Thermal decomposition of CuProp$_2$: In-situ analysis of film and powder pyrolysis

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ARTICLE INFO

Keywords:
- Copper propionate
- Pyrolysis
- Thermogravimetric analysis
- Evolved gas analysis
- Metal carboxylates
- CSD methods

ABSTRACT

The thermal decomposition of CuProp$_2$ in the form of film and powder was studied in different atmospheres by means of thermal analysis techniques (TG-MS, TG-IR, EGA), chemical-structural methods (FTIR, XRD, EA) and computational thermochemistry (VASP/PBE). The decomposition mechanism in terms of volatiles evolved was disclosed with the aid of ab-initio modeling; it was found to be dependent on the gas diffusion in and out of the sample and accelerated by a humid atmosphere. In films, the copper redox behavior showed sensitivity to the residual atmosphere. Finally, the role of the metal center is discussed in the frame of a general decomposition mechanism for metal propionates.

1. Introduction

The study of the thermal decomposition of metal carboxylates has been growing in the past years as a consequence of their role in the synthesis of functional oxide thin films. Among the latter, YBa$_2$Cu$_3$O$_7$ (YBCO) belongs to the II generation of High Temperature Superconductors (HTS) [1]; it is processed for tapes applications in fields such as energy transport and electrical power devices [2]. However, nowadays, its use for these applications is limited by its processing, so far Cu(CH$_2$CH$_2$CO)$_2$ (CuProp)$_2$ decomposition has only been studied for samples in the form of powder. Additionally, while several references can be found for the thermal analysis of CuProp$_2$ [15] or other carboxylates [18,19], an extensive analysis in terms of atmospheres and decomposition volatiles has not yet been done.

Volatile species evolved during decomposition can be detected by means of thermogravimetry (TG) coupled with Evolved Gas Analysis (EGA), which mainly works by infrared (EGA-FTIR or TG-FTIR) or by mass spectrometry (EGA-MS or TG-MS) detection. In fact, by EGA-FTIR and EGA-MS, it was shown that the decomposition of propionates in inert atmospheres involves formation of radicals CuH$_2$ and CuH$_2$CO (along with CO$_2$) and their recombination to form a symmetrical ketone [20–22]. For longer chain salts such as Ca(II)decanoate, techniques like pyrolysis coupled to gas chromatography (GC) revealed a more complicated scenario where normal alkanes were found along with several symmetrical ketones [23]. For odd chain length of Cu(II)carboxylates like Cu(II)propionate, only MS has been used for the in-situ EGA analysis and 2-pentanone (asymmetrical ketone) was reported to be the main decomposition product [11]. Formation of this asymmetrical ketone implies cleavage of a CeC(C=O) bond, instead of the expected CeC=O and C=O=O bonds. Conversely, no ketones at all were observed for even chain carboxylates of copper(II) [24], mercury(II) [25] and silver(I) [26]. In fact, references [24,25] report on the
decomposition of even chain length of Hg(II) and Cu(II) carboxylates as involving formation of ethylene and propionic acid along with CO₂. Clearly, there is no univocal decomposition pathway for metal carboxylates, on the contrary it depends on the surrounding atmosphere, the length of the carboxylate chain and the type of metal ion [27].

The aim of this paper is to disclose the thermal decomposition mechanism of CuProp. For this purpose, thermogravimetry combined with in-situ EGA analysis is used and different atmospheres are considered. Also, samples in the form of film and powder are analyzed and characterized by Fourier-transform infrared spectroscopy (FTIR), elemental analysis (EA) and X-ray diffraction (XRD). In situ TG-FTIR, TG-MS and EGA-MS allow an almost univocal identification of the volatiles, while the comparison of experimental enthalpies with theoretical data will show which of the proposed decomposition reactions takes place in a given atmosphere.

We will show that CuProp decomposes releasing the corresponding acid as major product and we propose a mechanism that is compatible with the observed phases and detected volatiles. We also show that the kinetics depends on sample preparation and atmosphere; in particular that decomposition is significantly enhanced in the presence of water vapor and in films.

2. Materials and methods

The initial solution was obtained by dissolving copper(II) acetate (Cu(CH₃COO)₂) Sigma Aldrich, 0.75 M), dried in vacuum at 60 °C, in a 1:1 mixture of methanol (VWR, ≥99.8%) and propionic acid (Merck, ≥99%). Deposition of that solution over 10 × 10 mm² or 5 × 5 mm² LaAlO₃ (LAO) substrates was followed by drying at 95 °C on a hot plate for a few minutes. The initial product after deposition is expected to be Cu(CH₃COO)₂ (CuProp), since in an excess of propionic acid, acetate groups are replaced by propionates [28]. The corresponding powder was recovered upon solvent evaporation in air at 90 °C. For comparison, CuProp in the form of powder and film was studied by FTIR, XRD and elemental analysis; no differences were observed (See Fig. S2). A single crystal was also obtained after a few-month storage of the solution at room temperature; it was recovered and dried at 30 °C and low vacuum (635 mmHg) prior to analysis.

The “nominal film thickness” h is determined from the mass (m) of the sample, the substrate area (A), and the particle density (p) of the final product (CuO) as h = m/A. Calculated thicknesses for the films used in this work ranged from 1.9 to 2.1 µm.

Since the density of the films is lower than the particle density, the nominal thickness is a lower-bound estimation of the real thickness of the pyrolyzed film.

TG analysis of CuProp as film and in the form of powder was carried out in a Mettler Toledo thermo-balance, model TGA/DSC1. Samples were heated at a constant rate of 5 °C/min under a dynamic atmosphere (flow rate of 55 ml/min of carrier gas and 15 ml/min of the protective gas). Water-saturated gas was obtained by bubbling the carrier gas in a water flask at standard temperature and pressure (25 °C, 1 atm). Under these conditions the saturated vapor partial pressure is 3.1% [29]. LAO substrates were used for films deposition, whereas uncovered 70-µl-Al₂O₃ pans were used for the powder sample. In-situ TG-FTIR or TG-MS were performed to monitor the evolution of the volatiles during sample decomposition. In the first case, the TG was coupled to the gas cell (Bruker ALPHA model) through a 40-cm-long steel tube kept at 200 °C. For TG-MS a steel capillary heated up to 200 °C was connected from the TG gas outlet to the MS chamber and the fragments were detected by means of a quadrupole mass analyzer (MKS model Microvision Plus).

In the case of samples decomposed directly in vacuum (EGA-MS), volatiles are detected without the need of a steel capillary: the film or crucible with the powder are placed on a platinum plate inside a quartz tube that is pumped to a total pressure of 10⁻⁶ bar by means of a rotary pump in series with a turbomolecular pump. The sample temperature is measured with a thermocouple soldered to the platinum plate and controlled with an external tubular furnace placed around the quartz tube. Interpretation of the MS spectra of the volatiles was done according to NIST reference spectra, but it should take into account that the fragmentation pattern of the anlyte can show temperature dependence in favor of lower molecular weight fragments [30,31].

X-ray diffraction curves (XRD) of powders and films (the latter were peeled off to be measured) were collected using an X-ray beam wave-length of 1.5406 Å (Cu-Kα) from two different diffractometers: a Siemens diffractometer (model D-5000) from Bruker, operating at a 35-ma current and at a voltage of 45 kV; and a D8 ADVANCE diffractometer from Bruker AXS, with an X-ray beam of Cu-Kα, operating at a 40-kV voltage and with a 40-ma current. Single-crystal X-ray diffraction was performed with a three-circle diffractometer (Bruker D8 QUEST ECO) system equipped with a Ceramic x-ray tube (Mo Kα, λ = 0.71076 Å) and a doubly curved silicon crystal monochromator. The measurements were made at 50 kV and 20 mA and x-rays collected by a PHOTON II (CPAD, Charge-Integrating Pixel Array Detector) detector.

Infrared analysis of solid samples was performed using a Fourier Transform Infrared Spectrometer (FTIR, Bruker model ALPHA) equipped with an attenuated total reflection (ATR) unit (model Platinum), while a Perkin Elmer 2400 series elemental analyzer was used for Elemental Analysis (EA).

Six TG-FTIR/MS experiments were conducted at atmospheric presSure-sure varying the flowing gas: CuProp as film was decomposed in humid O₂ (experiment A), humid N₂ (B) dry O₂ (C) and dry N₂ (D); the powder sample (labeled with ‘*’) was decomposed in dry O₂ (E*) and dry N₂ (D*). To avoid N₂ interference with CO detection (both with m/z = 28), D* was performed in argon for TG-MS. EGA-MS experiments were run on the film (E) and powder (E*, E”) samples, at 5 °C/min, 10 and 25 ºC/min, respectively.

Ab-initio theoretical calculations were performed within the framework of Density Functional Theory (DFT) as implemented in the VASP (Vienna Ab-initio Software Package) code [32,33]. All models are periodically replicated through space in all directions, in order to obtain a more accurate description of the solid phases investigated experimentally, as no boundary effects are to be taken into consideration. The theoretical investigation of CuProp and CuO was designed to ac-curately model their electronic structure as a bulk solid. Database structure files were converted into periodic models. The experimental unit cell was relaxed towards the computational minimum energy structure, identified using a built-in DIIS minimization algorithm with a convergence force threshold of 10⁻² eV/Å. The electronic structure was evaluated differently for gas phase and solid phase systems. For the former, a single K-point was considered sufficient to evaluate the electronic structure, but the unit cell size having been optimized to minimize the interaction between different replicas of the molecule. For the latter, a K-point grid was obtained using the Monkhorst-pack method and its size was optimized until further increases did not cor-respond to any change in total energy. More details can be found in Supp. info.

3. Results

3.1. Characterization of the initial product

In excess propionic acid, acetate groups are replaced by propio-nates. In fact, the elemental analysis results of the dry salt are in good agreement with the expected values for CuProp (Table 1), and with the EA results of CuProp obtained from an acetate-free solution (by re-acting CuCO₃ in excess propionic acid, see Supp. Info, Table S1). Ad-ditionally, the FTIR analysis of solid samples (Fig. 1) coincides with the reported assignments in [34]. The single-crystal analysis (see Supp. Info, Fig S1-a) performed on our single-crystal corresponds to the
Table 1
Weight percentages determined from Elemental Analysis of initial and decomposition products. (-) values inferior to detection limits.

| Compound                      | Found (Expected), mass % | %C  | %H  |
|-------------------------------|--------------------------|-----|-----|
| CuProp₂                       | 33.71 (34.37)            | 4.29 (4.81) |
| A (Film, hum O₂, 210 °C)      | 4.70                     | 0.38 |
| A (Film, hum O₂, 500 °C)      | –                        | –   |
| D (Film, dry N₂, 230 °C)      | 4.85                     | 1.32 |
| D (Film, dry N₂, 500 °C)      | –                        | –   |
| D’ (Powder, dry N₂, 500 °C)   | 3.56                     | –   |

Anhydrous CuProp₂ database structure (CCDC 1133510) [35], which consists of Cu dimers connected by bridging carboxylate ligands. Since the XRD powder-diffraction curve of the ground crystal (Fig. S1-c) coincides with that of our films (Fig. 1, inset), this structure (CCDC 1133510) was used later on for ab-initio calculations. In fact, our films’ magnetic behavior (see Supp. Info) is as expected for binuclear copper complexes where an antiferromagnetic super-exchange interaction exists between the Cu(II) atoms [36–38] due to bridging ligands, and which is responsible for the lower magnetic moment, μₑₐ, in CuProp₂ crystalline powders with respect to Cu(II) compounds in aqueous solutions (exp. range of 1.9–2.2 μₑ) [39,40]. In fact, by comparing the distance between the asymmetric and symmetric stretching band of the carboxylate bond [41,42] [Δν= νₘ(COO⁻) − νₛ(COO⁻)] of our CuProp with those of other carboxylates from literature, our ν = 165 cm⁻¹ value suggests a bridging ligand coordination [43,44].

Coordination water is expected to change the structure, but the OH stretching band of water does not increase significantly over time, indicating that CuProp₂ is not very hygroscopic.

3.2. Thermal decomposition of CuProp₂ at atmospheric pressure in film

During sample preparation, no water uptake is observed after drying the film at 95 °C. Therefore, no dehydration is observed in the thermogravimetric analysis and DSC (Fig. 2a and b, respectively); the first mass loss step corresponds to the ligand decomposition. The main volatiles evolved during this step are identified in the TG-FTIR spectra in Fig. 3.
and their detailed evolution as a function of atmosphere and temperature is shown in Fig. 4.

The general decomposition mechanism for films involves two stages (labeled as I and II in Fig. 2a) to yield CuO at 500 °C (Fig. 5a exp. A.C.D). Most of the mass loss (~92%) takes place during the first stage and involves formation of propionic acid (Fig. 3). From Fig. 2a, the first stage is not influenced by the presence of O₂ but it is accelerated by the presence of a humid atmosphere, and in fact step I in A (humid O₂) and B (humid N₂) ends at 190 °C, whereas it ends 10 °C above in C (dry O₂) and D (dry N₂). Therefore, H₂O vapor shifts the TG curve to low tem-peratures because it favors the hydrolysis of the salt through reaction scheme a (Table 2); this shift is enhanced for thinner films (Supp. Info., Fig. S9), indicating water vapor diffusion to be the reaction-rate lim-iting mechanism. So, for instance, in humid O₂, CuProp XRD refection-disappear and CuO crystallizes between 185 and 210 °C (Fig. 5b, A). Conversely, in dry N₂, CuO crystallization is delayed to 195 °C–215 °C (Fig. 5b, D). For any atmosphere, a change in the Cu-Prop: structure is observed during its decomposition as vcoo-de-creases from 165 to 125 cm⁻¹ (Fig. 6 and Supp. Info., Fig. S5). Similarly, the molar magnetic susceptibility at different decomposition stages (Supp. Info.) shows that the antiferromagnetic interactions decrease with increasing paramagnetic Cu(II) contribution, probably due to the breaking of the copper(II) carboxylate dimers [45].

After the first stage, a plateau in the TG curve is observed at m/m₀=42% (200–230 °C), corresponding to an intermediate product with a change of decomposition mechanism. The films quenched at 200–230 °C are dark, and in all atmospheres their FTIR absorptions (Fig. 6a and D) show the presence of residual carboxylate and copper oxide (bands below 500 cm⁻¹), with a high Cu/propionate ratio (combining EA and TG results: Cu:O=4:CuH₄), and a lower carbon content (Table 1) than the theoretical value for copper carbonates (6–10%) [46]. This harder-to-decompose intermediate may suggest the formation of CuO clusters stabilized by the remaining propionates, similarly to the way nanoparticles have been reported to form from solvent-less routes [47], although usually synthesized through wet chemistry [48,49]. This intermediate later decomposes producing mostly acetaldehyde (from oxidation of CH₃H₅ fragments of the ligand) and CO₂.

This second decomposition stage is accelerated by oxygen and it is not affected by humidity (see Fig. 2); in oxygen, decomposition occurs at 220 °C (solid lines in Fig. 2, A and C), whereas without oxygen, B and D, it is delayed to 240 and 260 °C, respectively (dashed lines in Fig. 2; this stage occurs at even higher temperatures in more inert atmosphere, Fig. S9). Apparently, stage II involves oxidation because the mass-loss step is accompanied by an exothermic DSC peak (Fig. 2b). This is in agreement with reaction scheme β (Table 2). Except for the experiment in dry N₂ (D), the sample mass is quite stable beyond 260 °C. Further-more, in all films at 500 °C, XRD analysis reveals only the presence of CuO (Fig. 5a, A), and EA (Table 1) confirms that neither carbon nor
hydrogen is left.

Only in the case of dry N₂, this second step coincides with Cu(II)/O reduction to Cu(Ii)/O (Fig. 5b, D) upon CO₂ evolution (Fig. 4, D) at 260 °C. The fact that Cu(Ii) is obtained out of the Cu(II)/Cu(I) thermodynamic equilibrium conditions suggests a redox reaction between the remaining carbon (4% in Table 1) and CuO. In fact, a sample heated in humid O₂ and quenched before the II stage begins, experiences CuO reduction (like in D) when heated in vacuum, releasing CO₂ and CO (shown in Supp. Info). However, by 350 °C, given the high sensibility of films to residual oxygen, the CuO film is oxidized again to CuO (Fig. 5a, D), and at 500 °C no carbon and hydrogen are left (Table 1). Finally, note that the presence of water vapor (B, humid N₂) is enough to prevent this reduction.

To further confirm these decomposition reactions and exclude any possible influence of remaining acetate ligands, the decomposition of a powder obtained from an acetate-free precursor solution (CuCO₂ in excess propionic acid) was studied as a comparison, spread on a LAO substrate to approach the conditions of experiment A. The decom-position positions were very similar and the volatiles were the same as in exp. A, confirming that the reaction schemes proposed are the expected ones for the decomposition of CuProp₂ (see Fig. S3 in Supp. Info).

### 3.3. Thermal decomposition of CuProp₂ at atmospheric pressure in powder

For “bulk” CuProp powder (m of tens of mg), the residual water vapor in nominally dry O₂ and N₂ (D') atmospheres has no effect on decomposition and, consequently, powders decompose at much higher temperatures (> 250 °C, Fig. 2a) than films.

In dry N₂ (D'), decomposition is endothermic (Fig. 2b), which is consistent with the reduction of copper (like in process y of Table 2). XRD and FTIR measurements (see Figs. 5b and 6 D'), respectively, on samples quenched at 250–265 °C, around the mass derivative discontinuity, indeed reveal the formation of CuO; just after quenching from 265 °C, a fraction of the sample is liquid, and green portions emerge as it quickly solidifies, resulting in an XRD peak at 2θ = 6.5 (* in Fig. 5b, D', 260 °C) and new C=O FTIR bands (* in Fig. 6 D') at 1740 cm⁻¹. In fact, the sample mass loss at this temperature (65% m/m) is in agreement with Cu(I)/Prop formation, but the XRD peak at low angles might suggest its oxidation to a basic copper propionate [50].
Fig. 6. Infrared evolution of the solid residue during decomposition of CuProp$_2$ as film and powder in different atmospheres. Details of the COO$^-$ stretching region can be found in the Supp. Info. (•) CuProp$_2$; (△) unknown phase. The vertical lines are drawn to show that $\nu_{as}$ gets closer to $\nu_{s}$ as pyrolysis proceeds.

during cooling, after coming into contact with atmospheric air. The volatiles consist of CO$_2$, acetaldehyde and propionic acid. Finally, the product at 500 °C consists of Cu and Cu$_2$O (Fig. 5a), and a significant 3% of carbon residues (Table 1, D'). This is probably due to the very inert local atmosphere when compared to the film (experiment D). This was also observed for other copper compounds (with long alkyl chains) in N$_2$ at 500 °C [51].

In dry O$_2$ (C'), the reactive atmosphere displaces decomposition to lower temperature (Fig. 2a) but it does not impede Cu reduction (Supp. Info, Fig. S8). In fact, the final abrupt mass-loss step observed at 250 °C in the TG curve (and sharp exothermic peak in the DSC curve) is a clear signature of combustion, and in such a fast reaction mode complete oxidation is usually not possible [52]. After combustion, the sample mass reaches a minimum below the expected mass for the formation of CuO, but the subsequent mass increase (and the final black color) shows that it is then oxidized again to the expected CuO final product.

3.4. Thermal decomposition of CuProp$_2$ in film and powder at low pressure

At low pressure (experiment E), volatilization of the film occurs: the substrate is left with a blackish color, consistent with both copper oxide residues and elemental carbon, but the mass loss is close to 99%, and XRD analysis could not detect anything. A blue compound is found on the cold parts of the quartz tube which by FTIR and XRD analysis (Fig. S6) is revealed to be similar to Cu(II)Prop$_2$. Volatilization of Cu(II)Prop$_2$ occurs simultaneously with decomposition, since EGA-MS (Fig. 7, E)

![Figures](image1.png)
detects a small amount of volatiles, mainly related to propionic acid ($m/z = 74, 45, 29, 28, 27$). Volatilization of copper propionate was already observed by [34] above 167 °C in vacuum, and it is consistent with a series of findings, for example that volatile binuclear Copper(I) carboxylates are among the decomposition products of Cu(II) carboxylates [53], and with the general strong tendency of copper complexes of even longer chains to sublime at low pressure [54,55].

On the other hand, when the powder is heated in vacuum to 500 °C ($E'$, $E''$), at 10–25 °C/min instead of 5 °C/min, the sample decomposes between 150 and 300 °C, yielding Cu(0). However, the crucible is almost empty, while the sample chamber has turned pink.

Unlike from the gas species are detected a small amount of volatiles, mainly related to propionic acid ($m/z = 29,28,27$), which could be in agreement with propionic acid ($\text{COOH}$), which could be in agreement with propionic acid ($m/z = 29,28,27$), acet-aldehyde ($m/z = 29,44$) and $\text{CO}_2$ found by TG-MS-FTIR experiments (Fig. 4, $D''$). In $E'$, the first peak at 240 °C follows scheme $\alpha$ and is due to residual water, as the water signal points inward ($m/z = 18$). The $\text{CO}_2$ peak at 260 °C is due to copper oxide's reduction to Cu(0) by elemental carbon, just like in $D$. Increasing mass and heating rate in $E''$, decom-position proceeds according to $\gamma$ as in $D''$. Thus, $E'E''$ can be the con-sequence of pushing experiment $D'D''$ to very low $O_2$ partial pressures, where only metallic copper is formed.

3.5. Thermochemistry Calculations

As a further confirmation of the reactions proposed in Table 2, the experimental enthalpies of reaction for film and powder decomposition obtained from DSC (Fig. 2b) were compared with the theoretical values (Supp. Info) of electronic energy for the same reactions.

Hydrolysis ($\alpha$) is predicted to be slightly endothermic, as observed for stage I in all experiments in film (A, B, C, D). This assignment is especially significant for the experiments in nominally dry atmosphere (C - O$_2$ and D - N$_2$), because it leads to the conclusion that residual humidity is enough to sustain this reaction.

Oxidation of the organic ligand ($\beta$) is exothermic and can occur at different degrees; thus, stage II in films in O$_2$ (A - humid, C - dry) is attributed to oxidation to acetaldehyde (entry iii and iv, in agreement with the detected volatiles) and its exothermicity decreases from C to A to B (humid N$_2$) as the degree of oxidation goes from iv, to iii to ii. D (dry N$_2$) represents an intermediate situation between films in O$_2$ and powders in N$_2$, since only partial reduction to Cu(0) occurs as the residual propionate decomposes in Stage II ($\gamma$). The latter (slightly endothermic) triggers the reduction of the CuO formed from stage I at the expenses of a stronger oxidation of the remaining organic ligand (probably to $\text{CO}_2$ and H$_2$O, see Fig. S7) according to $\text{vi-x}$ (slightly exothermic), and combined they could explain why stage II is not accompanied by a relevant DSC peak (Fig. 2b).

Concerning powder in N$_2$ ($D''$), the predicted slight endothermic character of the reduction reaction ($\text{vi of } \gamma$) is confirmed by experiment. In dry O$_2$ ($C''$), the heat evolved is lower than in films (A, C) indicating a lower degree of ligand oxidation ($\beta$) due to local $O_2$ depletion as reaction gases evolve [52], and as $\gamma$ compete with $\beta$. This fact does not impede combustion occurring at the end of decomposition because, in contrast with the good thermal contact with the substrate for films, heat dissipation is much less efficient for powder [52].

4. Discussion

CuProp$_2$ decomposition kinetics depends on the geometry (film or powder) and the furnace atmosphere. A humid atmosphere triggers decomposition at lower temperature (Fig. 2) in films, promoting the hydrolysis ($\alpha$) of the salt, and this shift is helped by reduced thickness. Films are so sensitive to humidity that the residual water vapor in nominally dry conditions (C and D) has a similar effect. Conversely, sensitivity to residual humidity or $O_2$ is lost for powders, which decompose at a much higher temperature. In fact, in bulk form, attention has to be paid to the local atmosphere created at the interparticle voids during decomposition [9,56,57]. This phenomenon results in inert or even reducing local conditions leading to similar behavior for powders in N$_2$ and in O$_2$: Cu(II) reduction (path $\gamma$) even in nominally pure O$_2$ ($C''$), and to a higher difficulty for carbon removal (Supp.Info, Fig. S8). Anyway, the nominal atmosphere has an effect on the decomposition process: combustion occurs in pure O$_2$. The difference in behavior between films and powders found for the particular case of CuProp$_2$, agrees with that of other metal organic salts [56].

Changing atmosphere affects the final product (Cu versus CuO) rather than the nature of the volatiles coming from decomposition (as in the case of YProp$_3$ [9]), which in both inert and oxidizing atmosphere consist of oxidized forms of the ligand. While in O$_2$ this is clearly understood from the high availability of oxygen, in inert atmosphere it can be explained by the oxidation of gaseous products contributing to the metal center reduction from Cu(II) to Cu(I) and Cu(0) according to reaction scheme $\psi$. In fact, step II in D is more delayed than $D$ due to decreasing oxygen availability (as $H_2O$ vapor), but oxidation of residual ligand to $\text{CO}_2$ can still be pushed by the CuO/CuO$_2$ reduction.

This is in agreement with the hypothesis that [27], in inert conditions, metal ions that easily undergo redox reactions, like Cu, Ag and Hg, promote the formation of the corresponding carboxylate acid, while the evolution of ketones is favored by the opposite class of metals (i.e. Y, Ca.,) emphasizing the role of the metal ion in affecting the salt decomposition. This would explain why metal carboxylates that would normally release ketones, yield an acid in the presence of iron or copper salts [23,58].

No alkene was found, unlike for the decomposition of even chain CuHg carboxylate [24,25], suggesting that the chain length may also play a role in the decomposition pathway or that the decomposition mechanism is a result of the general catalytic activity of copper, resulting in the oxidation of ethylene to acetaldehyde [59].

Lastly, reducing the pressure from atmospheric to $10^{-6}$ bar causes volatilization of the salt to occur, but at different stages of decom-position, according to the geometry of the sample and heating rate: the higher the surface to volume ratio (films, $E$), the faster the volatilization with respect to decomposition; conversely, at higher heating rates and thicker samples ($E''$), decomposition is more significant than volatilization. On the other hand, at atmospheric pressure, as the mass loss is never below than that of the expected compounds, any significant volatilization of copper species can be ruled out, until tested thicknesses of 300 nm. Additionally, the presence of water vapor is expected to pre-vent volatilization by shifting decomposition to low temperatures thanks to scheme $\alpha$. Indeed, the hydrolysis-driven decomposition of the carboxylate salts, already also found for the YProp$_3$ case [9], can ex-plain the advantage of using humid atmospheres in the framework of CSD-YBCO film pyrolysis; in fact, humidity is critical to reduce the decomposition temperatures of the thermal treatment.

5. Conclusions

In conclusion, the thermal decomposition of CuProp$_2$ was studied for samples in the form of powder and as film. By IR and MS detection of the volatiles, we were able to identify almost univocally the species evolved during decomposition (dominated by propionic acid), clarifying the mechanism with respect to the literature. While, as a consequence of the chemical nature of the metal center, the nature of the volatiles does not vary in response to the atmosphere change or in relation to the surface to volume ratio of the sample, the CuO-Cu redox behavior is found to be affected by it. Specifically, it is easier to obtain CuO and Cu from powders than from films. Additionally, the comparison of experimental and computational thermochemistry data helped clarify and support the decomposition reactions written based on the observed volatiles. Finally, the role of water vapor in the pyrolysis of YBCO films has been cleared. Humidity promotes decomposition at low temperatures, where volatilization would otherwise
prevail in a low-pressure environment. Therefore, the study of films’ thermal behavior is not only crucial for identifying the intrinsic thermal mechanisms that could pass unnoticed while studying powders, but also for the analysis of the real phenomena occurring during tape synthesis of high temperature superconductors and other functional oxide films.

Acknowledgments

This work was supported by Ministerio de Ciencia, Innovación y Universidades (grant numbers RTI2018-095853-B-C21 and RTI2018-095853-B-C22), by the Center of Excellence Severo Ochoa (SEV-2015-0496), the Generalitat of Catalonia (SGR753 and SGR948) and by the Universitat de Girona (UdG, contract number MPCUdG2016/059). SR and FS thanks the Universitat de Girona (UdG) and the Cardiff University School of Chemistry, respectively, for their fully-funded PhD scholarships. The authors also wish to thank the Scientific services of the UdG and of the Institute of Materials Science of Barcelona (ICMAB), Computing facilities were provided by ARCCA at Cardiff University, HPC Wales, and via the membership of the UK’s Materials Chemistry Consortium (MCC), which is funded by EPSRC (EP/F067496).

Appendix A. Supplementary data

Supplementary material related to this article can be found in the online version, at doi: https://doi.org/10.1016/j.jaap.2019.04.008.

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