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3 **Effect of drying on the composition of secondary**

4 **metabolites in extracts from floral parts of *Curcuma***

5 ***longa* L.**

6 **ABSTRACT**

7 **Background:** *Curcuma longa* is a rhizomatous herbaceous plant of Zingiberaceae family, originating
8 from South Asia and very widespread in hot and rainy regions of the globe. The rhizomes are very
9 popular spice and used as food additives for its coloring, aromatic, food preservation and nutritional
10 properties.

11 **Aim:** The aim of this study was to assess the drying effect of floral parts of *C. longa* at laboratory and
12 incubator temperature (35°C) on the composition of secondary metabolites in general and
13 polyphenolic compounds in particular.

14 **Place and Duration of Study:** The study was carried out at the Laboratory of Natural Products,
15 Department of Chemistry, Faculty of Sciences, University of Kinshasa between November 20 and
16 December 10, 2016.

17 **Methodology:** Different parts of *C. longa* floral parts were collected. Petals were collected every day
18 while sepals were collected 20 days after the first petal appeared. The phytochemical screening was
19 used as per the standards protocol and it was assessed between floral parts dried in the room
20 temperature and floral parts dried in the incubator at 35 °C.

21 **Results:** The findings revealed the presence of total polyphenols, flavonoids, tannins in petal extracts
22 dried in an incubator at 35°C. However, flavonoids and tannins were not detected in extracts from
23 petals dried at laboratory temperature. Phytochemical screening findings of the organic phase of *C.*
24 *longa* floral parts revealed the presence of free quinones and terpenes in sepal extracts dried in the
25 incubator at 35°C while those of extracts from the same part of the dried plant in the laboratory
26 temperature revealed the presence of terpenes and steroids.

27 **Conclusion:** Future studies should carry out a similar study using the spectrophotometry method to
28 determine polyphenolic compounds and confirm our hypothesis on the degradation of polyphenolic
29 compounds during the drying of *C. longa* floral parts at laboratory temperature.

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31 **Keywords:** *Curcuma longa*, Phytochemical screening, Effect of drying, Laboratory, Incubator

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33 **1. INTRODUCTION**

34 *Curcuma longa* or saffron is a rhizomatous herbaceous plant, perennial, 50 cm to 1m high, of the
35 Zingiberaceae family, originating from South Asia and very widespread in hot and rainy regions of the
36 globe such as Asia, Africa and Oceania [1]. It is widely cultivated in India but also to a lesser extent in
37 China, Taiwan, Japan, Burma, Indonesia and Africa. The intense cultivation of this plant is due to its
38 rhizomes [1-2]. The latter dried and powdered are a very popular spice and are used as food additives
39 for its coloring, aromatic, food preservation and nutritional properties. In addition, *C. longa* rhizomes
40 are a key ingredient in traditional South American and Asian medicines such as Ayurvedic medicine
41 (Indian medicine). Thus, *C. longa* is a remedy against gastrointestinal, digestive disorders,
42 inflammatory diseases, skin diseases [1-5].

43

44 Numerous studies correlated with ancestral knowledge have demonstrated the preventive action of *C.*
45 *longa* rhizomes on many diseases such as cancers, cardiovascular diseases [6]. Several studies have
46 shown that extracts from *C. longa* rhizomes possess a large pharmacological potential namely: anti-
47 cancer, anti-inflammatory, healing, cholesterol-lowering, hypoglycemic, anti-Alzheimer's,
48 antiplasmodial, anti-inflammatory, antioxidant, antibacterial, antifungal, anti-venomous, antipyretic,
49 analgesic, inhibits the action of HIV-1 integrase, and HIV-1 integrase protein replication, protects

50 against diabetic retinopathy and many other pathologies [2-5, 7-12]. In addition to rhizomes, the other
51 organs of the plant (leaves, roots and floral parts) are less used. The leaves of *C. longa* are
52 sometimes used in cooking or to extract essential oils [5]. Ritwiz *et al.* [13] have shown that extracts
53 from the leaves of *C. longa* have antioxidant, antibacterial potential and can modulate immunological
54 properties. Meanwhile Mbadiko *et al.* [11-12], report that total methanol extracts of rhizomes, roots,
55 leaves and floral parts (petals and sepals) possess an antisickling activity. Most of the studies carried
56 out so far focused only on the rhizomes of the plant, studies on the leaves, roots and floral parts of *C.*
57 *longa* remain poor or less reported. This justifies our interest in carrying out a phytochemical study on
58 the extracts of the floral parts (petals and sepals) of *C. longa*.

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60 In addition, Mbadiko *et al.* [11-12] reported that floral parts of *C. longa* contained low levels of
61 secondary metabolites (polyphenolic compounds) and thought this was related to the drying effect.
62 These authors reported that during the drying of *C. longa* floral parts, and in particular the petals at
63 laboratory temperature, the latter tended to soften and lose color, suggesting degradation of the
64 phytoconstituents under cellular conditions. The negative impact of post-harvest treatments of plant
65 samples on phytoconstituent composition has also been reported by Singh *et al.* [14]. The degradation
66 of polyphenolic compounds in plants under cellular conditions, i.e. fresh samples, has also been
67 reported by Yan *et al.*, [15] and Chang *et al.* [16]. Thus, this study is part of a context to assess the
68 effect of post-harvest treatments, in particular drying on the composition of secondary metabolites in
69 extracts from *C. longa* floral parts.

70 71 **2. MATERIALS AND METHODS**

72 **2.1 Collection of plant material**

73 As biological material, the floral parts of *C. longa* were used. It was observed that *C. longa* renews its
74 petals every 24 hours. Thus, petals were collected every day (between November 20 and December
75 10, 2016). The sepals were collected 20 days after the first petal appeared.

76 77 **2.2 Packaging of plant material**

78 After the collection, the petals and sepals were cleaned, washed quickly with tap water. Some of our
79 samples were dried at laboratory temperature and the other part at the incubator (Melag Nurfur
80 Wechselstrom brand) at 35°C.

81 82 **2.3 Preparation of aqueous and organic extracts**

83 The aqueous extracts were obtained by macerating 5g of the powder of our four samples each in 50
84 mL of distilled water during 24 hours at room temperature, then filtered using filter paper (Whatman
85 n°1). Maceration of 2g of the powder from our samples in 20 mL of ethyl acetate during 24 hours of
86 incubation then filtered using a filter paper (Whatman n°1).

87 88 **2.4 Phytochemical screening**

89 Phytochemical screening represents all the qualitative techniques used to determine or identify the
90 different chemical groups (secondary metabolites) contained in an extract. These chemical groups are
91 identified by means of coloring and precipitation reactions that take place by adding specific reagents
92 [11, 17].

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102 **3. RESULTS AND DISCUSSION**

103 The phytochemical screening of aqueous of *C. longa* floral parts dried in the laboratory temperature
 104 and at the incubator at 35 °C is presented in the following table.

105
 106 **Table 1 : Phytochemical screening of aqueous phase of floral parts dried in the laboratory**
 107 **temperature and in the incubator at 35 °C**
 108

Secondary metabolites	Laboratory temperature	Incubator (35 °C)	Laboratory temperature	Incubator (35 °C)
	Petals		Sepals	
Saponines	-	-	-	-
Total polyphenols	+	+	+	+
Flavonoids	-	+	-	+
Anthocyanins	-	-	+	+
Leuco-anthocyanins	-	-	-	-
Alkaloids	+	-	+	+
Bound Quinones	-	-	-	-
Tannins	-	+	+	+

109 Legend : + :presence of the phytoconstituent, - :absence of the phytoconstituent

110
 111 Table 1 shows that the aqueous phases of different floral parts (petals and sepals) of *C. longa* dried in the
 112 incubator at 35 °C differ in their composition in secondary metabolites. Total polyphenols,
 113 flavonoids, and tannins were detected in petal extracts. On the other hand, the analysis on sepal
 114 extracts revealed the presence of total polyphenols, flavonoids, anthocyanins, alkaloids and tannins. It
 115 should be noted that saponins were not detected in all extracts. **While the extracts *C. longa* petals**
 116 **dried in the laboratory** possess total polyphenols and alkaloids while the presence of total polyphenols,
 117 anthocyanins, tannins and alkaloids have been highlighted in sepal extracts.

118
 119 It should be noted that total polyphenols was detected in the petal extract dried in the laboratory
 120 temperature and in the incubator at 35 °C. For the sepal extracts, total polyphenols, anthocyanins,
 121 alkaloids and tannins were found in both settings.

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 123 **Table 2 : Phytochemical screening of organic phase of *C. longa* floral parts dried in the**
 124 **laboratory temperature and at the incubator at 35 °C**
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Secondary metabolites	Laboratory temperature	Incubator (35 °C)	Laboratory temperature	Incubator (35 °C)
	Petals		Sepals	
Terpenes	-	+	+	+
Steroids	-	-	+	-
Free Quinones	-	+	-	+

126 Legend : + :presence of the phytoconstituent, - :absence of the phytoconstituent

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 128 **Table 2** shows that the organic phase of extracts from different floral parts of *C. longa* dried in the
 129 incubator at 35 °C has the same chemical profile. The presence of terpenes and free quinones has
 130 been detected in both parts of the plant. **Meanwhile for the petal extracts dried in the laboratory**
 131 **temperature didn't show any compounds and the sepal extracts only terpenes and steroids were**
 132 **found. It should be noted that only terpenes was found in sepal extracts for both settings.**

133
 134 **DISCUSSION**

135
 136 Phytochemical analyses in this study revealed the presence of total polyphenols, flavonoids, tannins in
 137 petal extracts dried in an incubator at 35°C. However, flavonoids and tannins were not detected in
 138 extracts from petals dried at laboratory temperature. This implies their degradation under laboratory
 139 conditions. In fact, since biochemical reactions necessarily occur in an aqueous environment, drying at
 140 a temperature that does not remove moisture at a short time would favor the action of certain enzymes
 141 that are activated during drying [19], could be at the origin of the degradation of certain secondary
 142 metabolites. As mentioned above, the degradation of polyphenolic compounds in fresh plant samples
 143 has also been reported by Yan *et al.*, [15] and Chang *et al.* [16].

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145 In addition, phytochemical screening of sepal extracts dried in the incubator at 35°C revealed the
146 presence of total polyphenols, flavonoids, anthocyanins, tannins and alkaloids while those of sepals
147 dried at laboratory temperature showed the presence of total polyphenols, anthocyanins, tannins and
148 alkaloids. The absence of flavonoids in sepals and petals dried at laboratory temperature would
149 suggest their degradation during drying; this would at the same time justify the discoloration of these
150 samples during drying. Indeed, flavonoids are pigments responsible for the yellow, orange and red
151 discoloration of different plant organs [18]. Further studies are needed to confirm this hypothesis.

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153 Phytochemical screening findings of the organic phase of *C. longa* floral parts revealed the presence
154 of free quinones and terpenes in sepal extracts dried in the incubator at 35°C while those of extracts
155 from the same part of the dried plant in the laboratory temperature revealed the presence of terpenes
156 and steroids. Phytochemical screening of petal extracts dried in the incubator at 35°C showed the
157 presence of free quinones and terpenes. These compounds were not detected in petal extracts dried
158 at laboratory temperature. The absence of quinones in sepal and petal extracts dried at laboratory
159 temperature could justify the discoloration of these organs during drying at room temperature and
160 would affirm their degradation during drying. Boukri [18] reported that quinones are colored and bright
161 substances, usually red, yellow or orange.

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163 CONCLUSION AND RECOMMENDATIONS

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165 The purpose of this study was to assess the drying effect of floral parts (petals and sepals) at
166 laboratory or incubator temperature at 35°C on the composition of secondary metabolites in general
167 and polyphenolic compounds in particular. In light of the findings obtained, we believe that drying the
168 floral parts (petals and sepals) of *C. longa* at laboratory temperature would promote the degradation of
169 certain polyphenolic compounds (flavonoids and quinones). This would prevent the action of certain
170 enzymes that would be activated after harvest or during drying and would prevent the degradation of
171 polyphenolic compounds. Samples should also be cut into small pieces before drying to increase the
172 surface area of contact of the samples with heat and rapidly reduce moisture. The study on the drying
173 effect on the composition of secondary metabolites has not yet been reported in the literature.

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175 Future studies should carry out a similar study using the spectrophotometry method to determine
176 polyphenolic compounds and confirm our hypothesis on the degradation of polyphenolic compounds
177 during the drying of *C. longa* floral parts at laboratory temperature.

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180 COMPETING INTERESTS

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182 Authors have declared that no competing interests exist.

183

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