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# Propargyl carbamate-functionalized Cu(II)-MOF after reaction with chloroauric acid: an XPS data record

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A copper-containing metal organic framework (MOF) was prepared using the new organic linker  $5-(2-\{[(prop-2-yn-1-yloxy)carbonyl]-amino\}ethoxy)isophthalic acid (1,3-H<sub>2</sub>YBDC (where Y = alkYne and BDC = Benzene DiCarboxylate), and functionalized with gold particles by reaction with HAuCl<sub>4</sub> under thermal treatment in methanol. The resulting system was investigated by complementary techniques to obtain information on structure and morphology. In the present work, X-ray photoelectron spectroscopy (XPS) was employed to analyse the chemical composition of a representative specimen. Beside wide scan spectra, data obtained by the analysis of the C1s, O1s, N1s, Cu2p and Au4f signals are presented and critically discussed. The results highlight the reduction of Au(III) to Au(0) and to Au(I), with the latter oxidation state being the predominant one. Overall, the data presented herein may act as useful guidelines for the eventual tailoring of material properties and their possible implementation towards functional applications in heterogeneous catalysis.$ 

*Keywords:* Metal organic framework, copper, benzene dicarboxylate, propargyl carbamate, gold, X-ray photoelectron spectroscopy

#### INTRODUCTION

Metal-Organic Frameworks (MOFs) constituted by connecting metal ions with polytopic organic linkers have received enormous attention in the last decade. Their tuneable pore geometries and flexible frameworks have been successfully exploited in various research fields such as catalysis, gas absorption, gas storage or sensing (Refs. 1-9).

In recent years, our research group has shown that a propargyl carbamate  $[-N(H)C(O)O-CH_2-C\equiv CH]$  group anchored on different oxide supports (SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, TiO<sub>2</sub>, Fe<sub>3</sub>O<sub>4</sub>) is capable of reducing Au(III) to Au(0), yielding supported gold nanoparticles (AuNPs) without the addition of any external reducing and/or stabilizing agent (Refs. 10-13). Based on these results and considering that the chemical, structural, and functional behaviour of AuNPs are directly dependent on the physico-chemical environment dictated by the support, we envisaged that

Accession#:Enter Accession Number.

#### Technique: XPS

Host Material:  $[Cu(1,3-YBDC)] \cdot (C_{14}H_{11}NO_8Cu)$  and Au

**Instrument:** Perkin-Elmer Physical Electronics, Inc. 5600ci

Major Elements in Spectra: C, O, N, Cu

Minor Elements in Spectra: Au

Published Spectra: 6

Spectra in Electronic Record: 8

Spectral Category: comparison

the versatility demonstrated by the propargyl carbamate residue could be further exploited by anchoring it to solid supports different from oxides, for example within alkynyl-derivatized MOFs. In this regard, we showed that the reaction of the novel 5substituted organic linker 5-(2-{[(prop-2-yn-1-yloxy)carbonyl]amino}ethoxy)isophthalic acid (labelled 1,3-H<sub>2</sub>YBDC, where Y = alkYne and BDC = Benzene DiCarboxylate), bearing a propargyl carbamate substituent, with copper nitrate in refluxing 2-propanol leads to a new copper-based MOF Cu-YBDC in high yields. The novel material was fully characterized by complementary analytical techniques. X-ray diffraction (XRD) data revealed that Cu-YBDC contains a complex network of 5substituted isophthalate anions coordinated to Cu(II) centers belonging to the common paddlewheel dimeric structure with a Cu-Cu distance of 2.633 Å. Quite unexpectedly, the apical atom in the paddlewheel structure belongs to the carbonyl atom of the propargyl carbamate functionality, present with two equally populated alternative chain conformations. Such extracoordination by the propargyl carbamate groups drastically reduces the MOF porosity, as also confirmed by BET measurements. Despite this evidence suggests that the internal material pores are available only to a small extent to host reactive gold species, this system stands as an attractive candidate for the generation and anchoring of reduced gold species on the MOF surface. This capability can be, in turn, related to its uniform array of propargyl-carbamate residues branching out therefrom and acting as binding sites able to capture Au(III) ions and subsequently forming Au(0) clusters (Refs. 14-15).

In this context, the present study focuses on the reactivity of Cu-YBDC with HAuCl<sub>4</sub> under thermal treatment in refluxing methanol. The structure, morphology, and chemical composition of the developed system has been analysed by means of atomic absorption spectroscopy (AAS), thermogravimetric analysis (TGA), XRD, field emission scanning electron microscopy (FE-SEM). X-ray photoelectron spectroscopy (XPS) studies were performed on a selected representative sample using a standard AlK $\alpha$  X-ray excitation source. This investigation was carried out in order to gather information on the elemental composition, oxidation states and chemical environments characterizing the elements present in the investigated material.

## SPECIMEN DESCRIPTION (ACCESSION # FG82.)

Host Material:  $[Cu(1,3\mbox{-}YBDC)]\mbox{\cdot}(C_{14}H_{11}NO_8Cu)$  and Au

CAS Registry #: unknown

**Host Material Characteristics:** homogeneous; solid; polycrystalline; unknown conductivity; Powder

**Chemical Name:**  $[Cu(1,3-YBDC)] \cdot (C_{14}H_{11}NO_8Cu)$  and Au **Source:** In a 100 mL round-bottom flask, a suspension of 1,3-H<sub>2</sub>YBDC (0.120 g, 0.390 mmol) and Cu(NO<sub>3</sub>)<sub>2</sub>·2.5H<sub>2</sub>O (0.163 g, 0.701 mmol) in 2-propanol (15 mL) was refluxed at 83 °C under stirring (300 rpm) for 24 h and then cooled down to room temperature. The precipitate was collected by Buchner filtration and washed with 2-propanol (2×10 mL). The turquoise microcrystalline powder was subsequently dried in an oven at 70 °C for 24 h, subsequently kept under vacuum (0.02 bar) for 24 h and stored under nitrogen to give 0.142 g of [Cu(1,3-YBDC)]·xH<sub>2</sub>O (yield = 90% based on the organic acid and considering the Cu-MOF dihydrate).

# Host Composition: C, H, N, O, Cu and Au

Form: powder

**Structure:** The system structure was investigated by complementary techniques (AAS, TGA, XRD, FE-SEM) and the resulting data compared with the pristine Cu-MOF. The amount of adsorbed gold by was 0.8 wt.%. The TGA plot for the Aucontaining Cu-MOF showed that, similarly to the starting Cu-MOF, two decomposition steps occur in the range 200-450 °C range with a final residue of  $\approx$  30 wt%, *i.e.*, a 4 wt% higher than in the absence of gold. XRD patterns of Cu-YBDC and Cu-MOF/Au are identical, except for the fact that in Cu-MOF/Au the peak at 12.82° (attributed to Cu<sub>2</sub>(OH)<sub>3</sub>(NO<sub>3</sub>) contaminating species) is more intense. FE-SEM images of Cu-YBDC/Au (see

the inset in figure Accession # FG82-01) revealed the predominant presence of prismatic aggregates.

**History & Significance:** HAuCl<sub>4</sub>×3H<sub>2</sub>O (0.050 g, 0.127 mmol) was added to a suspension of Cu-YBDC (0.250 g) in MeOH (120 mL). The reaction mixture was stirred for 1 h at room temperature, and the solid was subsequently separated from the supernatant by centrifugation at 5400 rpm for 15 min. After three cycles of washing with MeOH (3×20 mL) and centrifugation, the product was stirred in MeOH (30 mL) at 65°C for 1 h. The solid was again separated by centrifugation and washed with MeOH.

As Received Condition: As grown.

Analyzed Region: Same as host material

**Ex Situ Preparation/Mounting:** Sample fixed on an adhesive tape and directly introduced into the analysis chamber through a fast entry.

In Situ Preparation: None

**Charge Control:** No flood gun was used during analysis. For further details on the charging correction procedure, see Data Analysis Methods, Energy Scale Correction.

Temp. During Analysis: 298 K

Pressure During Analysis: <10<sup>-4</sup> Pa

Pre-analysis Beam Exposure: 180 s.

# **INSTRUMENT DESCRIPTION**

Manufacturer and Model: Perkin-Elmer Physical Electronics, Inc. 5600ci

Analyzer Type: spherical sector

Detector: Channeltron

Number of Detector Elements: 16

# INSTRUMENT PARAMETERS COMMON TO ALL SPECTRA

Spectrometer

Analyzer Mode: constant pass energy

Throughput (T=E<sup>N</sup>): N=0

Excitation Source Window: 1.5 micron Al window

Excitation Source: Al Ka

Source Energy: 1486.6 eV

Source Strength: 200 W

**Source Beam Size:** > 25000 µm x > 25000 µm

Signal Mode: multichannel direct

■Geometry

Incident Angle: 9 °

Source-to-Analyzer Angle: 53.8 °

Emission Angle: 45 °

Specimen Azimuthal Angle: 0 °

Acceptance Angle from Analyzer Axis: 0 °

Analyzer Angular Acceptance Width: 14° × 14°

## ∎Ion Gun

Manufacturer and Model: PHI 04-303 A

**Energy:** 4000 eV

Current: 0.4 mA/cm2

Current Measurement Method: Faraday cup

Sputtering Species: Ar<sup>+</sup>

Spot Size (unrastered): 250 µm

Raster Size: 2000 µm x 2000 µm

Incident Angle: 40 °

Polar Angle: 45 °

Azimuthal Angle: 111 °

**Comment:** differentially pumped ion gun

## DATA ANALYSIS METHOD

**Energy Scale Correction:** Binding energy (BE) values were corrected for charging by assigning to the adventitious C1s peak a BE of 284.8 eV (Ref. 16).

# Recommended Energy Scale Shift: -2.4 eV.

**Peak Shape and Background Method:** Gaussian-Lorentzian functions were used for peak fitting, with a typical composition value of ~80 % Gaussian and ~20 % Lorentzian. No G/L constraints were applied during the fitting. A Shirley background function was used.

**Quantitation Method:** Atomic concentrations were calculated by peak area integration, using sensitivity factors provided by PHI V5.4A software.

#### ACKNOWLEDGMENTS

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## AVAILABILITY OF DATA

The data that supports the findings of this study are available within the article and its supplementary material.

#### **CONFLICT OF INTEREST**

The authors have no conflicts to disclose.

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SPECTRAL FEATURES TABLE							
Spectrum ID #	Element/ Transition	Peak Energy (eV)	Peak Width FWHM (eV)	Peak Area (eV x cts/s)	Sensitivity Factor	Concentration (at. %)	Peak Assignment
FG82-02ª	C 1s	284.8	2.1	12730.2	0.296	28.3	Aliphatic carbon arising from adventitious surface contamination
FG82-02ª	C 1s	286.2	2.2	9099.5	0.296	20.2	C-N and C-O bonds in the ligand skeleton
FG82-02ª	C 1s	288.1	1.9	2893.6	0.296	6.4	-COOH functionalities bonded to the aromatic ring
FG82-02ª	C 1s	289.5	2.1	3771.2	0.296	8.4	-NCOO- groups present in the ligand lateral chain
FG82-03 <sup>b</sup>	O 1s	530.2	1.8	2941.3	0.711	2.6	ligand oxygen atoms bonded to copper and O in Cu <sub>2</sub> (OH) <sub>3</sub> (NO <sub>3</sub> )
FG82-03 <sup>▶</sup>	O 1s	531.9	1.8	10814.81	0.711	9.7	C-O single bonds, chemisorbed hydroxyl groups and OH groups in Cu <sub>2</sub> (OH) <sub>3</sub> (NO <sub>3</sub> )
FG82-03 <sup>b</sup>	O 1s	533.3	2.6	18298.2	0.711	16.4	C=O groups in carboxylate functionalities and/or adsorbed water molecules
FG82-04	N 1s	400.4	2.6	2810.7	0.477	3.7	CuN coordination
FG82-05 FG82-05 FG82-05	Cu 2p Cu 2p <sub>3/2</sub> Cu 2p <sub>1/2</sub>	932.8 952.6	2.4 2.4	2380.0 1262.3	5.321	0.9 <sup>ª</sup>	Cu(I) species Cu(I) species Cu(I) species
FG82-05° FG82-05 FG82-05	Cu 2p Cu 2p <sub>3/2</sub> Cu 2p <sub>3/2</sub>	934.8 954 6	2.7 3 1	6189.8 2524 6	5.321	2.4 <sup>e</sup>	Cu(II) species Cu(II) species Cu(II) species
FG82-06 <sup>f</sup> FG82-06	Au 4f Au 4f Au 4f <sub>7/2</sub>	84.0	1.8	988.9	6.250	0.2 <sup>g</sup>	Au(0) Au(0)
FG82-06 FG82-06 <sup>f</sup>	Au 4f <sub>5/2</sub> Au 4f	87.6	1.8	741.7	6.250	0.5 <sup>h</sup>	Au(0) Au(I)
FG82-06 FG82-06	Au 4f <sub>7/2</sub> Au 4f <sub>5/2</sub>	85.3 88.9	1.7 1.7	2159.1 1619.2			Au(I) Au(I)
FG82-06 <sup>f</sup>	Au 4f				6.250	0.3 <sup>i</sup>	Au(ÌIÍ)
FG82-06	Au 4f <sub>7/2</sub>	86.4	1.8	1537.5			Au(III)
FG02-00	Cu LMM	90.0 915.0 <sup>j</sup>	1.0	1153.1			Cu species

<sup>a</sup>The sensitivity factor is referred to the whole C 1s signal.

<sup>b</sup>The sensitivity factor is referred to the whole O 1s signal.

<sup>c</sup>The sensitivity factor is referred to the whole Cu 2p signal.

<sup>d</sup>The atomic percentage is referred to the sole Cu(I) species. <sup>e</sup>The atomic percentage is referred to the sole Cu(II) species.

The sensitivity factor is referred to the whole Au4f signal.

<sup>9</sup>The atomic percentage is referred to the sole Au(0) species.

<sup>h</sup>The atomic percentage is referred to the only Au(I) species.

<sup>i</sup>The atomic percentage is referred to the only Au(III) species.

The peak position is given in KE.

**Footnote to Spectrum FG82-02:** High-resolution C1s spectra reveal the presence of different components: BE = 284.8 eV ( $\approx 45 \%$  of the total C signal), attributed to aliphatic carbon arising from adventitious surface contamination; BE = 286.2 eV ( $\approx 32 \%$  of the total C), assigned to C-N and C-O bonds in the ligand skeleton; BE = 288.1 eV ( $\approx 10 \%$  of the total C signal), related to -COOH functionalities bonded to the aromatic ring; and BE = 289.5 eV ( $\approx 13 \%$  of the total C signal), assigned to -NCOO- groups present in the ligand lateral chain (Refs. 17-19).

**Footnote to Spectrum FG82-03:** Three different peaks are concurring to the overall intensity of the O1s signal,: BE = 530.2 eV ( $\approx$  9 % of the total O signal), associated to ligand oxygen atoms bonded to copper (Ref. 17) and O in Cu<sub>2</sub>(OH)<sub>3</sub>(NO<sub>3</sub>) (Ref. 20); BE = 531.9 eV ( $\approx$  34 % of the total O signal), ascribed to C-O single bonds, chemisorbed OH hydroxyl groups and OH groups in Cu<sub>2</sub>(OH)<sub>3</sub>(NO<sub>3</sub>) (Refs. 18, 20-22); and BE = 533.3 eV ( $\approx$  57 % of the total O signal), attributed to the presence of C=O groups in carboxylate functionalities and/or adsorbed water molecules (Refs. 18-19). The broadening of this latter peak is due to the presence of water molecules in slightly different surface environments.

**Footnote to Spectrum FG82-04:** A single peak signal is determined in the N1s spectra, centered at BE = 400.4 eV. This binding energy value indicates a strong ligand coordination between nitrogen atoms and copper metal centers in the investigated material (Ref. 23).

**Footnote to Spectrum FG82-05:** Information on the Cu chemical state can be gained by a careful evaluation of the copper signal. As performed in Ref. 24, both Cu2p<sub>3/2</sub> and Cu2p<sub>1/2</sub> peaks can be fitted with two components. The features at BE values of 932.8 and 952.6 eV are attributed to Cu2p<sub>3/2</sub> and Cu2p<sub>1/2</sub> spin-orbit components of Cu(I) species, while those peaking at 934.8 and 954.6 eV are related to Cu(II) species into the copper-containing metalorganic framework (Refs. 17, 22, 25). Peak analysis indicates that Cu(II) are the dominant species (*i.e.*, more than 70 % of the overall copper). A further confirmation of this attribution is given by the presence of intense *shake-up* satellites peaking at BE values  $\approx$  9.5 eV higher than the main spin-orbit components, which are a finger-print for the predominant presence of *d*<sup>0</sup> copper (II) centers (Refs. 24, 26-27). When only Cu(I) species (*d*<sup>10</sup>, a closed-shell system) are present, such satellites are not detected (Refs. 16, 23). The evaluation of the Auger parameter, defined as  $\alpha_{Cu} = BE(Cu2p_{3/2}) + KE(CuLMM)$  (Ref. 16) (where KE = kinetic energy), yielded a value of 1850.0 eV, intermediate between those reported for Cu(I) and Cu(II) species (Ref. 26). **Footnote to Spectrum FG82-06:** The fitting of the Au 4f peak enabled to recognize the co-presence of: (i) Au(0) species (BE(Au4f<sub>7/2</sub>) = 84.0 eV and BE(Au4f<sub>5/2</sub>) = 87.6 eV,  $\approx$  21 % of the total Au4f signal) (Refs. 13, 18); (ii) Au(I) species (BE(Au4f<sub>7/2</sub>) = 85.3 eV and BE(Au4f<sub>5/2</sub>) = 88.9 eV,  $\approx$  46 % of the total Au4f signal) (Refs. 16, 18-19); and (iii) Au(II) species (BE(Au4f<sub>7/2</sub>) = 86.4 eV and BE(Au4f<sub>5/2</sub>) = 90.0 eV,  $\approx$  33 % of the total Au4f signal) (Refs. 16, 18-19).

ANALYZER CALIBRATION TABLE							
Spectrum ID	Element/	Peak Energy	Peak Width	Peak Area	Sensitivity	Concentration	Peak
#	Transition	(eV)	FWHM (eV)	(eV x cts/s)	Factor	(at. %)	Assignment
figCalib01 <sup>a</sup>	Au4f <sub>7/2</sub>	84.0	1.4	186403			Au(0)
figCalib02 <sup>a</sup>	Cu2p <sub>3/2</sub>	932.7	1.6	86973			Cu(0)

<sup>a</sup>The peak was acquired after Ar<sup>+</sup> erosion.

GUIDE TO FIGURES						
Spectrum (Accession) #	Spectral Region	Voltage Shift*	Multiplier	Baseline	Comment #	
FG82-01	Survey	+2.4	1	0		
FG82-02	C 1s	+2.4	1	0		
FG82-03	O 1s	+2.4	1	0		
FG82-04	N 1s	+2.4	1	0		
FG82-05	Cu 2p	+2.4	1	0		
FG82-06	Au 4f	+2.4	1	0		

\*Voltage shift of the archived (as measured) spectrum relative to the printed figure.



Publish i	n S <i>urface</i>	Science	Spectra:	Yes	$\boxtimes$	No	
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Accession #	FG82-01	
Host Material	[Cu(1,3-YBDC)]·(C₁₄H₁₁NO₀Cu) and Au	
Technique	XPS	
Spectral Region	survey	
Instrument	Perkin-Elmer Physical Electronics, Inc. 5600ci	
Excitation Source	Al Ka	
Source Energy	1486.6 eV	
Source Strength	200 W	
Source Size	> 25 mm x > 25 mm	
Analyzer Type	spherical sector analyzer	
Incident Angle	9°	
Emission Angle	45°	
Analyzer Pass Energy	187.85 eV	
Analyzer Resolution	1.9 eV	
Total Signal Accumulation Time	975.6 s	
Total Elapsed Time	1073.2 s	
Number of Scans	30	
Effective Detector Width	1.9 eV	







Publish in SSS: Yes ⊠ No □ ■ Accession #: FG82-04 Host Material: [Cu(1,3-YBDC)]-(C14H11NO8Cu) and Au Technique: XPS Spectral Region: N 1s Instrument: Perkin-Elmer Physical Electronics, Inc. 5600ci Excitation Source: Al Ka Source Energy: 1486.6 eV Source Strength: 200 W Source Size: >25 mm x >25 mm Analyzer Type: spherical sector Incident Angle: 9 ° Emission Angle: 45 ° Analyzer Pass Energy 58.7 eV Analyzer Resolution: 0.6 eV Total Signal Accumulation Time: 169.0 s Total Elapsed Time: 185.9 s Number of Scans: 21 Effective Detector Width: 0.6 eV Publish in SSS: Yes 🛛 No 🗆





