Copper sulfide derived nanoparticles supported on carbon for the electrochemical reduction of carbon dioxide

van Oversteeg, Christina H.M.; Tapia Rosales, Marisol; Helfferich, Kristiaan H.; Ghiasi, Mahnaz; Meeldijk, Johannes D.; Firet, Nienke J.; Ngene, Peter; de Mello Donegá, Celso; de Jongh, Petra E.

DOI
10.1016/j.cattod.2020.09.020

Publication date
2020

Document Version
Final published version

Published in
Catalysis Today

Citation (APA)
van Oversteeg, C. H. M., Tapia Rosales, M., Helfferich, K. H., Ghiasi, M., Meeldijk, J. D., Firet, N. J., Ngene, P., de Mello Donegá, C., & de Jongh, P. E. (2020). Copper sulfide derived nanoparticles supported on carbon for the electrochemical reduction of carbon dioxide. Catalysis Today. https://doi.org/10.1016/j.cattod.2020.09.020

Important note
To cite this publication, please use the final published version (if applicable). Please check the document version above.

Copyright
Other than for strictly personal use, it is not permitted to download, forward or distribute the text or part of it, without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license such as Creative Commons.

Takedown policy
Please contact us and provide details if you believe this document breaches copyrights. We will remove access to the work immediately and investigate your claim.
Copper sulfide derived nanoparticles supported on carbon for the electrochemical reduction of carbon dioxide

Christina H.M. van Oversteeg a,b,1, Marisol Tapia Rosales a,1, Kristiaan H. Helfferich a, Mahnaz Ghiasi a, Johannes D. Meeldijk a, Nienke J. Firet a, Peter Ngene a, Celso de Mello Donega b, Petra E. de Jongh a,c

a Inorganic Chemistry and Catalysis, Debye Institute for Nanomaterials Science, Utrecht University, Universiteitsweg 99, 3584 CG, Utrecht, the Netherlands
b Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, Princetonplein 1, 3584 CG, Utrecht, the Netherlands
c Materials for Energy Conversion and Storage (MECS), Department of Chemical Engineering, Faculty of Applied Sciences, Delft University of Technology, van der Maasweg 9, 2629 HZ, Delft, the Netherlands

ARTICLE INFO

Keywords:
Copper sulfide
Nanoparticles
Carbon support
Sulfidation
CO2 reduction
In situ X-ray absorption spectroscopy

ABSTRACT

The electrocatalytic reduction of CO2 to produce sustainable fuels and chemicals is attracting great attention. Cu-based catalysts can lead to the production of a range of different molecules, and interestingly the product selectivity strongly depends on the preparation history, although it is not fully understood yet why. We report a novel strategy that allowed us to prepare Cu nanoparticle on carbon catalysts with similar morphologies, but prepared by in-situ reduction of either supported CuS, Cu2S or CuO nanoparticles. For the first time the evolution of the Cu species was followed under CO2 and H+ reduction conditions using in-situ X-ray absorption spectroscopy. Excellent electrochemical contact between the Cu-based nanoparticles, the carbon support and the carbon-paper substrate was observed, resulting in metallic Cu as the predominant phase under typical electrochemical CO2 reduction conditions. Even covering less than 4% of the H2 producing carbon support with Cu-sulfide derived nanoparticles allowed to steer the selectivity to a maximum of 12% Faradaic efficiency for the production of formate. Clear differences between the catalysts derived from CuS, Cu2S or CuO nanoparticles were observed, which was ascribed to the presence of residual sulfur in the catalysts.

1. Introduction

When using renewable electricity, the electrochemical reduction of carbon dioxide (CO2) provides a promising route to produce chemicals and fuels in a sustainable manner. [1,2] The field of so called “solar fuels” has triggered the interest of many researchers, focusing on the development and understanding of electrocatalysts that can promote the electrochemical CO2 reduction efficiently and with high selectivity to the targeted product.

The majority of the electrocatalysts studied for the reduction of CO2 are based on transition metals. [3,4] As already described by Hori et al. in 1985, the catalytic performance of a catalyst highly depends on the metal used. [5] Metals such as gold [6,7], silver [8,9] and zinc produce mainly CO2, whereas metals such as platinum, nickel and iron reduce only small amounts of CO2, leading to H2 as the main product formed via the competing hydrogen evolution reaction.

Copper electrodes are extensively studied and stand out because of their unique ability to produce hydrocarbons and oxygenates, which is ascribed to their intermediate binding strength for the CO intermediate. [1,5,10] Interestingly, Cu can lead to a range of different H+ and CO2 reduction products, and much work is done in the field to obtain a better understanding of the catalytic activity of Cu and the selectivity of Cu electrodes. For example, the use of oxide-derived Cu electrodes promotes the production of CO and COOH at low overpotentials, even though the electrodes are operating at potentials where all oxide should be reduced to metallic copper. [11,12] The exact explanation for the influence of the origin of the copper electrodes remains under debate, it might be attributed to the formation of specific surface structures of these oxide-derived electrodes [1], although also some oxygen remaining in the material might be a factor. [13]

https://doi.org/10.1016/j.cattod.2020.09.020
Received 30 January 2020; Received in revised form 23 August 2020; Accepted 23 September 2020
Available online 1 October 2020
0920-5861/© 2020 The Authors. Published by Elsevier B.V. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).

Please cite this article as: Christina H.M. van Oversteeg, Catalysis Today, https://doi.org/10.1016/j.cattod.2020.09.020
More recently, several experimental and theoretical studies have demonstrated that the addition of sulfur modifies the performance of Cu catalysts. [14–19] Even though copper sulfide is unstable under CO2 and H2 reduction conditions, most studies show that some sulfur remains on the catalyst after reaction. [14,15,17,18,20,21] Many of these studies observed an increased selectivity towards formate, that is often attributed to a change in the binding energy between the sulfur-containing catalyst and key intermediates in CO oxidation.[14,15,17,18,20,21] Many of these studies indicate an increased production of CO and/or CH4 at low overpotentials when using sulfur derived Cu electrodes. [20,22] In addition, Zhuang et al. reported the production of ethanol, propanol and ethylene at intermediate overpotential (-0.95 V vs. RHE) using Cu2S derived nanoparticles. [21]

Inspired by these studies, in this work we investigated the catalytic performance and stability of copper sulfide (Cu2S) derived nanoparticles supported on carbon for electrochemical CO2 reduction. Copper sulfide consists of a family of chemical compounds with the formula Cu2Sx, where x represents Cu vacancies. [23] Cu2S can thus exist in a variety of different compositions and crystal structures, which have different chemical and physical properties and possibly also different catalytic performance. Therefore, two Cu2Sx catalysts with different starting compositions were studied, namely covellite (CuS) and chalcocite (Cu2S).

In thermal catalysts, techniques to prepare well-defined and uniformly distributed metal and metal oxide nanoparticles with tunable size have been well developed. [24] We explored a novel, two-step synthesis approach, building on known strategies for the preparation of supported CuO particles [25] followed by a liquid phase sulfidation. Carbon-supported CuS (CuS@C) or Cu2S (Cu2S@C) nanoparticles were synthesized by liquid phase sulfidation of carbon-supported CuO nanoparticles (CuO@C). This gave us a unique set of catalysts with very similar structural properties, that allowed us to investigate the influence of their chemical origin and of the presence of sulfur.

2. Experimental methods

2.1. Chemicals

XGNp500® graphene nanoplatelets (GNP) were purchased from XG Sciences. Copper(II) nitrate hydrate ([Cu(NO3)2]·2 H2O, 99 %), HNO3 (70 %), thiouacetamide (TAA, ≥99.0 %), Nafion® 117 solution, isopropanol (99.5 %), Potassium biphonate (KHCO3, ≥99 %) and 1-dodecanethiol (DDT, ≥98 %) were purchased from Sigma Aldrich. Nafion® Membrane XL was obtained from Ion Power, GmbH and carbon paper (TGP-H-060) was purchased from Toray.

2.2. Synthesis of carbon supported CuO nanoparticles

Carbon supported CuO (CuO@C) nanoparticles were prepared via incipient wetness impregnation followed by drying and heat treatment. [24–27] 1 g of graphite nanoplatelets (GNP500) was dried under vacuum at 120 °C for 2 h and subsequently impregnated with an aqueous Cu(NO3)2·3 H2O solution in 0.1 M HNO3. The volume used for impregnation corresponded to the total pore volume of the support, as determined with N2-physisorption. A weight loading of 20 wt% Cu was used for all samples. After impregnation, the sample was left to dry overnight at room temperature under vacuum. Then the sample was transferred into a tubular reactor and heat treated at 230 °C for 2 h under a N2 flow (200 mL/min) to decompose the Cu(NO3)2 into CuO nanoparticles.

2.3. Liquid-phase sulfidation of carbon-supported CuO nanoparticles

To obtain carbon supported CuS (CuS@C) nanoparticles, 400 mg of carbon supported CuO was dispersed into 40 mL deionized water in a 100 mL roundbottom flask and heated to 120 °C. At a temperature of 90 °C, 10 mL of an aqueous thioacetamide solution was gradually added to the mixture, using a Cu:S molar ratio of 1:1.1. The mixture was left to reflux for 2 h at 120 °C. After 2 h, the mixture was left to cool down and subsequently washed with demineralized water and acetone using vacuum filtration. The samples were then dried under vacuum for 2 h.

To obtain carbon supported Cu2S (Cu2S@C) nanoparticles, 400 mg of carbon supported CuO was dispersed in 15 mL 1-dodecanethiol in a 100 mL roundbottom flask and subsequently heated to 200 °C. The mixture was left to react for 2 h and then cooled down to room temperature. The sample was washed using toluene and acetone and subsequently dried under vacuum at room temperature.

2.4. Working electrode preparation

The working electrodes were prepared by spraying the CuO@C, CuS@C and Cu2S@C on a carbon paper substrate (Toray TGP-H-060). Prior to deposition of the catalyst, the carbon paper substrate was washed in ethanol by sonication for 30 min and subsequently rinsed with milliQ (MQ) water. Then, a catalyst ink was prepared by mixing 11 mg of carbon supported catalyst, 1120 μL isopropanol, 4470 μL MQ water and 44.4 μL Nafion solution. The ink was sonicated for 30 min in an ultrasonic bath to ensure good dispersion of the catalyst powder. Subsequently the ink was sprayed onto a round carbon paper electrode with a surface area of 4.9 cm2 using an Iwata HP-BP HI Performance Plus airbrush. A catalyst loading of 0.2 μg/cm2 (with 20 wt% Cu) was intended for all electrodes. The electrodes were dried overnight before electrochemical testing.

2.5. Characterisation of catalyst and electrodes

The GNP500 support was characterized using N2-physisorption performed with a Micromeritics TriStar instrument at a temperature of –196 °C. Prior to the measurements the powder was dried at 300 °C under N2-flow for 16 h. The pore diameter was determined using the Barrett-Joyner-Halenda (BJH) method and the pore volume was determined at p/p0 = 0.995.

Transmission electron microscopy (TEM) analysis was performed using a Thermo Fisher Scientific (formerly FEI) Tecnai12 microscope operating at 120 kV. The powder samples were suspended in ethanol and sonicated for 10 min. The suspension was then drop casted on a carbon coated 200 mesh copper TEM grid. Preparation of the TEM grids of the used catalyst was done by rubbing a TEM grid over the used electrode. Prior to this, the used electrodes were extensively rinsed with MQ water. TEM-EDX measurements were performed with a Thermo Fisher Scientific (formerly FEI) TalosF200X microscope, operating at 200 keV. For EDX measurements, gold TEM grids were used.

The electrodes before and after reaction were analyzed using scanning electron microscopy with energy dispersive X-ray spectroscopy (SEM-EDX) on a Thermo Fisher Scientific (formerly FEI) XL30SFESEM instrument.

X-ray diffraction (XRD) measurements were done on a Bruker D2 Phaser, equipped with a Cu Kα X-ray source with a wavelength of 1.79062 Å. The crystallite size of the copper sulfide nanoparticles was determined from the diffraction peak broadening using the Scherrer equation. [28]

2.6. Electrochemical measurements

A custom built, H-type electrochemical cell with two compartments separated by a Nafion membrane (Figure S1) was used for all electrochemical measurements. Each compartment of the electrochemical cell was filled with 11 mL electrolyte (0.5 M KNO3 or 0.5 M K2CO3 or 0.5 M KCl). The anolyte was purged with Argon and the catholyte was purged with CO2 at 10 mL/min. All electrochemical measurements were performed on an Autolab PGSTAT204 Potentiostat, with a Pt disk as counter electrode and a Ag/AgCl 3 M KCl reference electrode (Methrom). All
potentials were converted to the reversible hydrogen electrode (RHE) potential using the equation:

\[ E \text{(vs. RHE)} = E \text{(vs. Ag/AgCl)} + 0.209 + 0.059 \times \text{pH} \]

The catalyst supported on the carbon paper electrode was placed on a glassy carbon electrode for extra support and held in place by O-rings, leaving an electrode area of 3.8 cm² in contact with the electrolyte solution. Prior to all measurements, either CO₂ or Argon was bubbled through the solution for 20 min.

Cyclic voltammetry was performed in a 0.5 M KHCO₃ electrolyte, a scanning rate of 10 mV/s and a constant Ar or CO₂ flow of 10 mL/min. The selectivity of the catalysts was determined using chro- nopotentiometry at different current densities for 5 h (liquid products) or 1 h (gaseous products). Gaseous products were analyzed by connecting the outlet of the cathode compartment to a Global Analysis Solutions Microcompact GC 4.0. The GC system was equipped with three channels: The first channel has a Rt-QBond (10 m, 0.32 mm, Agilent) packed column and a FID detector for the detection of CH₄, C₂H₆ and CO₂, the second channel has Molecular Sieve 5A (10 m, 0.53 mm, Restek) packed column that separates small gaseous molecules such as CO and CH₄. This channel has a FID detector with a methanizer to increase the detection sensitivity of CO. The third channel has a Carbopack 1010 (8 m, 0.32 mm, Agilent) packed column which separates H₂ and CO₂ with a TCD. High purity nitrogen (99.999 %) was used as a carrier gas.

Liquid phase products were analyzed by analysis of the catholyte using a Varian HPLC equipped with a refractive index detector (RID) and a Bio-Rad Aminex HPX-87H column at 60 °C. 1 mM H₂SO₄ was used as the eluent with a flow rate of 0.55 mL/min. The retention time of formic acid was 15 min and the total analysis time was 20 min.

The Faradaic efficiency was calculated as:

\[ FE(\%) = \frac{n_e \times F \times [\text{moles of product}]_t}{Q} \times 100\% \]

In which \( n_e \) is the number of electrons needed to produce \( x \) (product) from CO₂ molecules and \( F \) is the Faradaic constant (96 485 s/A/mol).

For gaseous products (\( x = H_2, CO, CH_4, C_2H_6, \) or C₂H₄), the moles of product were determined via:

\[ \text{moles of product} = \frac{C_x \times q \times p}{RT} \]

in which \( C_x \) is the volumetric concentration of product \( x \) in ppm extracted from the GC calibration curve, \( q \) is the gas flow rate, \( p \) is the pressure, \( R \) is the ideal gas constant (8.314 m³·Pa·K⁻¹·mol⁻¹), \( T \) is the temperature, \( n_e \) is the number of electrons needed to produce \( x \) (product) from CO₂ molecules and \( F \) is the Faradaic constant (96 485 s/A/mol).

For liquid phase products (\( x = \text{formate} \)), the moles of product formed was determined via:

\[ \text{moles of product} = \frac{C_x \times V_{\text{catholyte}}}{1000 \times M_x} \]

in which \( C_x \) is the volumetric concentration of product \( x \) in ppm extracted from the HPLC calibration curve, \( V_{\text{catholyte}} \) is the volume of the catholyte (L) and \( M_x \) the molar weight of product \( x \) (g/mol).

2.7. XAS measurements

X-ray absorption measurements were performed at the Dutch-Belgian beamline DUBBLE, 26A at the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. Spectra were recorded at the Copper K-edge (8978.9 eV). Reference spectra of CuO@C, Cu₂S@C and Cu foil were recorded in transmission mode. The Athena XAS data processing software was used for the analysis of data.

For the in-situ XAS measurements, an electrochemical cell made by TU Delft was used (Figure S2). All in-situ experiments were performed in fluorescence mode, with an angle of 45° between the incoming X-rays and the sample. The time to acquire a spectrum was about 4.5 min and 2-5 scans were recorded at each potential. A CO₂-saturated 0.1 M KHCO₃ electrolyte was flown through the cell with a flowrate of 2 mL/min using a peristaltic pump. The Cu₂S@C electrodes were used as working electrode, a coiled platinum wire as counter electrode and a Ag/AgCl 3M KCl electrode as reference electrode. A BioLogic SP-240 potentiostat was used for all in-situ XAS experiments.

3. Results and discussion

3.1. Structural properties of the catalyst and electrodes

Supported CuS and Cu₂S catalysts were prepared via a novel, two-step synthesis route, in which carbon supported CuO nanoparticles (CuO@C) were converted into carbon supported CuS (CuS@C) or Cu₂S (Cu₂S@C) nanoparticles. This approach was chosen in order to investigate whether the nanoparticle size of the CuO@C sample could be maintained in the CuS@C and Cu₂S@C samples. Fig. 1 shows the X-ray diffractograms of CuO@C, CuS@C, Cu₂S@C and of bare carbon as a reference. For all four samples, diffraction lines are observed at 30.6° and 64.1°, which can be ascribed to the crystalline graphite support, which clearly was not damaged by the treatments. The CuO@C sample shows additional diffraction lines at 41.5°, 45.3° and 57.0°, which can be indexed with the monoclinic structure of CuO (PDF-01-007-6831 (ICDD, 2019) [30]). Upon sulfidation using thioacetamide, the diffraction pattern shows clear diffraction lines at 34.1°, 37.1°, 38.3° and 56.3°, which is consistent with the hexagonal CuS covellite structure (PDF-00-006-0464 (ICDD, 2019) [30]). The formation of metal sulfides using TAA has been described before and two different mechanisms are proposed [31-33]. CuS can be formed in a direct reaction, via the formation and subsequent decomposition of Cu-TAA complexes, or via a hydrolysis reaction, where the hydrolysis of TAA yields H₂S and H₂S subsequently reacts with Cu²⁺ (either dissolved or in CuO nanoparticles) to form CuS.

Upon sulfidation using 1-dodecanethiol, the reflections at 44.0°, 54.3° and 57.2° demonstrate the presence of hexagonal CuS chalcocite (PDF-00-053-0522 (ICDD, 2019) [30]). The formation of the supported Cu₂S nanoparticles from Cu²⁺ and DDT likely proceeded via the
formation of Cu-thiolate complexes, which were subsequently decomposed by thermal cleavage of the C–S bond, resulting in Cu–S monomers for nucleation and growth of Cu₂S nanoparticles.

For both the CuS@C and Cu₂S@C, no residual CuO reflections are observed, hence complete conversion of CuO to either CuS or Cu₂S was achieved. The relative broad peaks in the CuO@C, CuS@C and Cu₂S@C samples prove the presence of nanocrystallites and the absence of macrocrystalline material. The Scherrer equation was used to determine the crystallite size of the particles, resulting in crystallite sizes of 13, 14 and 12 nm for the CuO@C, CuS@C and Cu₂S@C samples, respectively.

Fig. 2 shows the transmission electron microscopy (TEM) images of the supported nanoparticles and the corresponding particle size histograms. Both the CuO@C and the Cu₂S@C samples show spherical particles with a particle size of 9 ± 3 nm for CuO and 17 ± 1 nm for Cu₂S. The supported CuS nanoparticles have a larger size of 25 ± 13 nm, with a few particles larger than 30 nm (Figure S3) and a platelet-like morphology (Fig. 2b). The CuO@C nanoparticles can thus be successfully converted to nanoparticles of either CuS@C or Cu₂S@C. However, especially for CuS@C significant particle growth is observed upon sulfidation. The increase in particle size upon sulfidation is likely caused by (partial) dissolution of the CuO@C nanoparticles caused by the high temperatures used during sulfidation (120 °C for CuS@C and 200 °C Cu₂S@C). The dissolved Cu²⁺ ions can react with either TAA or DDT and redeposit as CuS or Cu₂S, respectively, via the mechanisms described above.

The working electrodes CuO@C, CuS@C or Cu₂S@C were prepared by spraying an ink containing the catalyst powder onto carbon paper substrates. Fig. 3 shows representative scanning electron microscopy (SEM) images of the bare carbon paper substrate and of the CuS@C nanoparticles sprayed onto the carbon paper substrate, illustrating the distribution over and adherence of the CuS@C to the carbon paper substrate. CuS@C was also well distributed over the carbon paper (Figure S4).

3.2. Electrochemical characterization of the catalyst under eCO₂R reaction conditions

Fig. 4 shows the cyclic voltammograms of CuS@C and Cu₂S@C in a 0.5 M KHCO₃ electrolyte (pH = 7.1) under argon flow, obtained with a scan rate of 10 mV/sec. For each sample, the first 5 cycles are shown. For CuS@C (Fig. 4a), two reduction peaks are observed in the first cathodic scan at potentials of around -0.1 V and -0.5 V, with onsets around -0.0 V and -0.4 V vs. RHE, respectively. Figure S5 shows the Pourbaix diagram of an aqueous Cu-S system at 25 °C with Cu and S concentrations of 1 M. The diagram shows that under these conditions, CuS is not thermodynamically stable at potentials more negative than -0.4 V vs. SHE (0.0 V vs. RHE) in aqueous solutions at pH 7, and can be reduced, via several copper rich Cu₂S phases, to metallic Cu. Relating this to the reduction peaks observed in the voltammogram of CuS@C, the first reduction peak can be attributed to a reduction of the CuS nanoparticles to more copper-rich copper sulfide phases, such as djurleite(Cu₁.₉₄S) or chalcocite (Cu₂S). The second reduction peak can then be attributed to the reduction of the remaining Cu₂S phase to metallic Cu.

The Cu₂S@C sample (Fig. 4b) also shows a reduction peak around -0.5 V vs. RHE, with an onset around -0.4 V vs. RHE, which corresponds to a reduction of the Cu₂S nanoparticles to metallic Cu. In addition, a reduction peak is observed around a potential of -0.4 V, with an onset at -0.2 V. This reduction likely originates from a reduction of a more copper deficient Cu₂S phase or some CuO, which was formed during the storage of the electrode in air prior to the measurements. Figure S6 shows the voltammogram of the CuO@C sample, showing two reduction peaks with onsets above 0 V vs. RHE and around -0.2 V vs. RHE in the first cathodic scan. These reduction peaks correspond to the ready reduction of CuO to Cu₂O and subsequently metallic Cu. [34] The CVs thus show that CuS@C and Cu₂S@C are reduced to metallic...
Cu when scanning to potentials more negative than -0.4 V vs. RHE. However, no oxidation peaks are observed in the anodic cycles of the CVs (Fig. 4 a,b), indicating the reduced phases do not oxidize back to the original CuS or Cu$_2$S phase in this potential range. During reduction, the sulfur will likely react with protons from solution to form SH$^{(aq)}$ and H$_2$S$^{(g)}$. [15] The gaseous H$_2$S will escape from the cell, explaining the irreversibility of the reduction of CuS and Cu$_2$S.

X-ray absorption spectroscopy (XAS) was employed to probe the copper phases under operando reaction conditions. In-situ techniques are invaluable as the characterization of catalysts after reaction is challenging due to the limited amount of catalyst used in reactions. In addition, the catalyst is often highly susceptible to changes when a potential is no longer applied, and the catalyst is removed from the electrolyte solution (e.g. oxidation). XAS is a very suitable technique for in-situ studies of electrocatalysts, as X-rays can easily penetrate through air and water due to their high energy. [35–37]

Fig. 5 shows the ex-situ XAS spectra of the CuS@C and Cu$_2$S@C samples (dotted lines) and the in-situ XAS spectra of these samples (solid lines). The conditions under which the samples were measured are different. The ex-situ spectra were recorded on pellets of the as-synthesized powder samples. In contrast, the in-situ spectra shown in Fig. 5a are of the CuS@C and Cu$_2$S@C powders, deposited on the carbon paper substrate, in contact with a CO$_2$ saturated 0.1 M KHCO$_3$ electrolyte (pH = 6.8), without applying a potential. However, for both CuS@C and Cu$_2$S@C, the in-situ spectrum shows the same features as the corresponding sample measured under ex-situ conditions. This demonstrates that the CuS@C and Cu$_2$S@C were stable upon deposition of the catalyst powder on the carbon paper substrate and subsequent contact with the CO$_2$ saturated electrolyte, and hence validates the in-situ measurements.

Subsequently, the CuS@C and Cu$_2$S@C samples were analyzed under CO$_2$ and H$^+$ reduction conditions. Fig. 6a shows the in-situ XAS spectra of the CuS@C sample upon applying negative potentials of -0.9 V, -1.2 V and 1.6 V vs. RHE. Additional spectra of the sample at potentials of -0.05 V, -0.25 V and -0.6 V vs. RHE are shown in Figure S7. Up to a potential of -0.6 V, no clear changes are observed in the XAS spectra when compared to the spectrum of the ex-situ CuS@C sample (Figure S7). Upon applying potentials more negative than -0.9 V vs. RHE, the spectra show clear changes, features start to appear at energies of 9025 eV and 9070 eV. When going to more negative potentials, such as -1.2 V and -1.6 V vs. RHE, these features become even more pronounced. As is shown by the ex-situ XAS pattern of the Cu foil, the peaks at 9025 eV and 9070 eV are characteristic for metallic Cu. Although it is difficult to distinguish if the reduction occurs via any intermediate copper-rich Cu$_{2-x}$S phases, the in-situ XAS data thus clearly show the reduction of the CuS nanoparticles to metallic Cu under reaction conditions.

This technique thus allowed us to elucidate the chemical phase of the catalyst under electrochemical CO$_2$ reduction conditions. For both samples, the potential at which a reduction to Cu is observed by XAS (-0.9 V) is more negative than the potential of -0.5 V vs. RHE at which the reduction peak was observed in the CVs (Fig. 4 b,c). This is likely caused by the different experimental conditions used in the two experiments, such as the different cell geometries (see Figure S1, S2) leading to differences in ohmic resistance. In addition, an electrolyte...
concentration of 0.1 M KHCO$_3$ was used during the in-situ XAS experiments, as compared to a concentration of 0.5 M KHCO$_3$ and the resulting lower conductivity and pH is expected to shift the reduction to slightly more negative potentials. Therefore additional CV measurements were performed in the XAS cell. The reduction peaks of the reduction of the CuS@C and Cu$_2$S@C sample to metallic Cu are clearly visible at -0.63 V vs. RHE for both samples (supporting information, Figure S8). When comparing these values to the position of the reduction peaks in the H-type cell (supporting information, Table S1), we see that the reduction peak of the reduction of Cu$_2$S to metallic Cu is indeed shifted to more negative potentials when using the XAS cell.

Most importantly, XAS is a bulk technique and from the fact that the spectra at potentials more negative than -0.9 V correspond to metallic Cu, with very little or none of the original ex-situ spectrum remaining, it can be concluded that the vast majority of the nanoparticles is electrochemically active, and in good electrochemical contact with the carbon support and carbon-paper substrate as well as with the electrolyte solution. This is the first time that the chemical phases of Cu$_2$S@C- and Cu$_2$S@C-oxidized catalysts were followed in-situ under CO$_2$ and H$^+$ reduction conditions, giving valuable insight in the active catalyst phase and on the electrochemical contact between the nanoparticles and support material.

**3.3. Electrochemical CO$_2$ reduction**

The product selectivity of the CuS@C and Cu$_2$S@C derived catalysts in electrochemical CO$_2$ reduction were compared to that of the CuO@C derived catalyst. For all three catalysts, the main CO$_2$ reduction product formed was formate. Table 1 shows the amount of formate produced for all three catalysts and a carbon reference and two sulfidized carbon references, treated with either TAA or DDT, at different currents applied. The data is obtained by CP for 5 h with currents of -1.5, -3 and -4.5 mA/cm$^2$ performed in a 0.5 M KCl +0.5 M KHCO$_3$ electrolyte saturated with CO$_2$ (pH 7.5). The average potentials over 5 h of CP are also shown (with standard deviation).

| Applied current density (mA/cm$^2$) | Sample | Formate Production (μmol/hr) | Faradaic Efficiency Formate (%) | Resulting Potential (V vs. RHE) |
|------------------------------------|--------|-------------------------------|---------------------------------|-------------------------------|
| -1.5                               | CuS@C | 9                            | 8                               | -0.77 ± 0.02                 |
|                                    | Cu$_2$S@C | 13                        | 12                             | -0.78 ± 0.02                 |
|                                    | CuO@C | 3                            | 3                              | -0.66 ± 0.04                 |
|                                    | C     | 3                            | 3                              | -0.82 ± 0.02                 |
|                                    | C-TAA | 1                            | 1                              | -0.93 ± 0.01                 |
|                                    | C-DDT | 4                            | 4                              | -0.96 ± 0.05                 |
|                                    | Cu$_2$S@C | 14                       | 7                              | -0.90 ± 0.08                 |
|                                    | Cu$_2$S@C | 21                       | 10                             | -0.76 ± 0.01                 |
|                                    | CuO@C | 16                           | 8                              | -0.88 ± 0.10                 |
|                                    | C     | 4                            | 2                              | -0.96 ± 0.01                 |
|                                    | C-TAA | 3                            | 1                              | -1.13 ± 0.06                 |
|                                    | C-DDT | 4                            | 2                              | -1.00 ± 0.02                 |
|                                    | CuS@C | 11                           | 3                              | -1.05 ± 0.05                 |
|                                    | Cu$_2$S@C | 12                       | 4                              | -0.87 ± 0.03                 |
|                                    | CuO@C | 19                           | 6                              | -0.91 ± 0.02                 |
|                                    | C     | 6                            | 2                              | -1.65 ± 0.03                 |
|                                    | C-TAA | 4                            | 1                              | -1.20 ± 0.07                 |
|                                    | C-DDT | 4                            | 1                              | -1.06 ± 0.03                 |

The amount of formate produced and Faradaic efficiency of formate during CO$_2$ reduction using CuO@C, CuS@C and Cu$_2$S@C, a carbon reference and two sulfidized carbon references, treated with either TAA or DDT, at different currents applied. The data is obtained by CP for 5 h with currents of -1.5, -3 and -4.5 mA/cm$^2$, performed in a 0.5 M KCl +0.5 M KHCO$_3$ electrolyte saturated with CO$_2$ (pH 7.5). The average potentials over 5 h of CP are also shown (with standard deviation).

When a current density of -1.5 mA/cm$^2$ was applied, formate was the only reduction product, and the resulting concentration of 0.1 M KHCO$_3$ was used during the in-situ XAS experiments, as compared to a concentration of 0.5 M KHCO$_3$ and the resulting lower conductivity and pH is expected to shift the reduction to slightly more negative potentials. Therefore additional CV measurements were performed in the XAS cell. The reduction peaks of the reduction of the CuS@C and Cu$_2$S@C sample to metallic Cu are clearly visible at -0.63 V vs. RHE for both samples (supporting information, Figure S8). When comparing these values to the position of the reduction peaks in the H-type cell (supporting information, Table S1), we see that the reduction peak of the reduction of Cu$_2$S to metallic Cu is indeed shifted to more negative potentials when using the XAS cell.

When a current density of -1.5 mA/cm$^2$ was applied, formate was the only reduction product, and the resulting concentration of 0.1 M KHCO$_3$ was used during the in-situ XAS experiments, as compared to a concentration of 0.5 M KHCO$_3$ and the resulting lower conductivity and pH is expected to shift the reduction to slightly more negative potentials. Therefore additional CV measurements were performed in the XAS cell. The reduction peaks of the reduction of the CuS@C and Cu$_2$S@C sample to metallic Cu are clearly visible at -0.63 V vs. RHE for both samples (supporting information, Figure S8). When comparing these values to the position of the reduction peaks in the H-type cell (supporting information, Table S1), we see that the reduction peak of the reduction of Cu$_2$S to metallic Cu is indeed shifted to more negative potentials when using the XAS cell.

Most importantly, XAS is a bulk technique and from the fact that the spectra at potentials more negative than -0.9 V correspond to metallic Cu, with very little or none of the original ex-situ spectrum remaining, it can be concluded that the vast majority of the nanoparticles is electrochemically active, and in good electrochemical contact with the carbon support and carbon-paper substrate as well as with the electrolyte solution. This is the first time that the chemical phases of Cu$_2$S@C- and Cu$_2$S@C-oxidized catalysts were followed in-situ under CO$_2$ and H$^+$ reduction conditions, giving valuable insight in the active catalyst phase and on the electrochemical contact between the nanoparticles and support material.

**3.3. Electrochemical CO$_2$ reduction**

The product selectivity of the CuS@C and Cu$_2$S@C derived catalysts in electrochemical CO$_2$ reduction were compared to that of the CuO@C derived catalyst. For all three catalysts, the main CO$_2$ reduction product formed was formate. Table 1 shows the amount of formate produced for all three catalysts and a carbon reference and two sulfidized carbon reference, treated with either TAA or DDT, obtained from chronopotentiometry (CP) for 5 h in a 0.5 M KCl + 0.5 M KHCO$_3$ electrolyte saturated with CO$_2$ at current densities of -1.5, -3.0 and -4.5 mA/cm$^2$. Corresponding bar charts are shown in Figure S9. Here, the KCl was added as supporting electrolyte to increase the conductivity of the electrolyte and minimize ohmic resistances. Figure S10 shows the potentials over time obtained during the 5 h CP experiments.

When a current density of -1.5 mA/cm$^2$ was applied, formate was produced with yields of 9, 13 and 3 μmol/hr for the CuS@C-, Cu$_2$S@C- and CuO@C-derived catalysts, respectively. As the CuO@C-derived catalyst showed a similar production of formate (3 μmol/hr) to the bare carbon support and the two sulfidized carbon supports (3 μmol/hr, 1 μmol/hr and 5 μmol/hr for C, C-TAA and C-DDT, respectively), the production of formate on this catalyst is considerably less. The production of formate on the CuS@C- and Cu$_2$S@C-derived catalysts is however significantly higher, showing that the production of formate is clearly promoted on the sulfide derived catalysts when compared to the oxide derived catalyst and the carbon support. In addition, more formate is produced by the Cu$_2$S@C-derived catalyst than by the CuS@C-derived catalyst. This indicates that, even though both CuS@C and Cu$_2$S@C reduce to metallic Cu under reaction conditions, there is a difference between the two Cu-sulfide derived catalysts depending on whether they originate from CuS@C or Cu$_2$S@C. When applying more negative potentials, the formate production also on the oxide-derived catalyst
increases, making the differences less pronounced, although in all cases the CuS@C-derived catalyst produces more formate than the CuO@C-derived catalyst.

Besides formate, only small amounts of CO were produced with Faradaic efficiencies between 0–1.3 % (Table S2, Figure S11). The differences between the measured CO concentrations on the different catalysts were close to the experimental error. No other CO₂ reduction products were found in the liquid or gaseous phase and hence the rest of the current led to the production of H₂. The production of H₂ readily occurred on the bare carbon reference sample. In fact, the loading with Cu-based nanoparticles corresponds to a surface coverage of less than 4% of the carbon support (see Support Information for calculations). It is hence amazing that CO₂ reduction products with Faradaic efficiencies ranging from 3 to 12% can be obtained with such a low coverage of the carbon surface, which so readily produced H₂ instead. This means that the Cu-based nanoparticles clearly introduce a strong preference for CO₂ rather than H⁻ reduction.

The enhanced selectivity for formate on the sulfide derived catalysts at current densities of -1.5 mA/cm² is consistent with previous studies and can be explained by the different pathways possible for CO₂ reduction. [14,15,17,18] Norskov et al. proposed that the reduction of CO₂ to HCOO⁻ occurs via a *OCHO intermediate, which is further reduced to formate (HCOO⁻). [38] On the other hand, other studies suggest the formation of formate occurs via a direct reaction of physisorbed CO₂ with adsorbed H⁺. [18,39] In both mechanisms, CO and further reduced products are formed via a so-called CO pathway, in which CO₂ is reduced to *COOH, which can be further reduced to *CO, which in turn can desorb or be further reduced to various hydrocarbons and alcohols. Although no distinction can be made between the two mechanisms to form formate, the increased selectivity for formate observed for the sulfide derived catalysts at current densities of -1.5 mA/cm² suggests that the HCOO⁻ pathway is favored over the CO pathway. Which pathway is preferred highly depends on the binding strength of important intermediates such as CO on the catalyst, suggesting that the binding affinity for the CO₂ reduction intermediates is different on the sulfide-derived catalysts than on the oxide-derived catalyst. Nevertheless, elucidation of the exact role of sulfur on the CO₂ reduction mechanism will require additional studies.

### 3.4. Structure of catalyst after reaction

The increased formation of formate on CuS@C-derived catalyst when compared to CuS@C-derived catalyst suggests a difference in the chemical composition between the two. Hence, additional characterization of the catalysts after reaction was performed. After being used in the CO₂ and H⁻ reduction for 5 h at -3 mA/cm² in 0.5 M KCl + 0.5 M KHCO₃, the CuO@C-, CuS@C- and CuS@C-derived catalysts were analyzed to see if any changes occurred upon reaction. Fig. 7 shows TEM images of the CuO@C-, CuS@C- and CuS@C-derived catalysts after being employed in the CO₂ reduction reaction. For CuO@C, the particle size increased from 9 ± 3 nm to 33 ± 14 nm. For the CuS@C-derived catalyst, no significant change in particle size is observed, whereas for the CuS@C-derived catalyst, the particle size decreased from 17 ± 1 nm to 6 ± 2 nm upon reaction. Upon reaction, the CuO@C- and CuS@C-derived catalysts thus have a fairly similar particle size distribution, while the CuS@C-derived catalyst has smaller and more monodisperse particles. This difference in particle size could play a role in the different catalytic behavior discussed in section 3.3... [40,41] The larger size of the CuS@C-derived catalyst when compared to the CuS@C-derived catalyst can possibly explain the lower selectivity to formate for the CuS@C-derived catalyst, as the larger CuS-derived particles have a lower electrochemically active surface area.

In addition, TEM-EDX measurements were performed to analyze the presence of sulfur in the catalysts after reaction. The measurements showed remaining sulfur in the samples after reaction, where copper to sulfur ratios of 3:1 and 6:1 were found for the CuS@C- and CuS@C-derived catalyst, respectively (Table S3). This was much higher than the sulfur content detected in the CuO@C sample, with a Cu:S ratio of 32:1. This sample was used as a reference for the base sulfur content, as the Nafion binder used in the electrode preparation contains some sulfur as well. A small amount of sulfur, as well as K and Cl, (Figure S13) can also be ascribed to incomplete removal of the electrolyte. Although precise quantification is thus not straightforward, it is clear that sulfur content is higher for the CuS@C derived catalyst than for CuS@C-derived catalyst, which can be explained by the higher initial sulfur content of CuS@C, and for both sulfide derived catalysts much higher than for the oxide derived catalyst. Also the difference in sulfur content likely influences the product selectivities observed for the two sulfide-derived catalysts.

![Fig. 7. TEM figures of the a) CuO@C-, b) CuS@C-, and c) CuS@C-derived catalysts after 5 h of CO₂ and H⁻ reduction at -3 mA/cm² in 0.5 M KCl + 0.5 M KHCO₃. The lower panels show corresponding particle size histograms. For CuO@C, a few particles larger than 50 nm were also observed (Figure S12).](image-url)
4. Conclusions

In summary, carbon-supported, Cu$_2$S$_x$ derived nanoparticles were studied for electrochemical CO$_2$ reduction in aqueous media. First, CuS@C and CuS$_x$@C nanoparticles were successfully prepared via a liquid phase sulfidation of CuO@C nanoparticles. Subsequently the catalysts were deposited on carbon paper electrode substrates. Upon electrochemical CO$_2$ reduction, CuS@C and CuS$_x$@C nanoparticles undergo a reduction to metallic Cu, as was verified by both CV and in-situ XAS experiments. EDX measurements showed that some residual sulfur is left in both catalysts. The selectivity of the CuS@C- and CuS$_x$@C-derived catalysts was tested and compared to that of a CuO@C-derived catalyst. On all three catalysts, formate was produced as the main CO$_2$ reduction product, next to H$_2$. Electrochemical CO$_2$ liquid phase sulfidation of CuO@C nanoparticles. Subsequently theCuS@C and Cu$_2$S@C studiered for electrochemical CO$_2$ reduction, CuS@C and CuS$_x$@C nanoparticles. Subsequently the CuS@C and Cu$_2$S@C derived catalysts were tested and compared to that of a CuO@C-derived catalyst. On all three catalysts, formate was produced as the main CO$_2$ reduction product, next to H$_2$. Electrochemical CO$_2$ liquid phase sulfidation of CuO@C nanoparticles. Subsequently the CuS@C and Cu$_2$S@C derived catalysts were tested and compared to that of a CuO@C-derived catalyst. On all three catalysts, formate was produced as the main CO$_2$ reduction product, next to H$_2$. Electrochemical CO$_2$ liquid phase sulfidation of CuO@C nanoparticles. Subsequently the CuS@C and Cu$_2$S@C derived catalysts were tested and compared to that of a CuO@C-derived catalyst. On all three catalysts, formate was produced as the main CO$_2$ reduction product, next to H$_2$. Electrochemical CO$_2$ liquid phase sulfidation of CuO@C nanoparticles. Subsequently the CuS@C and Cu$_2$S@C derived catalysts were tested and compared to that of a CuO@C-derived catalyst. On all three catalysts, formate was produced as the main CO$_2$ reduction product, next to H$_2$. Electrochemical CO$_2$ liquid phase sulfidation of CuO@C nanoparticles. Subsequently the CuS@C and Cu$_2$S@C derived catalysts were tested and compared to that of a CuO@C-derived catalyst. On all three catalysts, formate was produced as the main CO$_2$ reduction product, next to H$_2$. Electrochemical CO$_2$ liquid phase sulfidation of CuO@C nanoparticles.
[24] P. Munnik, P.E. De Jongh, K.P. De Jong, Recent developments in the synthesis of supported catalysts, Chem. Rev. 115 (2015) 6687–6718, https://doi.org/10.1021/cr500486u.

[25] R. Van Den Berg, G. Prieto, G. Korpershoek, L.I. Van Der Wal, A.J. Van Bunningen, S. Legsgaard-Jørgensen, P.E. De Jongh, K.P. De Jong, Structure sensitivity of Cu and CuZn catalysts relevant to industrial methanol synthesis, Nat. Commun. 7 (2016), https://doi.org/10.1038/ncomms13057.

[26] P. Munnik, M. Wolters, A. Gabrielson, S.D. Pollington, G. Headdock, J.H. Bitter, P.E. De Jongh, K.P. De Jong, Copper nitrate redispersion to arrive at highly active silica-supported copper catalysts, J. Phys. Chem. C 115 (2011) 14698–14706, https://doi.org/10.1021/jp111778g.

[27] G. Wang, R. van den Berg, C. de Mello Donega, K.P. de Jong, P.E. de Jongh, Silica-supported Cu2O nanoparticles with tunable size for sustainable hydrogen generation, Appl. Catal. B Environ. 192 (2016) 199–207, https://doi.org/10.1016/J.APCATB.2016.03.044.

[28] A.L. Patterson, The scherrer formula for X-ray particle size determination, Phys. Rev. 56 (1939) 978–982, https://doi.org/10.1103/PhysRev.56.978.

[29] J. Li, F. Che, Y. Pang, C. Zou, J.Y. Howe, T. Burdyny, J.P. Edwards, Y. Wang, Z. Wang, P. De Lana, C.T. Dinh, T.T. Zhuang, M.I. Saidaminov, S. Cheng, T. Wu, Y. Z. Finfrock, L. Ma, S.H. Hsieh, Y.S. Liu, G.A. Botton, W.F. Pong, X. Du, J. Guo, T. K. Sham, E.H. Sargent, D. Sinton, Copper adparticle enabled selective electrocatalytic carbon dioxide reduction through operando techniques, Nat. Catal. 1 (2018) 922–934, https://doi.org/10.1038/s41929-018-0182-6.

[30] T. Cheng, H. Xiao, W.A.G. Iii, Reaction mechanisms for the electrochemical reduction of CO2 to CO and formate on the Cu(100) surface at 298 K from quantum mechanics free energy calculations with explicit water, J. Am. Chem. Soc. 138 (2016) 25, https://doi.org/10.1021/jacs.6b08534.

[31] A. Loiadice, P. Lobaccaro, E.A. Kamali, T. Thao, B.H. Huang, J.W. Ager, R. Buonsanti, Tailoring copper nanocrystals towards C2 products in electrochemical CO2 reduction, Angew. Chemie - Int. Ed. 55 (2016) 5789–5792, https://doi.org/10.1002/anie.201601582.

[32] R. Reske, H. Mistry, F. Behafarid, B. Roldan Cuenya, P. Strasser, Particle size effects in the catalytic electroreduction of CO2 on Cu nanoparticles, J. Am. Chem. Soc. 136 (2014) 6978–6986, https://doi.org/10.1021/ja500328L.