nm in diameter) in agglomerates containing from two to hundreds of nanoparticles (Figure 4). Some agglomeration is likely to occur during the catalyst precipitation, washing, and deposition required for TEM sample preparation, but the primary nanoparticles do appear to be intrinsically fused to form small clusters (Figure 4). Measurements of the lattice spacings in the high-resolution TEM images reveal that the nanoparticles consist of ZnO, Cu₀, and Cu₂O (Figure 4). Some nanoparticles showed a spacing of only ~2.47 Å, which corresponds to either ZnO(101) or Cu₂O(111). Such nanoparticles cannot be identified unambiguously. A lattice is not resolved in the remaining unidentified nanoparticles. Phases with larger lattice spacings are easier to resolve and, hence, easier to identify. ZnO(100) corresponds to a spacing 35% larger than Cu(111), which may be why the concentration of ZnO appears higher.

Table 2. Influence of the Organometallic Precursor Complexes on Methanol Catalytic Activity*4

<table>
<thead>
<tr>
<th>copper precursor</th>
<th>zinc precursor</th>
<th>methanol peak activity, mmol gZnOCu⁻¹ h⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu₂(stearate)₂</td>
<td>ZnEt₂</td>
<td>42</td>
</tr>
<tr>
<td>Cu₂(stearate)₂</td>
<td>ZnPb₂</td>
<td>32</td>
</tr>
<tr>
<td>Cu₂(laurate)₂</td>
<td>ZnEt₂</td>
<td>43</td>
</tr>
<tr>
<td>Cu₂(octanoate)₂</td>
<td>ZnEt₂</td>
<td>44</td>
</tr>
<tr>
<td>Cu₂(2-ethyl hexanoate)₂</td>
<td>ZnEt₂</td>
<td>39</td>
</tr>
<tr>
<td>Cu₂(diocyl phosphate)₂</td>
<td>ZnEt₂</td>
<td>50</td>
</tr>
</tbody>
</table>

*4Reaction conditions: 523 K, 50 bar (3:1, H₂/CO₂), in squalane/toluene (90:10) at a fixed total volume of 100 mL, a total gas flow of 166 mL min⁻¹, over 16 h. The weight ratio of ZnO/Cu was fixed at 55:45, assuming complete conversion of the precursor complexes, and the overall catalyst concentration (ZnO + Cu) was fixed at 0.4 gZnOCu/L. Error for each measurement is 4.5% as determined by runs in triplicate. Figure S10 shows the activity vs time data for all catalysts.
copper nanoparticles. The presence of a contact interface between Cu-containing and ZnO nanoparticles is clearly observed in Figure 4.

The Zn K-edge XANES spectrum of the postcatalysis solution (Figure 2) also indicates that ZnO is present. The main edge peak observed at 9669 eV and a feature at 9663 eV are entirely consistent with the 1s → 4pₓ,ᵧ transitions of ZnO.25 The Cu K-edge XANES spectrum of the same postcatalysis mixture indicates that the major copper-containing species continue to be Cu(0) nanoparticles (Figure 1). Indeed, the spectrum is essentially the same as that for the precatalysis mixture. On the other hand, the EXAFS data for the postcatalysis copper species demonstrate a small component of oxidic Cu species (as evidenced by the feature at ∼1.8 Å in the Fourier transform data) as well as the Cu(0) species. The distance (1.84 Å) is consistent with a Cu(I)−O bond length, suggesting there may be minor contamination by Cu₂O in the postcatalysis mixture, likely due to an increase in nanoparticle size after catalysis; however, because of the mixed speciation of the copper postcatalysis, it is not possible to accurately quantify the copper nanoparticle size.

Powder X-ray diffraction measurements, conducted on the samples after centrifugation, also confirmed the presence of ZnO and copper nanoparticles in the postreaction mixtures. Figures S13−S15 illustrate the XRD patterns obtained for different loadings of ZnO/Cu; an illustrative example at 55:45 ZnO/Cu is shown in Figure 5. In every case, there are clearly peaks that can be indexed against the patterns expected for ZnO and copper nanoparticles. In most spectra, some copper oxidic species are also present (both Cu₂O and CuO), although the oxidation is proposed to occur predominantly during sample preparation, because the quantities present differ nonuniformly, and in one case, these species are absent (30:70, ZnO/Cu). The Scherrer equation can be used to estimate the relative particle sizes, and Table S4 lists these values for the various different loadings. In all cases, ∼7 nm ZnO nanoparticles are estimated; on the other hand, the size of the Cu particles appears rather more variable with low loadings of either ZnO/Cu (relating to lower molar quantities of either Et₂Zn or Cu(stearate)₂), resulting in the formation of significantly larger copper nanoparticles (∼12−15 nm). When the relative loading of ZnO/Cu is more closely balanced (e.g., 55:45), then the sizes of the nanoparticles are also close, both being ∼7 nm. This value is slightly higher than in the precatalyst solution, in line with the TEM images. It is tentatively proposed that the matching between the ZnO and Cu nanoparticle sizes may in part rationalize the improved activity shown at these loadings, possibly by maximizing the active interface between the two components.

**CONCLUSIONS**

Highly active, quasi-homogeneous CO₂ hydrogenation catalysts were produced by reaction between discrete inorganic complexes, namely, bis(carboxylate) copper(II) and diethyl zinc, to form precatalyst solutions. These solutions were applied (at S23 K, 50 bar of a 1:3 mixture of CO₂/H₂) to produce methanol in a continuously stirred slurry reactor.
(squalane) with in-line GC detection equipment. The pre- and postcatalysis solutions were characterized using UV–vis, EXAFS, and XANES spectroscopy; microscopy (TEM); and, where possible, by 1H NMR spectroscopy. The combined techniques indicate that the precatalysis mixture contains small (1–4 nm) copper nanoparticles and organometallic zinc complexes (diethyl zinc and ethyl zinc stearate cluster). On the other hand, after catalysis, the organozinc reagents were hydrolyzed to produce zinc oxide nanoparticles and the copper nanoparticles of sizes <10 nm. These spectroscopic findings are in line with ethanol evolution (consistent with ethyl–zinc bond hydrolysis) during the initial phase of catalysis. The catalyst synthesis was easily tuned, and the influence of the relative loadings of ZnO/Cu, the nature of the precursor compounds, and the effect of overall catalyst concentration were reported.

The best systems, which were formed using equimolar ratios of diethyl zinc/bis(steareate) copper in dry toluene gave activities for methanol production equivalent to those observed with a commercially available heterogeneous catalyst benchmark (mostly comprising ZnO, CuO, and Al2O3 mixtures). Thus, it is clear that formation of the copper nanoparticles, by reduction with diethyl zinc and subsequent in situ hydrolysis of the organozinc complexes results in a highly active and selective catalyst mixture. It is tentatively proposed that this increased activity may result from a preferential formation of ZnO/Cu catalyst mixture. It is tentatively proposed that this increased activity may result from a preferential formation of ZnO/Cu interfaces using the in situ methodology (as compared with mixing together discrete solutions of nanoparticles). The facility with which variables can be changed as well as the ease of catalyst preparation make this route a rather attractive method with which variables can be changed as well as the ease of mixing together discrete solutions of nanoparticles. The facility

ASSOCIATED CONTENT

Supporting Information

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Experimental details, 15 figures, 4 tables of results; additional references (PDF)

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The authors declare no competing financial interest.

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