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3901

From molecular copper complexes to composite electrocatalytic materials for selective reduction of CO₂ to formic acid†

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The development of new energy storage technologies is central to solving the challenges facing the widespread use of renewable energies. An option is the reduction of carbon dioxide (CO₂) into carbon-based products which can be achieved within an electrochemical cell. Future developments of such processes depend on the availability of cheap and selective catalysts at the electrode. Here we show that a unique well-characterized active electrode material can be simply prepared *via* electrodeposition from a molecular copper complex precursor. The best performances, namely activity (150 mV onset overpotential and 1 mA cm⁻² current density at 540 mV overpotential), selectivity (90% faradaic yield) and stability for electrocatalytic reduction of CO₂ into formic acid in DMF/H₂O (97 : 3 v/v) have been obtained with the [Cu(cyclam)](ClO₄)₂ complex (cyclam = 1,4,8,11-tetraazacyclotetradecane) as the precursor. Remarkably the organic ligand of the Cu precursor remains part of the composite material and the electrocatalytic activity is greatly dependent on the nature of that organic component.

Received 19th December 2014
Accepted 5th January 2015

DOI: 10.1039/c4ta07022d

www.rsc.org/MaterialsA

CO₂ transformation into energy-dense organic compounds is currently emerging as an exciting strategy in the context of the development of new energy technologies.^{1–9} It indeed (i) allows storage of intermittent and diluted renewable energy sources (wind and sun) in the form of concentrated chemical energy (chemical fuels and industrial chemicals conventionally derived from petroleum), (ii) may contribute to limit the accumulation of CO₂, a gas with greenhouse effect, in the atmosphere, and (iii) generates a global carbon-neutral and circular energy system. Practically this can be achieved through electro-reduction of CO₂ within electrochemical cells ideally powered by wind turbines or photovoltaic panels.¹⁰ Unfortunately the reactions

involve multiple electrons. Therefore they are generally limited by very low kinetics which translate into large overpotentials (thus with significant energy losses) as well as by low selectivity since a variety of CO₂-derived products (CO, HCOOH, oxalate and, to lower extents, methanol or hydrocarbons) together with molecular hydrogen can be produced at the same time.⁶ The latter derives from the parallel reduction of protons required for CO₂ activation.

A key technological challenge thus resides in the development of active electro-catalysts, for reducing CO₂ with significant current densities at low overpotentials, in reactions displaying high selectivity and faradaic yield for a given CO₂-derived product. In particular, the production of liquid fuels such as formic acid¹¹ or methanol,¹² fitting into existing infrastructure for fuel transportation and distribution, is an attractive target. A number of electrocatalytic materials have been assayed under alkaline aqueous carbonate solutions.⁶ However, recent studies^{13–17} were carried out in non-aqueous media such as organic solvents and ionic liquids, since they offer larger CO₂ solubility and an easy control of H₂ evolution through tuning of water concentration. This allowed the development of novel materials for example based on MoO₂,¹⁶ Bi^{13,14} or B-doped diamond¹⁵ reducing CO₂ at low overpotential with good selectivity. Molecular metal complexes^{1,2,5,6} constitute another interesting class of such selective electrocatalysts¹⁸ operating at low overpotential in organic solvents. Their activity can be synthetically tuned through ligand variations. To date, only one copper

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/c4ta07022d

coordination compound has been reported as an electrocatalyst for CO₂ reduction.¹⁹ In the course of a study aimed at identifying novel Cu-based molecular CO₂-reducing catalysts, we found that a composite material could be electrodeposited on the electrode. This Cu-based material, which has been extensively characterized and shown to retain the organic ligand from the copper precursor, displays remarkable activity, selectivity and stability for electrocatalytic reduction of CO₂ into formic acid, a product of wide industrial applications.

Results and discussion

Continuous voltamperometric cycling at a fluorine-doped tin oxide (FTO) electrode dipped into a CO₂-saturated solution of 1.3 mM [Cu(cyclam)](ClO₄)₂ (cyclam = 1,4,8,11-tetraazacyclotetradecane) in DMF containing 3% water and supporting electrolyte (0.1 M *n*-Bu₄NBF₄) triggered the appearance and growth of a chemically-reversible signal with both an oxidation wave and a reduction wave, at $E_{pa} = 0.58$ V vs. Fc^{+/0} and $E_{pc} = -0.98$ V vs. Fc^{+/0}, (black trace in Fig. 1) respectively, and tentatively assigned to a Cu-based process.^{20,21} This feature revealed an electrodeposition reaction in agreement with the observation of a blue material at the surface of the electrode (inset in Fig. 1, in the following “blue material” will be used to name this novel Cu-based electrodeposit). In addition, the cyclic voltammograms display an intense current wave, with an onset at -1.60 V vs. Fc^{+/0}, reflecting an electrocatalytic process. Importantly, scanning over this electrocatalytic process was a requirement for the deposition of the blue material. No electrodeposition and no electrocatalytic wave could be observed when CO₂ or water was removed from the reaction mixture, thus strongly suggesting electrocatalytic CO₂ reduction. The cyclic voltammogram (red trace in Fig. 1) of the modified electrode was then recorded independently under similar conditions but in the

absence of [Cu(cyclam)]²⁺. It retained the two above mentioned features, which remained unchanged during tens of cycles. Remarkably, no change in the intensity of the electrocatalytic wave was observed upon repeated N₂/CO₂ saturation cycles (Fig. S1†).

The preparation of the blue material was also achieved from the same CO₂-saturated DMF/H₂O (97 : 3 v/v) solution containing 1.3 mM [Cu(cyclam)]²⁺ and 0.1 M *n*-Bu₄NBF₄ but using a potentiostatic method. This allows optimization of the catalytic properties of the blue material *via* variations of the applied potential and the charge passed during electrodeposition. Each modified electrode was then assayed for CO₂ reduction during electrolysis of a CO₂-saturated [Cu(cyclam)]²⁺-free DMF/H₂O (97 : 3 v/v) solution at -2.0 V vs. Fc^{+/0}, with subsequent monitoring of the formation of the following products: in the gas phase, CO, methane and other volatile hydrocarbons as well as H₂ were monitored by gas chromatography; in the liquid phase, methanol was monitored by gas chromatography, formaldehyde by a colorimetric test, while formic acid, oxalate and glyoxylate were assayed by ion-exchange chromatography (see the ESI†). In all cases only formic acid and H₂ could be detected and quantified. Working with 3% water during electrolysis was critical. Indeed, electrolysis carried out in the absence of water gave almost no current (Fig. S2†), while in the presence of 10% water a larger current density (Fig. S2†) but also a larger formation of H₂ (faradaic yields: 60% for H₂ and 40% for formic acid) were obtained. This reflects, in one hand, the requirement of water as a source of protons for CO₂ reduction and, in another hand, the competition between proton and CO₂ reduction processes. It is also consistent with the H atoms of H₂ and formic acid deriving from H₂O exclusively.

From these optimization experiments, it resulted that the best results in terms of current density, stability and faradaic yield for potentiostatic CO₂ conversion into formic acid (blue trace in Fig. 2a and blue dot in Fig. 2b) were obtained with a blue material resulting from passage of a total charge (*Q*) of 4 C cm⁻² during the electrodeposition step, using [Cu(cyclam)]²⁺ as the precursor, with an applied potential of -2.0 V vs. Fc^{+/0}. Lower and larger *Q* values resulted in lower faradaic yields (Fig. 2b) and less stable currents (Fig. 2a). Electrodeposition at lower potentials (Fig. S3†) resulted in less active deposits. Such an optimum might result from the balance between the charge needed to deposit enough active material and the decrease in conductivity expected for thick electrode films. Furthermore CH₃CN/H₂O mixtures were also tested as the solvent but resulted into less active deposits (Fig. S4†) and electrolysis in this solvent mixture gave very low faradaic yields in formic acid (<30%). Using two different copper complexes, namely [Cu(terpy)₂]²⁺ or [Cu(TACN)]²⁺ (TACN = 1,4,7-triazacyclononane) as precursors under the established optimal conditions also led to stable blue electrodeposits, however much less active and less selective (Fig. S5†). Furthermore, substitution of a simple copper(II) salt for [Cu(cyclam)]²⁺ during electrodeposition led to the formation of a blue deposit with similar electroactive surface area, however less active and stable (Fig. S6†).

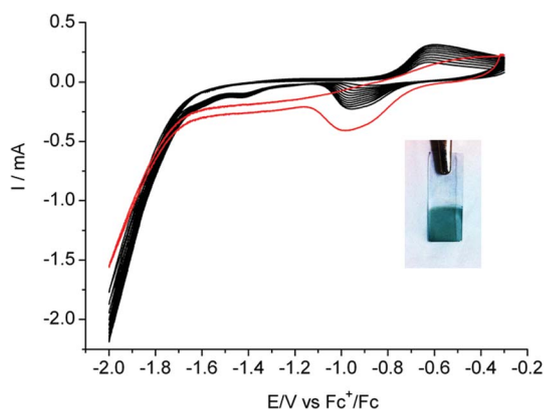


Fig. 1 Cyclic voltammograms recorded on a FTO electrode (1 cm²) in CO₂-saturated DMF/H₂O (97 : 3 v/v) solution and 0.1 M *n*-Bu₄NBF₄. The black trace was recorded in the presence of 1.3 mM [Cu(cyclam)](ClO₄)₂. The inset shows a picture of the electrode modified with the blue material obtained after recording the black trace. The red trace is obtained under similar conditions but in the absence of [Cu(cyclam)]²⁺ using the electrode after electrodeposition and washing of the blue material.

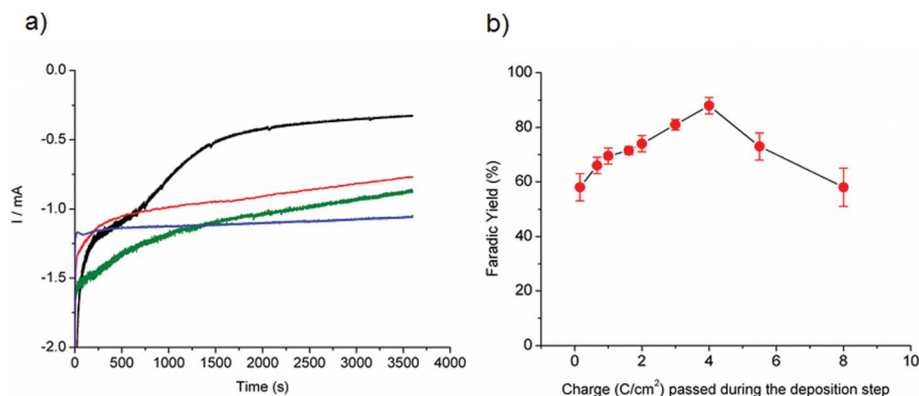


Fig. 2 (a) Evolution of the electrocatalytic current during potentiostatic CO_2 reduction assay at -2.0 V vs. Fc^+/Fc in DMF/ H_2O (97 : 3 v/v) solution measured on FTO electrodes modified with the blue material (1 cm^2) from $[\text{Cu}(\text{cyclam})]^{2+}$. The electrodes differed by the quantity of charge passed during the deposition step: 0.2 C (black trace), 1.6 C (red trace), 4.0 C (blue trace) and 8 C (green trace); (b) faradaic yield for formation of formic acid during potentiostatic CO_2 reduction assays (1 h) at -2.0 V vs. Fc^+/Fc in DMF/ H_2O (97 : 3 v/v) solution measured on modified FTO electrodes (1 cm^2). H_2 formation accounts for the complement of the faradaic yield to 100%. The blue dot shows the conditions for optimal yield. Errors bars indicate variations obtained during at least 5 experiments for each charge value.

In conclusion of these studies electrolysis of a CO_2 -saturated $[\text{Cu}(\text{cyclam})]^{2+}$ -free solution at -2.0 V vs. Fc^+/Fc using our best material generated a current density of about 1.3 mA cm^{-2} (blue trace in Fig. 2a) which only slowly decreased with time (0.8 mA cm^{-2} after 10 hours of continuous operation, Fig. S7†). Formic acid was produced with a faradaic yield of $88 \pm 3\%$, H_2 evolution accounting for the complement. No formic acid production could be detected when a blank FTO electrode was used under the same conditions (Fig. S7†). Interestingly, the electrodeposited material proved significantly more active, with larger current density values at steady state and higher faradaic yield for formic acid production, than an electro-polished polycrystalline Cu plate, a metallic Cu deposit²² or electro-deposited copper(i) oxide²³ with comparable electroactive surface areas, determined *via* standard methods (Fig. S8†).

Fig. 3 shows the Tafel plot of the blue material derived from linear sweep voltammetry measurement at a scan rate of 0.5 mV s^{-1} in a CO_2 -saturated DMF/ H_2O (97/3 v/v) solution. Determination of overpotential values requests calculation of the standard potential for CO_2/HCOOH in DMF/ H_2O solution, a parameter surprisingly not available in the literature value whereas the standard potential of the CO_2/CO couple in $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ and DMF/ H_2O media was recently reported.¹⁸ In the ESI† we indicate how our calculations led to a value of -0.78 V vs. NHE. For that purpose, solvation enthalpies for formic acid in various solvents were computed using the SMD implicit solvation method,²⁴ the M05-2X functional and the TZVP basis. The potential of the Fc^+/Fc couple in DMF ($E^{\text{DMF}}(\text{Fc}^+/\text{Fc})$) is 0.57 ± 0.02 V vs. NHE when taking into account the interliquid junction potential.¹⁸ This value was used to convert experimental electrochemical potential values into overpotential values in Fig. 3 (See the ESI†). Catalytic onset potential is -1.5 V vs. Fc^+/Fc which corresponds to an overpotential requirement of about 150 mV. The Tafel data are linear in the range $\eta = 200$ – 400 mV with exchange current density of $10^{-1.3} \text{ mA cm}^{-2}$ and slope of 120 mV per decade, a value consistent with a mechanism in

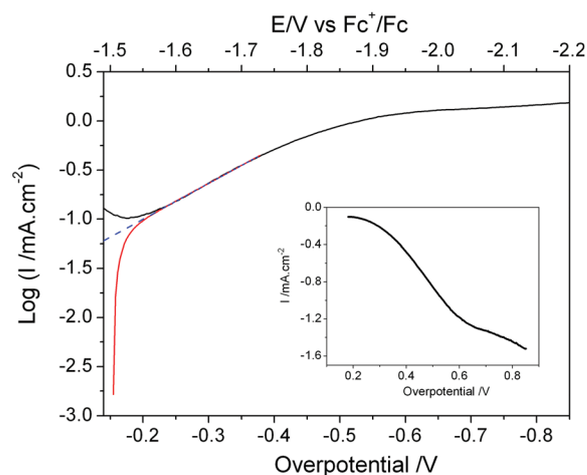


Fig. 3 Tafel plot obtained from linear sweep voltammetry experiments recorded at 0.5 mV s^{-1} on a blue material modified electrode (1 cm^2) obtained from $[\text{Cu}(\text{cyclam})]^{2+}$ in a CO_2 -saturated solution of DMF/ H_2O (97 : 3 v/v + $0.1 \text{ M } n\text{-Bu}_4\text{NBF}_4$). The corresponding polarization curve is shown in the inset. See the ESI† for the determination of the overpotential for formic acid production.

which the first electron transfer to adsorbed CO_2 is the rate-determining step.²⁵ The catalytic current then reaches a plateau ($\sim 1 \text{ mA cm}^{-2}$) for overpotentials larger than 540 mV and corresponding to the above-mentioned optimized conditions.

Characterization of the blue material on the most active electrode was carried out using a variety of methods. First, chemical analysis (see methods in the ESI†) revealed the presence of carbonate or bicarbonate ($2.0 \pm 0.2 \mu\text{mol cm}^{-2}$), formate ($1.6 \pm 0.1 \mu\text{mol cm}^{-2}$), Cu ions ($12.5 \pm 0.2 \mu\text{mol cm}^{-2}$) and cyclam ($4 \pm 2 \text{ nmol cm}^{-2}$).²⁶ When $[\text{Cu}(\text{terpy})_2]^{2+}$ or $[\text{Cu}(\text{TACN})]^{2+}$ were used as precursors, the deposits were also shown to contain a small amount of the organic ligand (2 ± 1 and $4 \pm 2 \text{ nmol cm}^{-2}$ respectively). Scanning electronic microscopy (SEM) showed regions where the FTO electrode is

coated by nanoparticles (~ 100 nm in diameter, Fig. 4) surrounded by significantly larger particles (few μm in diameter) consistent with a deposit thickness of $1.5 \mu\text{m}$ as determined by atomic force microscopy (AFM) (Fig. S9†). Optical microscopy confirmed this observation with the large particles and the background coating appearing as red and blue respectively. Energy Dispersive X-ray (EDX) spectroscopy indicated the presence of mainly Cu and O atoms in the large particles while the nanoparticles contained higher amounts of C, N and O with regard to Cu atoms (Fig. S10†). X-ray photoelectron spectroscopy (XPS) analysis (Fig. 4) confirmed the presence of carbon-based material: a broad peak with binding energies from 280 to 287 eV, in the C 1s orbital region, could be simulated with a mixture of C–N, C–O, C–H and C=O features. Nitrogen was also observed as a double peak between 394 and 400 eV in the N 1s region, likely reflecting a secondary amine as found in the cyclam ligand and quaternary ammonium ions from the supporting electrolyte. Finally analysis of the Cu 2p_{1/2} region

revealed the presence of Cu(I), in the form of Cu₂O, and/or metallic copper, together with Cu(II), potentially coordinated to formate or carbonate/bicarbonate as found in some coordination polymers.²⁷ X-ray absorption spectra collected at the copper K-edge indicate the presence of three contributions to the blue catalytic film. The main maximum of the XANES spectrum (around 8997 eV in Fig. 5A) is indicative of light ligand atoms (O/N/C); its low intensity renders octahedral coordination unlikely. The pronounced shoulder at 8982 eV suggests a major fraction of Cu(I) ions coordinated by two light ligands, as found in Cu₂O. By simulations we determined about 20% of copper metal (Fig. 5B) and about 30% of Cu(II) ions (Fig. 5A), likely coordinated by four light atoms (O/N/C); comparison with various CuO and Cu(II) compounds did not allow a more precise speciation of the latter (Fig. S11†). The remaining contribution was identified as Cu(I) ions, most likely coordinated by two light atoms (O/N/C). Regarding its local structure, the Cu(I) material could resemble copper(I) oxide (Cu₂O), but the long-range order

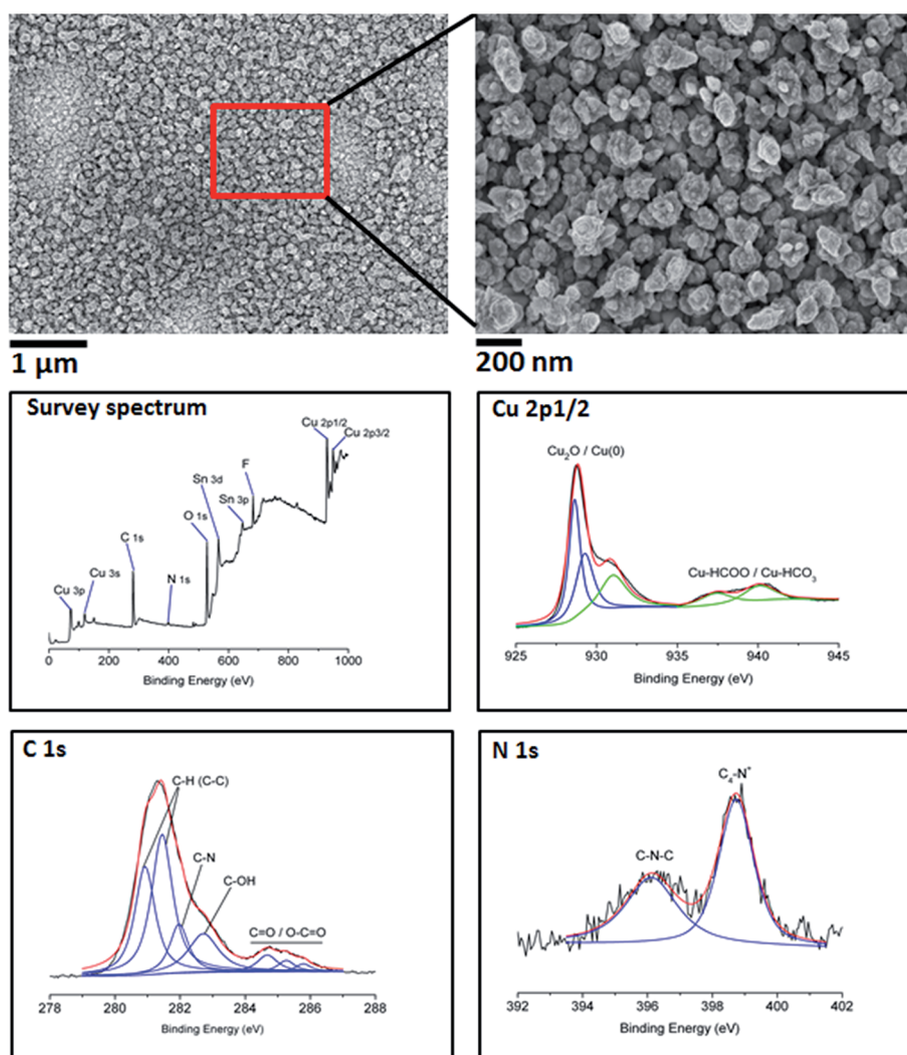


Fig. 4 Top: scanning electron micrograph of the blue material obtained from $[\text{Cu}(\text{cyclam})]^{2+}$ (left) with focus on the background nanoparticulate coating (right). Middle and bottom: XPS survey (middle-left), Cu_{2p} (middle-right), C_{1s} (bottom-left) and N_{1s} (bottom-right) core levels spectra of blue material deposited onto FTO; the black traces correspond to experimental data, the red traces to the simulated spectra obtained from the sum of the blue and green traces.

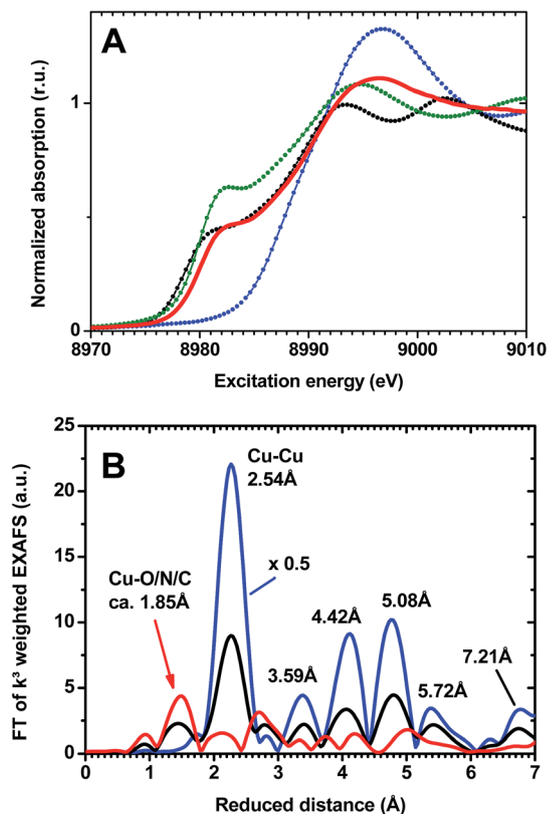


Fig. 5 Copper K-edge X-ray absorption spectra of the Cu-based blue material obtained from $[\text{Cu}(\text{cyclam})]^{2+}$ and of reference materials. (A) XANES region of the spectrum (red trace, Cu-based material, deposition of 1 C cm^{-2} ; black trace, Cu metal foil; green trace, Cu(I) oxide, Cu_2O ; blue trace, Cu(II) oxalate). The spectrum of the blue material can be described by a linear combination of the shown Cu references (about 20% Cu metal, 50% Cu(I) oxide and 30% Cu(II) salt). (B) Fourier-transforms of k^3 -weighted EXAFS spectra (black trace, blue material; blue trace, Cu metal foil, its FT amplitudes were multiplied by 0.5; red trace, blue material after correction for a metal contribution of 20%). The metal peaks are labeled by the corresponding Cu–Cu distance; the non-metallic peak of the blue material (red arrow) can be assigned to Cu–O/N/C coordination.

of crystalline Cu_2O was not detectable in the EXAFS (low-amplitude FT peaks at longer distance in metal-corrected spectra; Fig. S12[†]). The average distance of about 1.86 Å of the Cu–ligand distance as determined by EXAFS simulations of the metal-corrected spectra confirms a mixture of Cu(I) L_2 and Cu(II) L_4 coordination geometries ($\text{L} = \text{O/N/C}$, Table S1[†]).

We finally performed XRD measurements on the Cu-based blue materials (See Fig. 2 for corresponding catalytic activities). On the films grown with a passed charge less than 1 C cm^{-2} , no crystalline phase could be detected. By contrast, thicker films contain both crystallized metallic copper and copper(I) oxide. For example, films grown with a passed charge of 4 C cm^{-2} contain $\sim 20 \text{ nm}$ metallic copper crystallites and Cu_2O crystallites with size $< 5 \text{ nm}$ (Fig. S13[†]). This clearly suggests an influence of the thickness and structuration of the films on the performances.

The active blue material can thus be described as a composite mixture of metallic copper, copper(I) oxide (Cu_2O)

and an insoluble Cu(II)-based material as well as the organic ligand of the molecular Cu-based precursor. The most striking feature of this material resides in the fact that the activity is very much dependent on the choice of this ligand. On the other hand the best performances were obtained with films containing Cu and Cu_2O crystallites. Overall, this indicates that the blue material is unique in deriving its catalytic properties from synergetic contributions of its different, organic and inorganic, components. Considering the complex nature of the composite catalyst, further experiments are needed to understand its surface reactivity and reaction mechanisms. Whether a supported homogeneous catalysis, based on ligand-bound copper ions somehow integrated in the material or a metallic copper surface-based process, facilitated by the presence of organic molecules as reported in the case of cobalt-based water oxidizing catalytic materials,^{28–30} remains an open question. In the first case formic acid formation is likely to derive from CO_2 insertion into a metal-hydride intermediate.³¹ In the second case, the mechanism is likely to involve an adsorbed $\text{CO}_2^{\cdot-}$ intermediate.²⁵ More specifically, the exact role of the organic ligand in the activity of the blue material is intriguing. It might: (i) promote Cu active sites by coordination; (ii) participate in proton transfers as pendant amine functions do within electrocatalysts for H^+/H_2 interconversion³² or formate dehydrogenation;³³ (iii) capture CO_2 and increase its local surface concentration.^{34,35}

Conclusion

We have described a novel hybrid material, electrocatalytically active for the reduction of CO_2 . This material is promising for the following reasons, on route towards implementation of this new material into practical electrochemical cells:

- Its preparation through electrodeposition from a simple Cu(II) coordination complex is both straightforward and cost-effective.

- It exhibits high catalytic activity at moderate overpotentials. The overpotential requirement is $\sim 150 \text{ mV}$. It thus compares well with the recently described Bi-based material selectively forming CO from CO_2 in acetonitrile with similarly low overpotentials.^{13,14} We note that the thermodynamic CO_2 -reduction potential in common ionic liquids have been recently reported to be comparable to the value measured in CH_3CN .³⁶

- Remarkable selectivity for CO_2 reduction over H_2 evolution, high faradaic yield (90%) for the production of formic acid and significant stability upon turnover are observed under optimized conditions. Metallic electrodes have been previously extensively studied under alkaline aqueous conditions and some of them (Sn, Bi, ...) display similar selectivities for the production of formic acid.³⁷ By contrast, CO_2 reduction at metallic copper electrodes is known not to be as selective, whatever the nature of the electrolyte.³⁷ For example, metallic copper and Cu_2O electrodes assayed under DMF/ H_2O conditions (Fig. S7[†]) show lower selectivity for CO_2 -derived product as compared to the blue material, confirming previous studies carried out in MeOH.¹⁵ We thus report here on the first Cu-

based material displaying a high selectivity for formate production.

- The current density measured at $\eta = 540$ mV is about 1 mA cm^{-2} , which compares well to the activity of other Cu-based materials recently reported.^{38,39} Routes towards higher current densities, allowing for a cost-effective process, include nanostructuring of the catalytic layer as recently demonstrated for tin-based electrodes.^{40,41} Furthermore, the importance of the organic ligand derived from the precursor, even though its exact function is still ill-defined, provides another opportunity for optimization.

Methods

Preparation of [Cu(cyclam)]²⁺ solution

Cu(ClO₄)₂·6H₂O (19.6 mg, 53 μmol) was added to the solution of cyclam (10.4 mg, 53 mmol) in DMF (38.8 mL). Deionized water (1.2 mL) was added to the above solution to obtain DMF solutions with 3% water and 1.3 mM [Cu(cyclam)]²⁺. Then *n*-Bu₄BF₄ (1.316 g) was added to obtain 0.1 M electrolytic solutions. Solutions of [Cu(terpy)₂]²⁺ and [Cu(TACN)]²⁺ were prepared similarly from the corresponding ligands.

Preparation of blue material

Films of the blue material were grown on FTO electrode by controlled-potential electrolysis of freshly prepared [Cu(cyclam)]²⁺ solutions in DMF + 3% water (see above, 6 mL) saturated with CO₂ (CO₂ was continuously bubbled through the solution during electrodeposition). The electrolysis was typically performed at -2.0 V vs. Fc^{+/0} until a charge of 4 C has passed through the electrode. The blue material was then shortly immersed in fresh DMF for removing unreacted reagents on the surface.

Electro-catalytic CO₂ reduction assay

Electro-reduction of CO₂ was performed by applying a constant potential (typically -2.0 vs. Fc^{+/0}) to the working electrode dipped in a stirred, CO₂-saturated (through constant bubbling with a 5 mL min^{-1} flow) [Cu(cyclam)]²⁺-free DMF solution (3% water, 0.1 M *n*-Bu₄BF₄, 22 mL). A distinct procedure was used for detection and quantification of H₂ and CO: the solution was saturated (and the whole cell filled) with CO₂ though initial bubbling and then isolated during the electrocatalytic CO₂ assay. Some experiments have been performed with a smaller volume of electrolyte (8 mL) but, under such conditions, the catalytic current was less stable and the faradaic yield for production of formic acid dropped to 65%.

Acknowledgements

We acknowledge support from the French National Research Agency (ANR, Carbiored ANR-12-BS07-0024-03; Labex program ARCANÉ, ANR-11-LABX-0003-01 and DYNAMO, ANR-11-LABX-0011) and from Fondation de l'Orangerie for individual Philanthropy and its donors. We thank Jocelyne Leroy and Bruno Jousset (CEA/DSM/IRAMIS) for XPS measurements, Matthieu Fumagalli for AFM measurements. The XAS

experiments were carried out at beamline KMC-1 of the BESSY synchrotron operated by the Helmholtz Zentrum Berlin (HZB); we thank M. Mertin and Dr F. Schäfers (both HZB) for their support. JH and HD gratefully acknowledge financial support by the Berlin cluster of excellence on Unifying Concepts in Catalysis (UniCat).

References

- 1 A. M. Appel, J. E. Bercaw, A. B. Bocarsly, H. Dobbek, D. L. DuBois, M. Dupuis, J. G. Ferry, E. Fujita, R. Hille, P. J. A. Kenis, C. A. Kerfeld, R. H. Morris, C. H. F. Peden, A. R. Portis, S. W. Ragsdale, T. B. Rauchfuss, J. N. H. Reek, L. C. Seefeldt, R. K. Thauer and G. L. Waldrop, *Chem. Rev.*, 2013, **113**, 6621–6658.
- 2 C. Costentin, M. Robert and J. M. Saveant, *Chem. Soc. Rev.*, 2013, **42**, 2423–2436.
- 3 G. Centi, E. A. Quadrelli and S. Perathoner, *Energy Environ. Sci.*, 2013, **6**, 1711–1731.
- 4 E. V. Kondratenko, G. Mul, J. Baltrusaitis, G. O. Larrazabal and J. Perez-Ramirez, *Energy Environ. Sci.*, 2013, **6**, 3112–3135.
- 5 A. J. Morris, G. J. Meyer and E. Fujita, *Acc. Chem. Res.*, 2009, **42**, 1983–1994.
- 6 J. L. Qiao, Y. Y. Liu, F. Hong and J. J. Zhang, *Chem. Soc. Rev.*, 2014, **43**, 631–675.
- 7 R. Reske, M. Duca, M. Oezaslan, K. J. P. Schouten, M. T. M. Koper and P. Strasser, *J. Phys. Chem. Lett.*, 2013, **4**, 2410–2413.
- 8 H. Mistry, R. Reske, Z. Zeng, Z.-J. Zhao, J. Greeley, P. Strasser and B. R. Cuenya, *J. Am. Chem. Soc.*, 2014, DOI: 10.1021/ja508879j.
- 9 R. Reske, H. Mistry, F. Behafarid, B. R. Cuenya and P. Strasser, *J. Am. Chem. Soc.*, 2014, **136**, 6978–6986.
- 10 J. J. Wu, F. G. Risalvato, P. P. Sharma, P. J. Pellechia, F. S. Ke and X. D. Zhou, *J. Electrochem. Soc.*, 2013, **160**, F953–F957.
- 11 A. S. Agarwal, Y. M. Zhai, D. Hill and N. Sridhar, *ChemSusChem*, 2011, **4**, 1301–1310.
- 12 D. P. Summers, S. Leach and K. W. Frese, *J. Electroanal. Chem.*, 1986, **205**, 219–232.
- 13 J. L. DiMeglio and J. Rosenthal, *J. Am. Chem. Soc.*, 2013, **135**, 8798–8801.
- 14 J. Medina-Ramos, J. L. DiMeglio and J. Rosenthal, *J. Am. Chem. Soc.*, 2014, **136**, 8361–8367.
- 15 K. Nakata, T. Ozaki, C. Terashima, A. Fujishima and Y. Einaga, *Angew. Chem., Int. Ed.*, 2014, **53**, 871–874.
- 16 Y. Oh, H. Vrabel, S. Guidoux and X. L. Hu, *Chem. Commun.*, 2014, **50**, 3878–3881.
- 17 M. Asadi, B. Kumar, A. Behranginia, B. A. Rosen, A. Baskin, N. Reppin, D. Pisasale, P. Phillips, W. Zhu, R. Haasch, R. F. Klie, P. Král, J. Abiade and A. Salehi-Khojin, *Nat. Commun.*, 2014, **5**, 4470.
- 18 C. Costentin, S. Drouet, M. Robert and J. M. Saveant, *Science*, 2012, **338**, 90–94.
- 19 E. Bouwman, R. Angamuthu, P. Byers, M. Lutz and A. L. Spek, *Science*, 2010, **327**, 313–315.

- 20 C. Bucher, J. C. Moutet, J. Pecaut, G. Royal, E. Saint-Aman, F. Thomas, S. Torelli and M. Ungureanu, *Inorg. Chem.*, 2003, **42**, 2242–2252.
- 21 C. Bucher, J. C. Moutet, G. Pecaut, G. Royal, E. Saint-Aman and F. Thomas, *Inorg. Chem.*, 2004, **43**, 3777–3779.
- 22 W. Shang, X. Shi, X. Zhang, C. Ma and C. Wang, *Appl. Phys. A: Mater. Sci. Process.*, 2007, **87**, 129–135.
- 23 A. Paracchino, V. Laporte, K. Sivula, M. Gratzel and E. Thimsen, *Nat. Mater.*, 2011, **10**, 456–461.
- 24 A. V. Marenich, C. J. Cramer and D. G. Truhlar, *J. Phys. Chem. B*, 2009, **113**, 6378–6396.
- 25 M. Gattrell, N. Gupta and A. Co, *J. Electroanal. Chem.*, 2006, **594**, 1–19.
- 26 Following the suggestion of a reviewer, we have submitted the blue material to a 30 min air plasma treatment in order to specifically destroy organic molecules and in particular the cyclam ligand. HPLC titration indeed indicates almost quantitative removal of the ligand from the blue material during this process. The catalytic activity is concomitantly reduced to the level of an electrode fabricated without any ligand (see Fig. S14†), in agreement with the specific role of the ligand in promoting catalysis.
- 27 K. L. Hu, M. Kurmoo, Z. M. Wang and S. Gao, *Chem.–Eur. J.*, 2009, **15**, 12050–12064.
- 28 Y. Yamada, T. Miyahigashi, H. Kotani, K. Ohkubo and S. Fukuzumi, *Energy Environ. Sci.*, 2012, **5**, 6111–6118.
- 29 D. Hong, J. Jung, J. Park, Y. Yamada, T. Suenobu, Y.-M. Lee, W. Nam and S. Fukuzumi, *Energy Environ. Sci.*, 2012, **5**, 7606–7616.
- 30 D. Shevchenko, M. F. Anderlund, A. Thapper and S. Styring, *Energy Environ. Sci.*, 2011, **4**, 1284–1287.
- 31 M. Cokoja, C. Bruckmeier, B. Rieger, W. A. Herrmann and F. E. Kuhn, *Angew. Chem., Int. Ed.*, 2011, **50**, 8510–8537.
- 32 D. L. DuBois, *Inorg. Chem.*, 2014, **53**, 3935–3960.
- 33 B. R. Galan, J. Schöffel, J. C. Linehan, C. Seu, A. M. Appel, J. A. S. Roberts, M. L. Helm, U. J. Kilgore, J. Y. Yang, D. L. DuBois and C. P. Kubiak, *J. Am. Chem. Soc.*, 2011, **133**, 12767–12779.
- 34 I. Niedermaier, M. Bahlmann, C. Papp, C. Kolbeck, W. Wei, S. Krick Calderón, M. Grabau, P. S. Schulz, P. Wasserscheid, H.-P. Steinrück and F. Maier, *J. Am. Chem. Soc.*, 2013, **136**, 436–441.
- 35 Y. Jiao, Y. Zheng, S. C. Smith, A. Du and Z. Zhu, *ChemSusChem*, 2014, **7**, 435–441.
- 36 D. C. Grills, Y. Matsubara, Y. Kuwahara, S. R. Golisz, D. A. Kurtz and B. A. Mello, *J. Phys. Chem. Lett.*, 2014, **5**, 2033–2038.
- 37 Y. Hori, Electrochemical CO₂ Reduction on Metal Electrodes, *Modern Aspects of Electrochemistry*, Vol. 42, ed. G. Vayenas *et al.*, Springer, New York, 2008, pp. 89–189.
- 38 C. W. Li and M. W. Kanan, *J. Am. Chem. Soc.*, 2012, **134**, 7231–7234.
- 39 Z. F. Chen, P. Kang, M. T. Zhang, B. R. Stoner and T. J. Meyer, *Energy Environ. Sci.*, 2013, **6**, 813–817.
- 40 Y. H. Chen and M. W. Kanan, *J. Am. Chem. Soc.*, 2012, **134**, 1986–1989.
- 41 S. Zhang, P. Kang and T. J. Meyer, *J. Am. Chem. Soc.*, 2014, **136**, 1734–1737.