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| Original Influence of sonication on co-precipitation synthesis of copper oxide catalyst for CO2 electroreduction / Roldán, Daniela; Guzmán, Hilmar; Russo, Nunzio; Hernández, Simelys ELETTRONICO (2021), pp. 1-1. ((Intervento presentato al convegno 2021 VIRTUAL MRS SPRING MEETING & EXHIBIT tenutosi a Online nel From 17th to 23th April. |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Availability: This version is available at: 11583/2898460 since: 2021-05-06T15:27:35Z |
| Publisher: Materials Research Society |
| Published DOI: |
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16 July 2022









Influence of sonication on coprecipitation synthesis of copper oxide catalyst for CO₂ electroreduction

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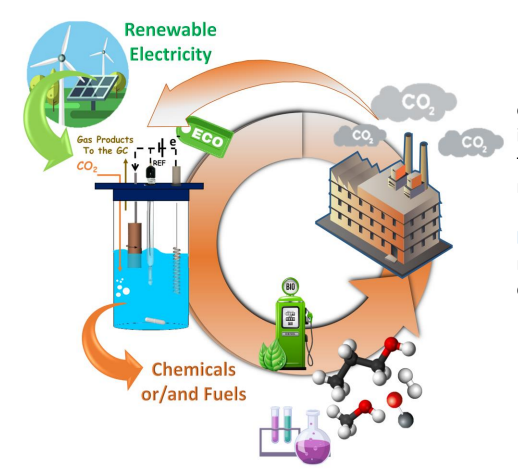
Overview



The need to reduce greenhouse gas emissions and increase our energy supply makes the **electrochemical reduction of** CO_2 (CO_2R) a very attractive alternative to produce non-fossil-based fuels or chemicals. Copper-based catalysts is one of the catalyst that most efficiently promote the formation of species with one or more carbon-carbon bonds from the electrochemical reduction of CO_3 .

It was decided to evaluate the effect of the ultrasound application (US) on the shape and size of the particles obtained, its electrocatalytic activity and its selectivity to products of interest.





Physical characterization was carried out by using different techniques including X-ray diffraction, BET and filed-emission scanning electron microscopy (FESEM).

Electrochemical tests for CO₂ reduction were done under ambient conditions.



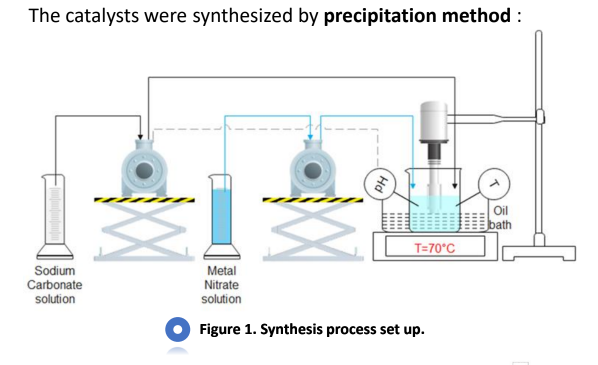
Materials and Methods

The catalysts were synthesized by precipitation method with different ultrasound conditions to evaluate its effect on the performance of the catalysts. All catalysts are copper-based and with the **same precursor concentration**. **Different ultrasound amplitude** values were applied:

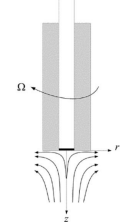
Table 1. Conditions of the different catalysts.

| Catalyat | Precursor concer | ntration, | | |
|-------------|---------------------------------------|----------------------------------|------------|--|
| Catalyst | M | Ultrasound conditions | %Amplitude | |
| name | Cu(NO ₃) ₂ .3I | H_2O | | |
| Cu-06 | 0.6 | Without ultrasound | - | |
| Cu-06-%23-A | 0.6 | Ultrasound in the ageing process | 23 | |
| Cu-06-%30-A | 0.6 | Ultrasound in the ageing process | 30 | |
| Cu-06-%37-A | 0.6 | Ultrasound in the ageing process | 37 | |

4 catalysts with different US conditions



The tests were carried out in a threeelectrode electrochemical cell using a rotating disk electrode (RDE) system.



Results

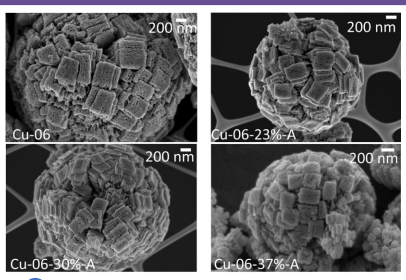
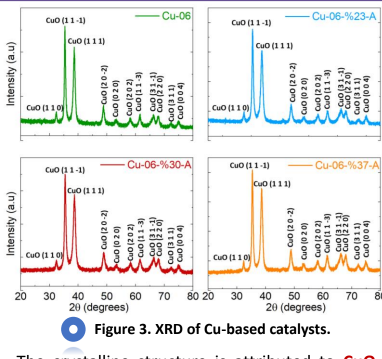
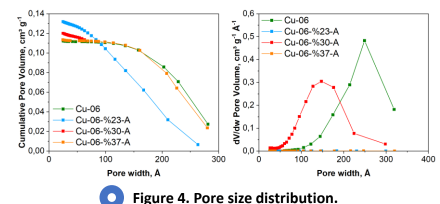


Figure 2. FESEM images of Cu-based catalysts.

The US during the ageing process do not influence the final morphology of the catalyst at the different applied amplitudes. The highest US amplitude produced less spherical particles.



The crystalline structure is attributed to **CuO** and does not change with the US irradiation.



Pore size distribution is narrower by assisting the co-precipitation synthesis with US

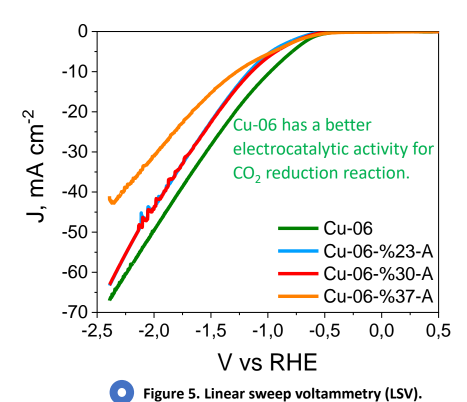
Table 2. Physical-chemical properties.

| Catalyst | BET surface area, | Total pore | Crystallite size, nm |
|-------------|--------------------------------|-----------------------------------------|-----------------------|
| Cuturyst | m ² g ⁻¹ | volume, cm ³ g ⁻¹ | (1 1 -1) facet of CuO |
| Cu-06 | 18.40 | 0.107 | 13.97 |
| Cu-06-%23-A | (34.07) | 0.135 | 11.38 |
| Cu-06-%30-A | 32.07 | 0.125 | 11.39 |
| Cu-06-%37-A | 18.68 | 0.124 | 16.78 |

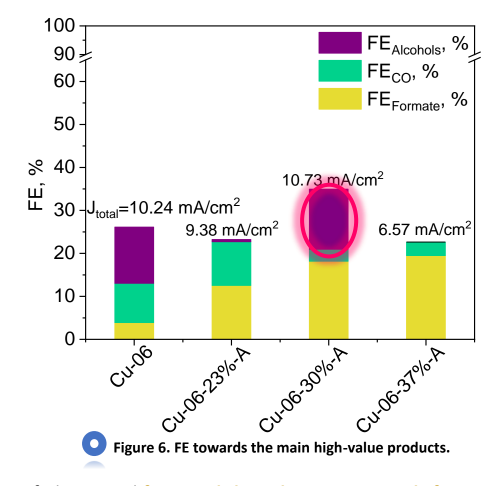
The US has increased the surface area of the sonicated sample at %23 and 30% in comparison to the non-sonicated one. Probably the smaller BET surface area of the %37 sample is due to the higher crystallites size. On the other hand, there is an increase of pores volume by using US compared to the original sample, Cu-06.

Results

Cu-06-23%-A and Cu-06-30%-A exhibit the performance, indicating that the materials similar prepared have electrocatalytic activity for the CO₂ reduction reaction.



The most promising catalyst among the new US-prepared catalysts is the Cu-06-%30-A with 14% of FE to alcohols.



The use of ultrasound favoured the selectivity towards formate. 5

Conclusion

Cu-based catalysts with different physicochemical properties were prepared by ultrasound assisted coprecipitation method.

Regarding the physical characteristics, we found that **pore size distribution is narrower by increasing the US amplitude**. On the other hand, there is no significant difference in morphology and dimension of particles.

The surface area increased with the use of ultrasound (23 and 30%), which is attributed to an improved dispersion created by acoustic cavitation. However, there is an optimal amplitude (30%) over which the advantages of the use of ultrasound are diminished.

Ultrasound has also an effect on Copper-based catalysts performance; in this case, the selectivity towards alcohols and C₁ products (Formate) was enhanced.

This work has been performed with the financial support of Eni SpA and the R&D Program Energy Transition (Cattura e Utilizzo CO₂).