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Insights on the Stability and Cationic Nonstoichiometry of CuFeO$_2$ Delafossite

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Supporting Information

ABSTRACT: CuFeO$_2$, the structure prototype of the delafossite family, has received renewed interest in recent years. Thermodynamic modeling and several experimental Cu$-$Fe$-$O system investigations did not focus specifically on the possible nonstoichiometry of this compound, which is, nevertheless, a very important optimization factor for its physicochemical properties. In this work, through a complete set of analytical and thermostructural techniques from 50 to 1100 °C, a fine reinvestigation of some specific regions of the Cu$-$Fe$-$O phase diagram under air was carried out to clarify discrepancies concerning the delafossite CuFeO$_2$ stability region as well as the eutectic composition and temperature for the reaction L = CuFeO$_2$ + Cu$_2$O. Differential thermal analysis and Tammann’s triangle method were used to measure the liquidus temperature at 1050 ± 2 °C with a eutectic composition at Fe/(Cu + Fe) = 0.105 mol %. The quantification of all of the present phases during heating and cooling using Rietveld refinement of the high-temperature X-ray diffraction patterns coupled with thermogravimetric and differential thermal analyses revealed the mechanism of formation of delafossite CuFeO$_2$ from stable CuO and spinel phases at 1022 ± 2 °C and its incongruent decomposition into liquid and spinel phases at 1070 ± 2 °C. For the first time, a cationic off-stoichiometry of cuprous ferrite CuFe$_{1-y}$O$_2$ was unambiguous, as evidenced by two independent sets of experiments: (1) Electron probe microanalysis evidenced homogeneous micronic CuFe$_{1-y}$O$_2$−δ areas with a maximum y value of 0.12 [i.e., Fe/(Cu + Fe) = 0.47] on Cu/Fe gradient generated by diffusion from a perfect spark plasma sintering pristine interface. Micro-Raman provided structural proof of the existence of the delafossite structure in these areas. (2) Standard Cu additions from the stoichiometric compound CuFeO$_2$ coupled with high-temperature X-ray diffraction corroborated the possibility of obtaining a pure Cu-excess delafossite phase with y = 0.12. No evidence of an Fe-rich delafossite was found, and complementary analysis under a neutral atmosphere shows narrow lattice parameter variation with an increase of Cu in the delafossite structure. The consistent new data set is summarized in an updated experimental Cu$-$Fe$-$O phase diagram. These results provide an improved understanding of the stability region and possible nonstoichiometry value of the CuFe$_{1-y}$O$_2$−δ delafossite in the Cu$-$Fe$-$O phase diagram, enabling its optimization for specific applications.

1. INTRODUCTION

The present work focuses on the description of equilibrium phases of the Cu$-$Fe$-$O system, which has, for instance, a high technological interest in the copper (Cu) industry because iron (Fe) is a major contaminant of Cu scrap and ore. In air, this system contains two mixed oxide phases, namely, cuprospinel and delafossite. A large amount of research effort during the last decade has focused on the cuprospinel phase Cu$_2$Fe$_{y-2}$O$_4$, a well-known inverse spinel structure with cubic (space group Fd$\bar{3}$m) or tetragonal (space group I$4$_1/amd) symmetry depending on the temperature and consisting of a solid solution between copper ferrite (z = 1, i.e., CuFe$_2$O$_4$) and magnetite (z = 0, i.e., Fe$_3$O$_4$). Fe-based spinels have been widely described in the literature because of their magnetic, electrical, and optical properties, with applications in catalysis among others.
Conversely, the CuFeO$_2$ phase, which is the structure prototype of the delafossite family (first described by Friedel$^{12}$ in 1873 and named in honor of the French mineralogist and crystallographer Delafosse), did not attract much attention until the discovery of the properties of simultaneous transparency and p-type conductivity in this delafossite crystallography$^{6}$ as well as its photovoltaic$^{20}$ and/or thermoelectric$^{21}$ properties. This highly anisotropic phase crystallizes with the rhombohedral (space group R$\overline{3}$m) or hexagonal (space group P6$_3$/mmc) polytype. It is composed of compact double layers of O atoms, with octahedral sites occupied by Fe$^{3+}$ ions, and the cohesion between layers is ensured by Cu$^+$ ions in linear coordination.

1.1. Cu–Fe–O Phase Diagram in Air. Experimental investigations of the phase equilibria in the Cu–Fe–O system have been carried out for decades and led to the proposal of several phase diagrams$^{1,24–30}$ Figure 1 (dashed blue lines) presents the phase diagram assessed by Perrot in 2007$^{31}$ based on an exhaustive literature analysis. Recently, two thermodynamic models of this system have been established with the CALPHAD (CALculations of PHase Diagrams) method, which is based on an assessment of the Gibbs energy functions for all phases of the system and allows the computation of phase diagrams. Khvan et al.$^{32}$ assessed the Cu–Fe–O system in 2011, and the model covers a temperature range between 650 and 1000 °C (not considered in this work). Shishin et al.$^{33}$ published another model in 2013 (solid black lines, Figure 1), which covers the high-temperature equilibria.

One can note that several areas of the diagram are consistent according to the different studies. First, the transition temperatures in the binary subsystems Cu–O and Fe–O are described similarly. Second, the stability region of the cuprospinel phase selected by Perrot et al. and based on the experimental results of Mexmain$^{34}$ and Kenfack and Langbein$^{35}$ is well represented by the thermodynamic model of Shishin et al.$^{33}$ Third, delafossite is always considered to be a stoichiometric compound CuFeO$_2$. However, there are also noticeable differences between the two diagrams plotted in Figure 1. The first is the stability range of the delafossite compound. Indeed, Perrot et al. selected a formation temperature of 1015 ± 2 °C and a decomposition temperature of 1090 ± 5 °C, based on the measurements done by Yamaguchi$^{36}$ in 1966. The same transition temperatures

Table 1. Published Values for the Lattice Parameters of Some Delafossite CuFeO$_2$ at Room Temperature

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<thead>
<tr>
<th>lattice constant</th>
<th>form</th>
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<th>ref</th>
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<td>(c) (Å)</td>
<td>(c/a)</td>
<td>(V) (Å$^3$)</td>
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<td>17.156</td>
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<td>5.66</td>
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<td>5.66</td>
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</tr>
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<td>17.16(2)</td>
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<td>17.14</td>
<td>5.66</td>
<td>136.28</td>
<td>powder</td>
</tr>
</tbody>
</table>

![Figure 1. Phase diagrams of the Cu–Fe–O system in air, according to Perrot et al.$^{31}$ (dashed blue lines, drawn from the literature review) and Shishin et al.$^{33}$ (solid black lines, calculated from the thermodynamic model).](image-url)
calculated by the model of Shishin et al.\textsuperscript{33} are 1019 and 1077 °C, respectively. The second flagrant inconsistency for this system is the composition and temperature of the eutectic L = CuFeO\textsubscript{2} + Cu\textsubscript{2}O in air, highlighted by a star symbol in Figure 1. Yamaguchi\textsuperscript{25} suggested a eutectic temperature at 1080 °C, while the one announced by Buist et al.\textsuperscript{25} is lower than 1036 °C. The eutectic temperature obtained from the thermodynamic model is 1049 °C. Similarly, the composition of the eutectic is not unanimously defined and ranges from 0.13 to 0.33 (based on the Fe molar ratio). It is particularly difficult to select one or the other coordinates because of their limited number and the variability of available experimental data.

1.2. Delafossite CuFe\textsubscript{O}\textsubscript{2} Stoichiometry. The exact composition of the delafossite compound has been controversial for a long time. In the early 1900s, Rogers\textsuperscript{36} determined the formula CuFeO\textsubscript{2} (Cu\textsubscript{2}O:Fe\textsubscript{2}O\textsubscript{3}) after leaching in sulfuric acid and chemical analysis. In the 1960s, Buist et al.\textsuperscript{24} proposed a new formula (Cu\textsubscript{6}Fe\textsubscript{3}O\textsubscript{7}), deduced from weight loss on thermal analysis. Three years later, a new investigation was performed by Wiedersich et al.\textsuperscript{37} coupling X-ray diffraction (XRD), Mössbauer spectroscopy, and chemical analytical techniques. The Cu\textsubscript{3}Fe\textsubscript{2}O\textsubscript{7} (3Cu\textsubscript{2}O:Fe\textsubscript{2}O\textsubscript{3}) compound could not be stabilized and was thus ruled out. CuFeO\textsubscript{2} was finally established as the only composition for delafossite. Hey\textsuperscript{38} confirmed this result using natural delafossite, and Schaefer et al.\textsuperscript{26} provided more data under low O\textsubscript{2} pressure (1 × 10\textsuperscript{−4}−5 × 10\textsuperscript{−4} atm) in the 900−977 °C temperature range. Because of the lack of accuracy of the analytical devices available for these early studies, small deviations in the cationic stoichiometry of the delafossite phase were not truly investigated.

In the past decade, small deviations of the cationic stoichiometry have been evidenced in a few compounds of the delafossite family, such as Cu\textsubscript{i−δ}Ga\textsubscript{O} (−0.01 ≤ y ≤ 0.02),\textsuperscript{39} Cu\textsubscript{i+δ}Mn\textsubscript{i−δ}O\textsubscript{2} (0 ≤ y ≤ 0.2),\textsuperscript{40} and CuCr\textsubscript{i−δ}O\textsubscript{2} (0 ≤ y ≤ 0.1).\textsuperscript{41} To a larger extent, this can even considerably improve the properties of certain materials such as, for example, the p-type transparent conducting Cu\textsubscript{i−δ}Cr\textsubscript{i+δ}O\textsubscript{3} thin film.\textsuperscript{42} Significant quantities of O (δ) can also be intercalated into the Cu planes of the CuMO\textsubscript{2} delafossite structure according to CuMO\textsubscript{2} δ.\textsuperscript{43} This requires a cation M\textsuperscript{3+} exceeding a certain size and leads to an off-stoichiometric value δ, which can reach 0.66 when M\textsuperscript{3+} = La\textsuperscript{3+}, corresponding to a Cu\textsuperscript{2+} /Cu\textsuperscript{3+} (and even Cu\textsuperscript{4+} /Cu\textsuperscript{3+}) mixed valence.\textsuperscript{44} To date, because of the relatively small size of the Fe\textsuperscript{3+} cation, only limited O nonstoichiometry (0.08 ≤ δ ≤ 0.18) has been reported on CuFeO\textsubscript{2−δ}\textsubscript{δ} delafossite-type powders\textsuperscript{34,45} or single crystals.\textsuperscript{46,47} No work has so far provided real evidence of the existence of a cationic nonstoichiometry in CuFeO\textsubscript{2}, but some indications available in the literature, however, suggest this eventuality. Indeed, in a recent work, Wuttig et al.\textsuperscript{14} reported Cu nonstoichiometry in Cu\textsubscript{i−δ}Fe\textsubscript{O}\textsubscript{2}, with y values as low as 0.005 and 0.02, but they did not provide elemental or structural proof of the existence of the delafossite structure. Furthermore, in a recent chemical measurement of an inclusion phase in a Cu matrix,\textsuperscript{38} the chemical formula CuFe\textsubscript{0.8}O\textsubscript{1.79} was proposed. Once again, no structural analysis was provided to confirm the existence of this Cu-rich delafossite.

To complete this review on the CuFeO\textsubscript{2} delafossite phase stoichiometry, an exhaustive list of the available crystallographic data is reported in Table 1 for powders and single-crystal samples. Thin-film materials were intentionally excluded because of the potential stress and strain induced by the growing method and substrate nature. Despite the various synthesis conditions, the accuracy of the measurement techniques, and the possible O off-stoichiometry, remarkably grouped values are obtained for these lattice parameters. Whatever the single-crystal or powder samples, the standard deviation of the lattice constants is very narrow, i.e., 1.3% along the [100] direction, 3.5% along [001], and 0.6% in volume. Because there is no significant difference between all of the reported lattice constants, it is hard to suspect a cational nonstoichiometry, and up to now, the cationic molar ratio Fe/(Cu + Fe) has always been considered to be equal to 0.5 in the delafossite phase.

1.3. Research Questions. According to our bibliographic review, it appears that several important elements are missing to describe accurately the Cu−Fe−O system in air. The first concern is the stability region and phase boundaries of the delafossite phase, which are substantially different in the various available phase diagrams and models and need to be revisited. The second is related to the eutectic composition and temperature for the equilibrium L = CuFeO\textsubscript{2} + Cu\textsubscript{2}O. Finally, the possibility of a degree of cationic nonstoichiometry in delafossite needs to be closely investigated.

In the present work, we address these open questions through the implementation of an experimental study combining in situ high-temperature XRD (HT-XRD), thermal analysis, and accurate compositional analysis on various oxide mixtures. Precise monitoring of the delafossite composition range was also realized through a diffusion couple between copper and iron oxides.

Such a better description of the Cu−Fe−O phase diagram is crucial because it will permit one not only both to minimize the experimental effort and to improve the general understanding of the processes for producing these phases with the delafossite structure,\textsuperscript{60,61,67−69} but also to optimize the remarkable properties of CuFeO\textsubscript{2} by a better understanding of the complex relationships between the structure and properties.\textsuperscript{15−17,70}

One can note that, because some major applications of CuFeO\textsubscript{2} materials concern the surface properties, it also should be interesting to consider similar analysis on film. Even if thin-film materials could be characterized by an ad-hoc method, the equilibrium conditions are generally not satisfied because of, for instance, an almost instantaneous transition of the elements of the very high-temperature plasma to a crystalline solid phase at low temperature in the physical vapor deposition process. It is, however, possible to establish experimental “metastable” phase diagrams, representing the nature of the phases formed depending on the composition of the system and the deposition conditions of the thin films.\textsuperscript{71} In recent work, Schneider et al.\textsuperscript{72} developed a predictive approach to calculating metastable phase diagrams according to the power applied, based on the determination, by ab initio calculations, of the surface diffusion properties of the element constituents of the system. These calculations strongly rely on the most accurate description of the system at equilibrium, such as that presented in the present work for the Cu−Fe−O phase diagram.

2. EXPERIMENTAL SECTION

2.1. Materials. Oxide Powder Precursors. Two different batches of CuO−Fe\textsubscript{2}O\textsubscript{3} powders were used: (i) CuO (Acros Organics; >99%), α-Fe\textsubscript{2}O\textsubscript{3} (Alfa Aesar; 99.5%), and Cu\textsubscript{2}O (Alfa Aesar; 99%) for
Table 2. Samples Compositions Measured by Weighing, XRF, and ICP-AES

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<th>XRF</th>
<th>ICP-AES</th>
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<td>0.00</td>
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<td>0.25</td>
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<td></td>
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Table 3. Rietveld Refinement Correction Parameters

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<th>ref</th>
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“Mass absorption coefficients were determined at 8.05 keV [I(γ)/Kα] using physics.nist.gov/PhysRefData/FFast/html/form.html.”

XRD investigations; (ii) CuO (Alfa Aesar; 99.998%) and α-Fe2O3 (Alfa Aesar, 99.995%) for thermal analysis. All samples were prepared by weighing the starting binary oxides in the desired proportions and mixing them in an agate mortar until a homogeneous mixture was obtained. The nominal composition of five mixed samples is presented in Table 2. The cationic molar ratio of the samples was controlled by X-ray fluorescence (XRF; SR 2 Ranger, Bruker) and inductively coupled plasma-atomic emission spectroscopy (ICP-AES; Ultima 2, Jobin Yvon) after dissolution in aqua regia. The three techniques provide very similar compositional values for all samples in the limit of their accuracies. In the following, all samples are identified by their nominal compositions in the scale of molar fractions x = Fe/(Cu + Fe), with the ratio obtained from the weighing data, which are accurate enough to evaluate this ratio x with two decimals.

Delafossite CuFe2O4 Powder. A polycrystalline sample of CuFe2O4 was prepared by a standard solid-state reaction according to a process described in a previous work:21 CuO and Fe2O3 powders of batch (i) were mixed in stoichiometric quantities and then treated in a nitrogen atmosphere between 900 and 1000 °C for 30 h with intermittent grinding.

Diffusion Couple. A spark plasma sintering (Fuji 632 Lx) device was used to prepare a bilayer sample, where each layer corresponds to one of the binary oxides of batch (i). The following conditions were applied: pressure of 50 MPa; C mold under an argon atmosphere; heating rate of 100 °C/min, dwell time of 7 min at 750 °C; cooling rate of 150 °C/min. After a dense bilayer sample (96% of theoretical maximum value) was obtained, the bulk was cut and annealed in a muffle furnace (Nabertherm) under an air atmosphere at 1045 °C for 30 min (heating rate of 5 °C/min; cooling rate of 10 °C/min).

2.2. Characterization Methods. Composition Analysis. The stoichiometries of copper–iron mixed oxides formed in the diffusion couple experiment were measured by field-emission-gun electron probe microanalysis (EPMA; SX Five FE, CAMECA).

Microstructural Analysis. The distribution sizes of the CuO and Fe2O3 precursors were determined using a laser diffraction particle size analyzer (Malvern Instruments, Mastersizer 3000) with an Aero S dry powder dispersion attachment. Control of the diffusion couple interface was carried out by field-emission-gun scanning electron microscopy (SEM) using a Joel JSM 7800F microscope.

Thermal Analysis. Thermogravimetric analysis (TGA) and differential thermal analysis (DTA) were carried out using a Setaram TGA/DTA 92 instrument in flowing air, in order to measure the transition temperatures. Samples of around 20 mg placed in alumina crucibles were heated from 25 to 1100 °C, at heating and cooling rates of 5 °C/min. Different rates of 1 and 10 °C/min to heating and cooling, respectively, were also applied to resolve some close transitions. Gold (Tmelting = 1064.18 °C) was used as the reference material to calibrate the setup around the working temperature. The resulting accuracy for temperatures is ±2 °C.

Structural Analysis. In order to perform local structural characterization at room temperature in the diffusion couple samples, micro-Raman spectroscopy was carried out using a Jobin Yvon-Horiba LabRam infinity spectrometer (from 100 to 1000 cm⁻¹ with a 532 nm laser). For global structural analysis, HT-XRD measurements were performed using a Bruker D8 diffractometer [λ(Cu Kα1) = 1.54056 Å and λ(Cu Kα2) = 1.54443 Å radiation; nickel filter] equipped with an Anton Parr HTK1200N high-temperature chamber under an air, a nitrogen, or a primary vacuum atmosphere. A Bruker LynxEye detector was used in a 1D mode for rapid acquisition. The HT-XRD patterns were collected from 50 to 1100 °C, under isothermal conditions in steps of 50 °C, and with a heating rate of 30 °C/min between dwells. The acquisition conditions were 2θ within 15–75°, with a step size of 0.02° and a counting time of 1.11 s. Crucibles in alumina were used. Some tests have also been done in a ZrO2 crucible to confirm that no problem with aluminum diffusion (which can form the parent compound CuAl2O4 in a Cu-rich region) occurs with the Al2O3 crucibles. A second set of measurements was made from 975 to 1045 °C, under isothermal conditions in steps of 5 °C and a heating rate of 30 °C/min between dwells. The conditions were 2θ within 17–69°, with a step size of 0.02° and a counting time of 1.85 s. The total acquisition duration was 80 min per diffraction pattern.

Phase identification was performed with the DiffraPlus EVA software combined with the PDF database and subsequently refined by the Rietveld method using the Full Prof-Win Plot program.22,23 The Rietveld refinement method was used to determine the lattice constants as well as quantify the amount of crystalline phases at each temperature. The lattice constant standard deviations were systematically balanced by the quality of the refinement. In order to obtain an accurate quantification whatever the mass fraction of the constituent phases, i.e., to be in agreement with the pristine values reported in Table 2, intensity corrections had to be applied in the Rietveld refinement. First, an internal standard correction was applied using the reference intensity ratio (RIR) from the sample to corundum (I/I0) obtained in the PDF database. Second, differential microabsorption between phases was taken into account by applying the Brindley correction method.22,23 The Brindley intensity correction factor τm of phase i in mixture m is defined by eq 1:

\[
\tau_m = \frac{1}{V_i} \int_0^V e^{-n_i(n - n_m)} \, dV_i
\]

where V, n, and n are the volume, linear absorption coefficient, and density of phase i, respectively. The average linear absorption coefficient of the mixture μm is defined by eq 2:

\[
\mu_m = \rho_m \sum_i \frac{\mu_i}{\rho_i}
\]

where ρm is the density of the mixture m and wi is the mass fraction of phase i.

The expression of τm was approximated using the average radius (R) of the spherical particles composing phase i according to the following equation:

\[
\tau_m = 1 - 1.450(\mu_m - \mu_\text{ref})R_i + 1.426(\mu_m - \mu_\text{ref})R_i^2
\]
3. RESULTS AND DISCUSSION

3.1. Formation Temperature of the Delafossite Phase. The solid reactions between CuO (tenorite, monoclinic structure, and space group C2/c) and α-Fe₂O₃ (hematite, trigonal structure, and space group R3c) were evaluated via in situ HT-XRD and Rietveld refinement for all of the samples listed in Table 2. The corresponding HT-XRD patterns are shown in Figure 2a except for x = 1.00 because α-Fe₂O₃ is stable in the whole range of temperature. The temperature evolution of the amount (in mole percent) of each crystalline phase obtained from Rietveld analysis is plotted in Figure 2b.

For the x = 0.00 sample (pure CuO), a change in the broadening of the tenorite Bragg peaks is visible from 600 °C because of the increase of the particle size (from 45 to >100 nm estimated by Scherrer’s law on the one side and laser diffraction particle size analysis on the other side). The complete reduction of tenorite CuO into cuprite Cu₂O occurs between 1000 and 1050 °C according to eq 4:

\[ 2\text{CuO(s)} \rightarrow \text{Cu}_2\text{O(s)} + \frac{1}{2}\text{O}_2(\text{g}) \]  

(4)

For all of the samples in the 0.25 ≤ x ≤ 0.75 composition range, the single oxide precursors CuO and Fe₂O₃ are stable up to 800 °C. From 850 to 1000 °C, the precursors react and...
form a spinel phase (green square symbols in Figure 2b),
according to eq 5:

$$\text{CuO(s)} + \text{Fe}_2\text{O}_3(s) \rightarrow \text{CuFe}_2\text{O}_4(s)$$

The progressive increase of the amount of the spinel phase
from 850 to 1000 °C, which is thermodynamically stable at
lower temperature according to the phase diagram (Figure 1),
indicates a thermally activated diffusion process. For the
samples $x = 0.25, 0.40, 0.50,$ and $0.61$, the refined lattice
parameter of the spinel phase is constant whatever the Cu
content such as, for instance, $a = 8.499(2)$ Å at 950 °C. This
corresponds to formation of the stoichiometric CuFe$_2$O$_4$
cuprospinel phase, in accordance with the phase diagram.

The delafossite phase CuFeO$_2$ forms between 1000 and
1050 °C for $0.25 \leq x \leq 0.61$. The proposed reaction is the
concomitant reduction of the Cu$^{2+}$ cations present in the
cuprite and spinel phases, according to eq 6. This is confirmed
by the significant decrease of the amount of spinel and cuprite
phases at this temperature as observed, for instance, for $x = 0.40$.

$$\text{CuFe}_2\text{O}_4(s) + \text{CuO}(s) \rightarrow 2\text{CuFeO}_2(s) + \frac{1}{2}\text{O}_2(g)$$

Because the diffusion process is very active at this
temperature, formation of the delafossite phase could also be
due to the simultaneous reduction of Cu$^{2+}$ from CuO to Cu$_2$O
(eq 4) and its diffusion within Fe$_2$O$_3$ according eq 7.

$$\text{Cu}_2\text{O}(s) + \text{Fe}_2\text{O}_3(s) \rightarrow 2\text{CuFeO}_2(s)$$

Because no trace of Cu$_2$O was detected by HT-XRD in the
whole series of samples (except for $x = 0.00$), it can be
deduced that the reaction scheme that leads to the formation
deafosite is eqs 5 and 6 rather than eqs 4 and 7.

Finally, at 1100 °C, the delafossite phase fully decomposes.
In the samples $x = 0.25, 0.40, 0.50,$ and $0.61$, the presence of a
liquid phase was observed on top of a spinel phase, as reported
in Figure 2b. Even if HT-XRD cannot directly evidence and
Figure 4. Mass variation (dotted lines) and heat flow (solid lines) over the temperature range 1000–1100 °C: (a) $x = 0.03, 0.07, 0.128,$ and $0.145$ in air, with a heating rate of 1 °C/min; (b) $x = 0.03, 0.07, 0.128,$ and $0.145$ in air, with a cooling rate of 10 °C/min. The integral of the peaks associated with eutectic solidification is the red area with a sigmoidal baseline.

Figure 5. Summary of the experimental information obtained by TGA/DTA: temperature of formation to CuFeO$_2$ at $1022 \pm 2$ °C; transformation of CuO into Cu$_2$O at $1031 \pm 2$ °C; liquidus formation at $1050 \pm 2$ °C; solidus thermal effect of the eutectic formation during cooling (Tammann’s triangle).
Table 4. Relationship between the Structural Parameters of Delafossite and Composition

<table>
<thead>
<tr>
<th>Parameter Cell (Å)</th>
<th>CuFeO2 (x = 0.50)</th>
<th>CuFe0.88O2 (x = 0.47)</th>
<th>D + Cu2O (x = 0.43)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>3.0348(5)</td>
<td>3.0349(5)</td>
<td>3.051(5)</td>
</tr>
<tr>
<td>c</td>
<td>17.157(3)</td>
<td>17.159(3)</td>
<td>17.161(3)</td>
</tr>
<tr>
<td>V</td>
<td>136.85(2)</td>
<td>136.88(2)</td>
<td>136.91(2)</td>
</tr>
<tr>
<td>O</td>
<td>0.1139(7)</td>
<td>0.1113(5)</td>
<td>0.1112(7)</td>
</tr>
<tr>
<td>Angle (deg)</td>
<td>90.14(12)</td>
<td>90.11(12)</td>
<td>90.09(12)</td>
</tr>
<tr>
<td>Cu–O</td>
<td>1.954(12)</td>
<td>1.910(9)</td>
<td>1.908(12)</td>
</tr>
<tr>
<td>Fe–O</td>
<td>1.972(6)</td>
<td>1.993(4)</td>
<td>1.994(6)</td>
</tr>
<tr>
<td>O–Fe–O,Fe</td>
<td>0.100(2)</td>
<td>0.092(2)</td>
<td>0.091(2)</td>
</tr>
<tr>
<td>Cu–O–Fe</td>
<td>7.94(5)</td>
<td>8.08(4)</td>
<td>8.09(5)</td>
</tr>
<tr>
<td>Cu–O–Fe–O</td>
<td>117.3(4)</td>
<td>118.5(4)</td>
<td>118.5(4)</td>
</tr>
</tbody>
</table>

In order to determine more accurately the temperature of formation of the delafossite phase, additional HT-XRD measurements were carried out for three mixtures (x = 0.25, 0.50, and 0.61) with temperature steps of 5 °C between 975 and 1045 °C. The value of 5 °C corresponds to the minimum applicable temperature step given the accuracy of the temperature probe and thermal regulation in the HT-XRD chamber. The compilation of the resulting phase assemblage is presented in Figure 3, with the HT-XRD patterns in Figure 3a and the temperature evolution of the amount (in mole percent) of each crystalline phases obtained from Rietveld analysis in Figure 3b.

On the basis of these accurate HT-XRD measurements, the formation temperature of delafossite CuFeO2 can be refined in the temperature range 1010–1015 °C, which is in good agreement with the value of 1015(2) °C published by Yamaguchi. Compared to the first set of experiments (Figure 2), the delafossite formation occurs in a very narrow temperature range (less than 25 °C). This is attributed to the decrease of the overall heating rate from 1.67 °C/min in Figure 2 to 0.0625 °C/min in Figure 3. Another consequence of the decrease of the heating rate is the evidence of CuO formation at around 1030–1040 °C, for the Cu-rich sample x = 0.25.

For the sample x = 0.50, in the 1020–1045 °C range, the delafossite is not the only phase because around 15 mol % of the spinel phase is present. The origin of this spinel phase may be due to either a slight Fe enrichment on the nominal composition (precisely, xreal = 0.54) or the presence of an excess of Cu in the delafossite phase (Cu1+x,FeO). These two hypotheses, leading to an excess of Fe in the extra phase, could be at the origin of stabilization of the spinel phase. This also could be due to an equilibrium between CuFeO2 and CuFe5O8, a spinel phase composed of Cu+.

The determination of transition temperatures by HT-XRD exhibits a significant uncertainty (±10 °C) because of the thermal regulation, temperature probes, and kinetics of phase transitions. As a consequence, these experiments were completed by TGA/DTA measurements performed between room temperature and 1100 °C, in air, with a heating rate of 5 °C/min. Pure delafossite CuFeO2 was used to determine its temperature of formation in air. The recorded TGA/DTA signals during heating at above 950 °C are presented in Figure 3c. At low temperature (about 400 °C, not shown here), the initial CuFeO2 powder oxidizes into a stoichiometric spinel and CuO according to eq 8.

$$\text{CuFeO}_2(s) + \frac{1}{4}\text{O}_2(g) \rightarrow \frac{1}{2}\text{CuFe}_2\text{O}_3(s) + \frac{1}{2}\text{CuO(s)}$$

(8)

As illustrated in Figure 3c (TGA, blue solid line), the weight gain measured around 1000 °C, related to eq 8, is 5.2%, which corresponds exactly to the theoretical mass gain (5.23%). A first endothermic peak associated with the start of a weight loss is evidenced at 1022 ± 2 °C. This peak is attributed to the delafossite phase formation according to eq 6. However, a residual mass gain of 0.70 wt % can be noted after this reaction. It can be explained by an increase of the O off-stoichiometry δ = 0.066 in CuFeO2+xδ (see a preliminary discussion in section 1.2). This hypothesis is supported by the larger a lattice parameter observed at high temperature (a = 3.035 Å at 50 °C and 3.074 Å at 1050 °C), which allows an easier insertion of the O anions into the Cu planes of the structure. A second endothermic peak, attributed to CuFeO2 incongruent melting into spinel and liquid phases, starts at 1070 ± 2 °C and ends at 1093 ± 2 °C. It is associated with a total weight gain of about 1.8% at 1093 °C.

In summary, coupled HT-XRD and TGA/DTA allow one to determine the delafossite formation temperature at 1022 ± 2 °C and its decomposition at 1070 ± 2 °C. These new experimental data strengthen the selection of Shishin et al.33 for the formation temperature (1019 °C), while we
recommend a slightly lower value than their selected decomposition temperature (1077 °C).

3.2. Eutectic Coordinates. Cu-rich samples ($x = 0.00$–0.20) were carefully analyzed in order to determine the formation temperature of CuFeO$_2$, Cu$_2$O, and liquid phases and provide a reliable measurement of the eutectic coordinates. The weight losses and heat flows recorded by TGA/DTA of samples $x = 0.03$, 0.07, 0.128, and 0.145 in air, with a heating rate of 1 °C/min and a cooling rate of 10 °C/min in Al$_2$O$_3$ crucible, are presented in Figure 4. For these measurements, the CuFeO$_2$ and Cu$_2$O starting materials were used to avoid any kinetic limitation that could lead to a shift of the transition temperatures. Because it is known that the presence of liquid CuO in contact with Al$_2$O$_3$ crucibles can form the parent delafossite compound CuAlO$_2$, it is important, in the present study, to ensure that the formation of this phase is limited. Recent works on the formation of CuAlO$_2$ in air$^{77}$ and the solubility of Al$_2$O$_3$ in CuO$^{78}$ indicate that CuAlO$_2$ is solid below 1238 °C and that the solubility of Al$_2$O$_3$ in CuO is limited to 0.11 wt % at 1150 °C. The very limited reactivity between the liquid phase and crucible was confirmed by the complete absence of CuAlO$_2$ in the XRD pattern and subsequent EPMA of the samples.

Parts a and b of Figure 4 show the TGA/DTA signals recorded for Cu-rich samples ($x = 0.03$, 0.07, 0.128 and 0.145), which contain an increasing amount of CuFeO$_2$ in the starting

![Figure 7](image-url)
powders. Three endothermic peaks are evidenced in all thermograms. The first endothermic peak at around 1022 °C is attributed to the delafossite formation in good agreement with the pure CuFeO$_2$ phase ($x = 0.50$) shown in Figure 3c. This transition temperature is clearly visible on the Fe-rich samples ($x = 0.128$ and 0.145), in contrast to the Cu-rich samples ($x = 0.03$ and 0.07), where the quantity of produced phase is too low to generate an intense signal. The second endothermic peak at (1031–1034) ± 2 is also associated with a weight loss. According to the literature data, it corresponds to the reduction of Cu$^{2+}$ to Cu$^+$ from CuO to Cu$_2$O (eq 4).

The third endothermic peak is detectable at (104–10515) ± 2 °C, with a slight weight gain. This latter peak is associated with formation of the liquid phase, at a temperature considerably lower than that previously measured (1090 ± 525). The slight weight gain, previously reported by Wolf et al., is similar to the system where the liquid contained two oxidation states of Cu (Cu$^{2+}$/Cu$^+$), attributed to the partial oxidation of Cu$^+$ to Cu$^{2+}$ and the associated O incorporation. The liquid phase was also analyzed during cooling (Figure 4b); values between 1048 and 1051 °C were measured to the exothermic picture of eutectic solidification, in good agreement with the values recorded upon heating. In the case of the eutectic’s temperature determination, one value was discarded out of the eight obtained (which corresponds to one measured temperature during heating and one during cooling for each of the four samples). The average of the seven selected measurements led to $T = 1050.1$ °C with an uncertainty of 1.7 °C. Our recommended value is $T_{\text{eutectic}} = 1050 ± 2$ °C because it is in the reliability range of expanded uncertainty, using a coverage factor of 2.45 with a level of confidence of 95%.

The other information extracted from the cooling curves is the solidus temperature measured at 1105, 1077, 1064, and 1067 ± 2 °C for samples $x = 0.03$, 0.07, 0.128, and 0.145, respectively. Tammann’s triangle method, which is based on analysis of the thermic effect of the eutectic formation, was used to determine the eutectic composition. The integrated surfaces of the DTA peaks represented as the red area in Figure 4b were plotted versus composition for the samples $x = 0.03$, 0.07, 0.128, and 0.145. The experimental information is summarized in Figure 5 with extrapolation of the thermic effect of eutectic formation in cooling. The intersection of both sides of the triangle at $x = 0.105$ mol % represents our best estimation of the eutectic composition.

3.3. Cationic Nonstoichiometry in Delafossite. HT-XRD Characterization of Stoichiometric CuFeO$_2$ + Minor Additions of CuO. Compositions close to the delafossite phase, i.e., with $x ≈ 0.50$, were also analyzed by HT-XRD with steps of 5 °C between 975 and 1045 °C in order to study the possible cationic nonstoichiometry range Cu$_{1-x}$Fe$_{1-x}$O$_2$. Starting from the CuO and Fe$_2$O$_3$ precursors, a spinel phase is systematically formed together with the delafossite phase when the initial stoichiometry contains an excess of Fe, i.e., when $x > 0.50$, such as, for instance, for $x = 0.60$ (Figure 2) or $x = 0.52$ (not shown here). In contrast, for $x < 0.50$, the delafossite phase is the only phase detected by XRD, such as, for instance, with the sample $x = 0.40$ at 1050 °C (Figure 2). However, as previously mentioned, in this part of the diagram, the phase assemblage can not be precisely quantified by Rietveld analysis because of the formation of a Cu-rich liquid phase ($x ≈ 0.105$). Because quantification of the amount of liquid phase is not possible, the amount and composition of delafossite cannot be determined either.

To overcome this difficulty, we carried out three successive HT-XRD analyses on (1) a stoichiometric delafossite CuFeO$_2$ ($x = 0.50$) sample, (2) the same sample with the addition of Cu$_2$O, leading to the composition $x = 0.47$, and (3) the same sample once again enriched in Cu$_2$O, leading to the composition $x = 0.43$. Furthermore, during the HT-XRD recordings, a nitrogen atmosphere was continuously maintained in the chamber in order to stabilize the Cu$^+$ valence and to avoid any decomposition of the delafossite phase at 1050 °C.

Figure 6a shows the XRD patterns at 50 °C of various compositions after thermal cycles at a maximal temperature of 1050 °C: the pristine stoichiometric $x = 0.50$ CuFeO$_2$ material after 1 cycle, the mixture $x = 0.47$ (i.e., CuFeO$_2$ + 6 mol % Cu$_2$O) before and after 2 cycles, and the mixture $x = 0.43$ [i.e., CuFeO$_2$ + (6 + 7) mol % Cu$_2$O] before and after 1 cycle. Figure 6b shows the evolution of the intensity of the Cu$_2$O(111) characteristic Bragg peak at 1050 °C during the dwells. The error bars correspond to the accuracy of the intensity measurement in the HT-XRD setup for scans of 30 and 45 min. This (111) peak appears at around 36.5° in 2θ at room temperature, as shown in Figure 6a.

The pure delafossite was first submitted to a thermal cycle up to 1050 °C. It confirms its stability under a nitrogen atmosphere, over the temperature range 50–1050 °C. Furthermore, the delafossite phase is pure, without any trace of Cu$_2$O or any other phase (black pattern in Figure 6a). The resulting Cu$_2$O intensity (20 cps) measured at 1050 °C and reported with a black diamond in Figure 6b is not significant and is attributed to statistical accumulation. The addition of 6 mol % Cu$_2$O is detected on the XRD pattern at 50 °C (the first blue pattern in Figure 6a) and confirmed by a measurable intensity at 1050 °C (blue squares in Figure 6b). After two temperature cycles and 180 min of cumulative dwell time at 1050 °C, the peak disappears, indicating that Cu$_2$O diffused in the delafossite phase.

A second addition of 7 mol % Cu$_2$O was then realized. The characteristic Cu$_2$O(111) peak is again clearly identifiable (the first red pattern in Figure 6a). A dwell of 240 min at 1050 °C leads to a partial extinction of the signal (red circles in Figure 6b). However, even after a total dwell time of 400 min, the peak intensity remains constant, indicating the presence of Cu$_2$O as a stable phase in the system. Back to 50 °C, the Cu$_2$O(111) peak is still visible (the second red pattern in Figure 6a).

This set of experiments proves that the delafossite can accept an excess of Cu in a range of compositions between $x = 0.47$ and 0.43.

Rietveld analysis carried out on the samples $x = 0.50$, 0.47, and 0.43 after their respective annealing at 1050 °C led to the structural parameters listed in Table 4. These data are fully consistent with the literature data reported in Table 1, and no clear change in the lattice constant can be associated with the Cu enrichment of the system. One can only notice some slight changes in the Cu–O and Fe–O distances (decrease of the Cu–O bond length from 1.95 to 1.91 Å and concomitant increase of the Fe–O length from 1.97 to 1.99 Å) in the delafossite structure from $x = 0.50$ to 0.47. No changes are evident from $x = 0.47$ to 0.43. This confirms the hypothesis of Cu enrichment into the delafossite phase up to $x = 0.47$. 
**EPMA on the Diffusion Couple.** EPMA was carried out in order to improve quantification of the nonstoichiometry range of the delafossite phase, thanks to a specific diffusion couple between CuO and Fe$_2$O$_3$ oxides.

The diffusion couple (see section 2.1 for the preparation protocol) allows one to obtain a material containing different zones, whereas the composition varies from the initial pure CuO and Fe$_2$O$_3$ to various mixtures of both oxides. Figure 7a shows the SEM image of the diffusion couple in a chemical contrast (backscattered electron, BSE) mode with the initial CuO/Fe$_2$O$_3$ interface in the lower part. Because of the elaboration conditions close to the melting temperature of the Cu compounds, the specimen exhibits large porosities in the Cu-rich region, which appears as big dark areas. Three representative zones containing the delafossite phase are also presented, which are all located in the Cu-rich region. The first region (zone 1) is located at the initial CuO/Fe$_2$O$_3$ interface, the second one (zone 2) is located around 400 μm from the interface, and the last one (zone 3) is located at more than 1300 μm from the interface. The overall Cu (Fe) content is increasing (decreasing) from zone 1 to zone 3.

The BSE image of the Cu-richest zone (zone 3) is shown in Figure 7c. It exhibits three different chemical contrasts, i.e., three different phases (delafossite, cuprite, and tenorite) because of variation of their average atomic numbers. A 0D-point EPMA measurement of the dark-gray area leads to 26.3(6) mol % Cu, 23.2(6) mol % Fe, and 50.5(6) mol % O. It corresponds to the formula CuFe$_{0.88}$O$_{1.92}$. EPMA indicates that the light-gray and white areas correspond to pure CuO and CuFe$_2$O$_4$, respectively, in good agreement with the electronic density. The BSE image of zone 2 (Figure 7d) shows similar areas, i.e., a homogeneous grain of the delafossite phase surrounded by the Cu$_2$O and CuO phases. The average composition of delafossite (Figure 7e) determined by 1D-line analysis in EPMA is 25.9(4) mol % Cu, 23.2(4) mol % Fe, and 50.9(13) mol % O. The normalized formula for delafossite in this region is CuFe$_{0.89}$O$_{1.96}$. This composition is homogeneous from side to side on the grain. The interface zone (zone 1) presents a Cu$_2$O grain (white area) with around 1.9% Fe solubilized (analysis from 0 to 6 μm; see Figure 7f). At 8 μm from the starting point, a delafossite grain is analyzed. In this grain, a clear evolution of the x ratio is observed from x = 0.48 on the Cu-rich side (distance = 8 μm) to x = 0.50 on the Fe-rich side (distance = 20 μm). The latter reaches the stoichiometric composition x = 0.50 at the interface with a grain composed of a spinel solid solution Cu$_{5/3}$Fe$_{x}$O$_{4}$. The maximal off-stoichiometry detected in the sample was in zone 3. For this composition, the Fe/(Cu + Fe) molar ratio is x = 0.47. This minimal Fe content is in accordance with HT-XRD experiments.

In order to determine if the area in which the chemical analysis shows a deficient Cu content corresponds to a pure nonstoichiometric delafossite or a mixture of stoichiometric delafossite (CuFeO$_2$) and copper oxide phase (CuO or Cu$_2$O), micro-Raman characterizations have been carried out in the same area. Beforehand, various reference spectra have been acquired on pure CuO, Cu$_2$O, and CuFe$_2$O$_4$ powders (Figure 7b). In parts c, d, and f of Figure 7 respectively related to zones 3, 2, and 1, the label “a” indicates the areas identified as the delafossite phase in which the Raman measurements were made. Raman spectra have been acquired very closed to the area where the chemical analysis has been carried out. The resulting spectra are shown in Figure 7b and compared to that obtained on the pure CuFeO$_2$ powder. This analysis confirms the presence of pure delafossite in the three zones, with no presence of copper oxides (CuO or Cu$_2$O). In zone 2, other Raman measurements have been made in the areas b and c, which confirms that the CuO and Cu$_2$O phases are clearly identified and fit well with the chemical analysis.

These coupled EPMA/SEM/Raman analyses prove that the delafossite structure could accommodate some Cu excess up to x = 0.47, i.e., with a CuFe$_{0.88}$O$_{2−δ}$ formula.

### 4. CONCLUSION

In this work, we reinvestigated the Cu–Fe–O system in air with a special focus on the delafossite phase. The starting point of this study was the phase diagrams available in the literature (Perrot et al.31 Khvan et al.32 and Shishin et al.33). Some inconsistencies were highlighted, especially regarding the delafossite stability range and the eutectic point coordinates. Besides, considering the composition of CuFeO$_2$, no work had so far provided real evidence about the existence of a cationic off-stoichiometry in this compound.

We performed a systematic experimental study of the full system under air between 50 and 1100 °C by in situ HT-XRD and TGA/DTA, and EMPA was performed to investigate the delafossite composition stability range in the copper–iron mixed oxide regions.

The stability region of delafossite was successfully determined in air, with a formation temperature at 1022 ± 2 °C and an incongruent decomposition at 1070 ± 2 °C. DTA curves were used to measure the liquidus temperature at 1050 ± 2 °C with a eutectic composition at x = 0.105 obtained by Tammann’s triangle method.

For the first time, a cationic off-stoichiometry of cuprous ferrite CuFe$_{1−y}$O$_{2−δ}$ was evidenced, with a maximum value of y = 0.12 measured by EPMA, this value being supported by HT-XRD analysis. No evidence of an Fe-rich delafossite was found, and complementary analysis under a controlled atmosphere shows narrow lattice variation with an increase of Cu in the delafossite structure. The updated experimental phase diagram obtained as a result of this work is presented in Figure 8 with experimental points.
ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorg-chem.9b00651.

Supplementary figure of XRD patterns for the sample x = 0.52 over the temperature range 975–1045 °C (PDF)

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Notes
The authors declare no competing financial interest.

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Supporting information
for

Insights on stability and cationic non-stoichiometry of CuFeO$_2$ delafossite
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Figure S1: a) X-ray diffraction patterns for the sample x=0.52 over the temperature range 975-1045 °C from simple oxides. b) Corresponding amount of crystalline phases quantified by Rietveld refinement.