

Chemical composition and antioxidant activity of *Dialiumguineense* fruit (Fabaceae) from Côte d'Ivoire

Abstract : This study is devoted to the phytochemical analysis of *Dialiumguineense* fruit, which is particularly popular in urban areas in Côte d'Ivoire. Despite its high consumption, there is still little information about its potential. To assess and identify the levels of various secondary metabolites, hydromethanolic extracts of the pulp, kernel, and shell of *D. guineense* fruit were analyzed via qualitative and quantitative methods (colorimetric and TLC tests) : total phenols (40.201 ± 1.8 ; 509.023 ± 14.5 and 194.368 ± 2.2 μg EAG/g dry extract), flavonoids (58.958 ± 6.1 ; 216.771 ± 21.7 and 238.385 ± 6.2 μg EC /g dry extract), condensed tannins (264.167 ± 13.9 ; 411.667 ± 25.8 and 282.500 ± 5.2 μg EC/ mg dry extract) and hydrolysable tannins (0.44 ± 0.03 %; 0.21 ± 0.02 % and 0.30 ± 0.03 %). Furthermore, the analysis of antioxidant potential (DPPH and FRAP) showed satisfactory activity overall, with special attention given to the kernel and shell. These results indicate that the fruit of *D. guineense* could be viewed as a potential source of antioxidant compounds that can help prevent or fight various pathologies associated with oxidative stress.

Key words: *Dialiumguineense*, fruit, chemical composition, antioxidant activity

1. Introduction

Dialiumguineense, commonly referred to as black tamarind or velvet tamarind, is an endangered plant species that is widely used by people in Côte d'Ivoire [1, 2]. Despite the high level of consumption, this species has not undergone any studies. [3] has conducted research on the

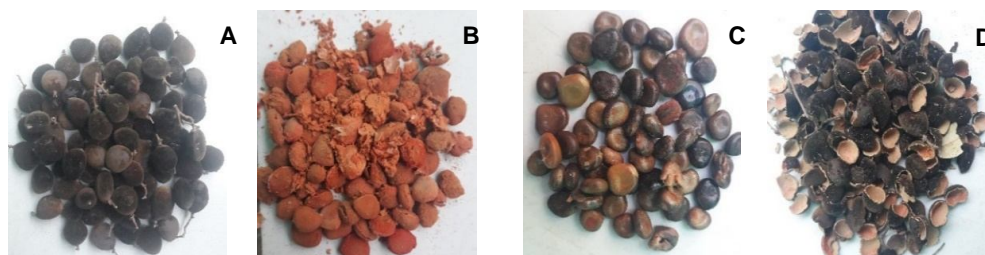
48 shell, pulp, and kernel of this fruit, which is why. The results reveal the presence of various
49 minerals, with a particularly high concentration of calcium (Ca), potassium (K) phosphorus (P)
50 and magnesium (Mg). Despite not being factored into men's eating habits, the kernel is a
51 significant protein source ($23.58 \pm 1.18\%$) in comparison to other parts of the fruit. In terms of
52 energy utilization, the pulp, kernel, and shell can generate 317.01 ± 0.14 kcal/100g, $364.47 \pm$
53 0.35 kcal/100g, and 351.02 ± 0.32 kcal/100g, respectively. The kernel contained 56.79 ± 1.22
54 g/100 g of carbohydrates, while the pulp had $70.15 \pm 0.9\%$, and the shell had $79.44 \pm 0.7\%$. The
55 hull had a significant amount of total fiber ($30.14 \pm 0.02\%$). The anti-nutritional elements were
56 found to be only in the kernel (14.83 ± 0.07 mg/100g), and oxalates were found in all parts of the
57 fruit through the evaluation of the anti-nutritional elements. With this information, the
58 understanding of *D. Guineans* in Côte d'Ivoire is enhanced and new opportunities for this fruit's
59 development are opened up [4–6] Incorporating antioxidant-rich food supplements into our diet
60 is essential to combat these conditions [7]. Synthetic antioxidants have been criticized for their
61 potential toxic risks [8]. Many researchers are investigating antioxidant molecules that come
62 from nature for this reason. *D. guineense* has been studied in various regions, particularly in
63 Nigeria, and it has been found to have phytochemicals with interesting potential [9, 10]. *D.*
64 *guineense* has the potential to be a promising source of antioxidant substances. The aim of this
65 study is to examine the shell, kernel, and pulp of the fruit of *D. guineense* from Côte d'Ivoire's
66 chemical composition and antioxidant potential.

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2. Materials and methods

2.1. Plant material

69 *Dialiumguineense* fruits (Figure 1A) were collected in March 2020 in Dimbokro ($6^{\circ}39'$ north,
70 $4^{\circ}42'$ west), a town in the N'Zi region of central Côte d'Ivoire. They were identified by an
71 ethnobotanist from Nangui ABROGOUA University and then certified at the Centre National de
72 Floristique in Abidjan in accordance with existing herbariums (N^oUCJ009331). After collection,
73 the fruits were cleaned and shelled. The various resulting parts (pulp, kernel and shell) (Figure
74 1(B-D)) were dried in an air-conditioned room (18°C) for 21 days, then reduced to powder using
75 an electric grinder.
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**Figure 1: Unhulled fruit (A), pulp (B), kernels (C) and shells (D) of *Dialiumguineense*
(Photo taken by ADOPO in 2020)**

2.2. Methods

2.2.1. Preparation of samples

83 100 g of powder from each part of the fruit was macerated in 600 mL of methanol (MeOH, 3×24
84 h) at room temperature (25°C) with continuous stirring. After vacuum filtration, the filtrates were

85 combined and concentrated using a rotary evaporator (Büchi R11) at 40°C. The aqueous extracts
86 obtained were kept in a refrigerator (4°C) for 24 h, in order to precipitate the lipophilic
87 compounds. Once the lipophilic compounds had been removed by decantation, part of the
88 aqueous extracts were oven-dried at 45°C until the dry hydromethanolic crude extracts were
89 obtained, which were used for phytochemical screening using colour reactions, quantitative
90 analysis and antioxidant potential studies. The second part of the aqueous extracts was
91 successively exhausted in solvents of increasing polarity (3×50 mL): hexane (n-C₆H₁₄),
92 chloroform (CHCl₃), ethyl acetate (AcOEt) and n-butanol (n-BuOH). The selective extracts
93 obtained were used for phytochemical screening by thin-layer chromatography (TLC).

94 **2.2.2. Qualitative analysis**

95 The chemical composition of every part of the *D. guineense* fruit was determined through
96 qualitative analysis. To achieve this, colorimetric tests and TLC were employed [11–18].

97 **2.2.3. Quantitative analysis**

98 The content of certain compounds in each part of the *D. guineense* fruit was evaluated through
99 quantitative analysis using spectrophotometry. The Folin-Ciocalteu colorimetric method was
100 used to determine the total phenols [12, 19, 20]. Aluminum chloride method was used to
101 estimate the total flavonoid levels [21, 22]. The vanillin hydrochloride assay was used to
102 determinate condensed tannins [22-24] method was used to evaluate the level of hydrolysable
103 tannins.

104 **2.2.3.1. Study of antioxidant potential**

105 DPPH test [25-27] and FRAP method [28, 29] were used to assess the free radical scavenging
106 capacity of the various parts of *D. guineense* fruit.

107 **2.2.4. Statistical analysis**

108 Statistical analyses were performed using IBM SPSS Statistics version 22 software. Assay
109 results, expressed as the mean of six trials ± standard deviation, were subjected to a one-factor
110 ANOVA at the 5% threshold. Tukey's test (verified homoscedasticity) and Tamhane's T2 test
111 (unverified homoscedasticity) were used as post hoc tests, when the ANOVA test led to
112 significant differences between means. These tests make it possible to group the different
113 means into homogeneous subsets. Finally, the Pearson test was applied to establish the
114 correlations between the different levels obtained.

115 **3. Results and discussions**

116 **3.1. Chemical composition of the different parts of the *D. guineense* fruit**

117 Several secondary metabolites were discovered during phytochemical screening using
118 colorimetric tests (Table I). FeCl₃ solution was used to identify polyphenols in all parts of *D.*
119 *Guineans* fruit. Kernel and pulp extracts had intense blue-black coloration, while the shell
120 extracts had green-black coloration, suggesting a significant presence of these compounds in
121 these parts. Different classes of flavonoids could be identified by detecting flavonoids in all three-
122 study media using Shinoda, ammonia, and vanillin tests. According to [17], flavones and
123 flavonols can be detected by a yellow color at room temperature or a red-orange color when
124 heated by ammonia test, which can be verified by Shinoda's test. The presence of flavonoids is
125 confirmed by a positive result observed with vanillin. All parts of the fruit contain tannins,

126 coumarins, sterols, and turbanes. Saponins were detected with foam indices of 100, 125 and 200
 127 for the pulp, kernel and shell respectively. The kernel and shell showed positive results from
 128 Borntraeger reagent, which suggests that quinones are present, as well as alkaloids detected by
 129 the Dragendorff and Wagner reagents. In spite of this, cardiotoxic glycosides were not present in
 130 all parts of the fruit According to studies on the species in Nigeria, which [30] observed in
 131 ethanolic, methanolic, and aqueous extracts, these results agree with those of other studies.

132 **Table I: Compounds detected in different parts of *D. guineense* fruit**

		Pulp	Kernel	Shell
Polyphenols		+	+	+
	<i>Shinoda's test</i>	+	+	+
Flavonoïds	<i>Ammonia test</i>	+	+	+
	<i>Vanillin test</i>	+	+	+
Coumarins		+	+	+
Tannins		+	+	+
	<i>Dragendorff's test</i>	-	-	+
Alcaloïds	<i>Wagner's test</i>	-	+	+
	<i>Picric acid test</i>	-	-	-
Sterols and terpenes		+	+	+
Quinones		-	+	+
Saponins		+(Im = 100)	+(Im = 125)	+(Im = 200)
Cardiotonic glycosides		-	-	-

133 + : Positive test; - : Negative test

134 In the end, the fruit of *D. guineense* has phytochemicals that have the potential to be beneficial
 135 for the body. The kernel and shell, which are often ignored by the public, could have potential
 136 uses in other fields because of their intriguing chemical composition.

137 3.2. TLC profiles of selective extracts

138 The presence of various phytochemicals, including sterols, terpenes, coumarins, flavonoids,
 139 tannins, and anthocyanins, was uncovered through TLC screening of *D. Guineans* shell (Coq),
 140 kernel (Noy), and pulp (Pul) (Tables II and III). To study the migration of compounds, present in
 141 hexane and chloroform extracts, solvent gradients containing n-C₆H₁₄/CHCl₃/AcOEt (4:10:1,
 142 V/V/V) and (6:10:6, V/V/V) were used. To determine what compounds were in the ethyl acetate
 143 and n-butanol extracts, the AcOEt/CHCl₃/EtOH/ (V/V/V) mixture was employed. Different
 144 developers were used to identify the compounds. Liebermann-Bürchard reagent was utilized to
 145 view sterols, which appeared brown and green to the naked eye and yellow and yellow-green
 146 under UV at 365 nm (Figure 2A). So, these compounds were revealed in hexanolic and
 147 chloroform extracts from shells (R_f = 0.34) and (R_f = 0.19; 0.26; 0.4 and 0.8), kernels (R_f = 0.32;
 148 0.6 and 0.92) and (R_f = 0.12; 0.23; 0.62 and 0.88) and pulps (R_f = 0.3 and 0.81) and (R_f = 0.72;
 149 0.8 and 0.92) respectively. The same developer was used to detect terpenes, in particular
 150 triterpene genins[18, 31], which appear as blue and purple spots to the naked eye, and orange-
 151 yellow spots under UV light at 365 nm for lupane-type triterpene genins. Terpenes are revealed
 152 in hexanolic extracts: Coq (R_f = 0.08; 0.14; 0.2; 0.34; 0.4; 0.5; 0.6; 0.7 and 0.9), Noy (R_f = 0.08;
 153 0.14; 0.2; 0.3; 0.45; 0.6; 0.7 and 0.81) and Pul (R_f = 0.08; 0.2; 0.3; 0.41; 0.5; 0.54; 0.56; 0.64;

154 0.7 and 0.9) (Figure 2A) and chloroformic : (Coq (Rf = 0.19; 0.22; 0.3; 0.4; 0.54; 0.62; 0.71; 0.86;
155 0.89 and 0.94), Noy (Rf = 0.71; 0.88; 0.91 and 0.94) and Pul (Rf = 0.12; 0.27; 0.35; 0.57; 0.67;
156 0.72; 0.81; 0.88 and 0.92)). The sulphuric vanillin solution was employed to show the presence
157 of terpenes, which can be detected by looking at them in violet, pink, and orange and under UV
158 light at 365 nm. In hexane extracts, terpenes were observed at the following Rf values: 0.08;
159 0.14; 0.2 and 0.46 (Coq), 0.08; 0.27; 0.5; 0.6; 0.7 and 0.92 (Noy) and 0.08; 0.3; 0.41; 0.5; 0.56;
160 0.64 and 0.8 (Pul). In chloroform extracts, they are found at Rf values: 0.08; 0.16; 0.62; 0.71 and
161 0.86 (Coq) and 0.12 and 0.88 (Pul). Sulphuric vanillin was also used to confirm the existence of
162 sterols [31] in the various parts of the *D. guineense* fruit, visible as blue spots to the naked eye
163 and under UV light at 365 nm. The Rf values in the hexanic extracts are: 0.21; 0.62 and 0.8
164 (Coq), 0.27 and 0.8 (Noy) and 0.32; 0.64 and 0.8 (Pul). In chloroform extracts, they are at Rf:
165 0.26; 0.37 and 0.88 (Coq), 0.84 and 0.88 (Noy) and 0.8 (Pul). Flavonoids were identified using
166 ammonia vapour and aluminum chloride solution (1 The flavonoids are made yellow by this
167 solution, and it can vary in coloration from blue to brown or yellow-green under UV 365 nm [11,
168 31]. In chloroform extracts, they are visible at Rf values of 0.06; 0.17; 0.4; 0.49; 0.54; 0.86 and
169 0.89 (Coq), at 0.12; 0.23; 0.29; 0.59 and 0.82 (Noy) and at 0.12; 0.35; 0.4; 0.57; 0.85; 0.88 and
170 0.9 (Pul (Rf) (Figure 2B), in ethyl acetate extracts, the Rf values are: 0.1; 0.19; 0.27; 0.34; 0.42;
171 0.56; 0.6; 0.7 and 0.79 (Coq), 0.19; 0.29; 0.4; 0.46; 0.61 and 0.79 (Noy) and 0.08; 0.19; 0.55;
172 0.66; 0.8 and 0.9 (Pul) and in the n-butanolic extracts the Rf values are 0.06; 0.11; 0.16; 0.2;
173 0.25; 0.31; 0.45; 0.59; 0.67; 0.74 and 0.8 (Coq), 0.17; 0.31; 0.41 and 0.49 (Noy) and 0.1; 0.1;
174 0.2; 0.6 and 0.8 (Pul). In the presence of ammonia vapour, flavonoids are visible as yellow and
175 green to the naked eye and under UV 365 nm [32]. This is the case in the chloroform extracts:
176 Coq (Rf = 0.06; 0.21; 0.49 and 0.59), Noy (Rf = 0.12; 0.25; 0.4; 0.87 and 0.91) and Pul (Rf =
177 0.12; 0.22; 0.29; 0.35; 0.75 and 0.88) (Figure 2B); ethyl acetate: Coq (Rf = 0.07; 0.17; 0.34; 0.49
178 and 0.62), Noy (Rf = 0.12 and 0.69) and Pul (Rf = 0.1; 0.45; 0.56; 0.71; 0.77 and 0.9) and n-
179 butanol extracts : Coq (Rf = 0.16; 0.31; 0.5 and 0.75), Noy (Rf = 0.19; 0.41; 0.5 and 0.6) and Pul
180 (Rf = 0.75)). In addition, ammonia vapour revealing blue or violet coloration indicates the
181 presence of anthocyanins [11] in chloroform extracts at the following Rf values: 0.87 and 0.91
182 (Noy) and 0.88 (Pul) (Figure 2B); in ethyl acetate extracts, they are at Rf 0.22 and 0.86 (Coq),
183 0.19; 0.36 and 0.42 (Noy) and 0.66 (Pul) and in n-butanolic extracts, they are at the following Rf
184 values: 0.07; 0.12 and 0.21 (Coq), 0.06; 0.08; 0.14 and 0.27 (Noy) and 0.12 and 0.89 (Pul (Rf).
185 Yellow spots observed with the naked eye using a 5% methanolic KOH solution indicate the
186 presence of coumarins. This yellow coloration can intensify or diversify under UV/365 nm light.
187 In chloroform extracts, coumarins are visible at the following Rf values: 0.06; 0.12; 0.3; 0.3; 0.47;
188 0.54 and 0.69 (Coq), 0.06; 0.12; 0.25; 0.5 and 0.82 (Noy), and 0.06; 0.12; 0.35; 0.47; 0.85 and
189 0.9 (Pul) (Figure 2B). In ethyl acetate extracts, Rf values were: 0.14; 0.19; 0.22; 0.3; 0.41; 0.51
190 and 0.6 (Coq), 0.5; 0.7 and 0.85 (Noy), and 0.08; 0.12; 0.22; 0.26; 0.82 and 0.9 (Pul). In n-
191 butanol extracts, coumarins appeared at the following Rf values: 0.11; 0.16; 0.2; 0.25; 0.32;
192 0.45; 0.6; 0.69 and 0.75 (Coq), 0.27; 0.4 and 0.5 (Noy), and 0.07; 0.29; 0.4 and 0.76 (Pul).
193 Tannins were detected with a 2% (w/v) FeCl₃ aqueous solution, appearing as grey or brown
194 stains [31, 33] in ethyl acetate extracts at Rf: 0.08; 0.19; 0.22; 0.31 and 0.71 (Coq), 0.24 (Noy),
195 and 0.71 (Pul) (Figure 2C). In n-butanolic extracts, the Rf values for tannins are: 0.16 and 0.25
196 (Coq), and 0.27 and 0.4 (Noy).

197 **Table II: Compounds identified in the hexanolic and chloroform extracts of the different**
 198 **parts of the *D. guineense* fruit.**

Solvents	Extracts	Possible compounds : [Rf], Colour, Developer
Hexane	Pul	Sterols : [0.3],Br ^e ; [0.32 ; 0.64],B ^a ; [0.81],Ve ^e , B ^a Terpenes : [0.08],J-Or ^e , Ve ^d , Vi ^a ; [0.2],J-Or ^e , [0.3],J-Or ^e , Vi ^a ; [0.41],J-Or ^e , Or ^a ; [0.5],J-Or ^e , Ve ^a ; [0.54],R ^e ; [0.56],R ^e , Ro ^a ; [0.64],J-Or ^e , Or ^a ; [0.7],J-Or ^e ; [0.8],Ro ^a ; [0.9],J-Or ^e
	Noy	Sterols : [0.32 ; 0.6 ; 0.92],Br ^e ; [0.27 ; 0.8], B ^a Terpenes : [0.08],J-Or ^e , Vi ^a ; [0.14],J-Or ^e ; [0.2],J-Or ^e ; [0.27],Vi ^a ; [0.32 ; 0.45],J-Or ^e ; [0.5],Vi ^a ; [0.6],J-Or ^e , Vi ^a ; [0.7],Vi ^{e,a} ; [0.81],J-Or ^e ; [0.92],J-Or ^a
	Coq	Sterols : [0.34], Br ^e ; [0.21 ; 0.62 ; 0.8],B ^a Terpenes : [0.08 ; 0.14 ; 0.2],J-Or ^e , Vi ^{ea} ; [0.34 ; 0.4],J-Or ^e ; [0.46],Vi ^a ; [0.5 ; 0.6 ; 0.7],J-Or ^e ; [0.9],J-Or ^e
Chloroform	pul	Sterols : [0.72 ; 0.8 ; 0.92],Br ^e ; [0.8],Br ^e , B ^a Terpenes : [0.12], J-Or ^e , B ^a ; [0.27 ; 0.35],J-Or ^e ; [0.57 ; 0.67 ; 0.72 ; 0.81 ; 0.92],B ^e ; [0.88],B ^e , B ^a Flavonoïds : [0.12],Ve ^b , J ^f ; [0.22 ; 0.29], J ^f ; [0.35],B ^b , Ve ^f ; [0.47], B ^b ; [0.57],J ^b ; [0.75], Ve ^f ; [0.88], B ^b , B ^f ; [0.9],J ^b Anthocyanins : [0.87],B ^f Coumarins : [0.06],Ve ^c ; [0.12 ; 0.35],J ^c ; [0.47],Ve ^c ; [0.54],B ^c ; [0.85 ; 0.9],J ^c
	Noy	Sterols : [0.12],J-Ve ^e ; [0.23; 0.62],Ve ^e ; [0.84],B ^a ; [0.88],J ^e , B ^a Terpenes : [0.71 ; 0.88], J-Or ^e ; [0.91],B ^e ; [0.94], J-Or ^e Flavonoïds : [0.12],Ve ^b ,J ^f ; [0.23],Ve ^b ; [0.25],J ^f ; [0.29],Ve ^b ; [0.4],Ve ^f ; [0.59],J ^b ; [0.82],B ^b ; [0.87],B ^f ; [0.91],B ^f Anthocyanins : [0.87 ; 0.91],B ^f Coumarins : [0.06],Ve ^c ; [0.12 ; 0.25 ; 0.5],J ^c ; [0.4],B ^c ; [0.82],Ve ^c ; [0.91],B ^c
	Coq	Sterols : [0.19],Br ^e ; [0.26],Ve ^e , B ^a ; [0.37 ; 0.88],B ^a ; [0.4 ; 0.8],Br ^e Terpenes : [0.08 ; 0.16],B ^e ,Or ^a ; [0.19 ; 0.22 ; 0.3 ; 0.4],B ^e ; [0.62 ; 0.71 ; 0.86], B ^e , Or ^a ; [0.54 ; 0.89 ; 0.94],J-Or ^e Flavonoïds : [0.06],J ^{b,f} ; [0.17],B ^b ; [0.21],J ^f ; [0.4],B ^b ; [0.49],Ve ^{b,f} ; [0.54],J ^b ; [0.59],J ^f ; [0.86],J ^b ; [0.89],B ^b Coumarins : [0.06 ; 0.12],Ve ^c ; [0.3 ; 0.35],J ^c ; [0.47 ; 0.54 ; 0.69],J ^c ; [0.88],R ^c

199 a : compounds developed with sulfuric vanillin; b: compounds developed with aluminum chloride (AlCl₃); c:
 200 compounds developed with 5% potassium hydroxide (KOH); e: compounds developed with 1% Liebermann-Bürchard
 201 reagent ; f: compounds developed with ammonia vapour (NH₃).
 202 B: blue, Br: brown, J: yellow, Or: orange, R: red, Ro: pink, Ve: green, Vi: violet or purplish.
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208 **Table III: Compounds identified in ethyl acetate and n-butanol extracts of different parts**
 209 **of *D. guineense* fruit.**

Solvents	Extracts	Possible compounds : [Rf], Colour, Developer
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Ethyl acetate	Pul	Flavonoïds : [0.08 ; 0.19],J ^b ; [0.1],J ^f ; [0.45 ; 0.56],J ^f ; [0.55],Ve ^b ; [0.66],Ve ^b , P ^f ; [0.71 ; 0.77],Ve ^f ; [0.8],Ve ^b ; [0.9],Ve ^{b,f} Anthocyanins : [0.66],Vi ^f Coumarins : [0.08 ; 0.12 ; 0.22 ; 0.26],J ^C ; [0.71],P ^C ; [0.82],J ^C ; [0.9],Ve ^C Tannins : [0.7],Gr ^g
	Noy	Flavonoïds : [0.12],Ve ^f ; [0.19],J ^b , P ^f ; [0.29],J ^b , P ^f ; [0.36],P ^f ; [0.4],P ^{b,f} ; [0.46],J ^b ; [0.54],P ^f ; [0.61 ; 0.79],J ^b ; [0.69],Ve ^f Anthocyanins : [0.19 ; 0.36 ; 0.42],Vi ^f Coumarins : [0.06 ; 0.15 ; 0.29 ; 0.4],P ^C ; [0.5],Ve ^C ; [0.6],Gr ^C ; [0.7],Ve ^C ; [0.81],J ^C Tannins : [0.11 ; 0.15],Ro ^g ; [0.24 ; 0.42],Gr ^g
	Coq	Flavonoïds : [0.07 ; 0.17],J ^f ; [0.1],J ^b ; [0.19 ; 0.27],Ve ^b ; [0.22],B ^f ; [0.34],Ve ^{b,f} ; [0.42],J ^b ; [0.56],B ^b ; [0.49 ; 0.62],J ^f ; [0.6],Ve ^b ; [0.66 ; 0.75],P ^f ; [0.7],J ^b ; [0.79],B ^b ; [0.86],B ^f Coumarins : [0.14 ; 0.19 ; 0.22],J ^C ; [0.31],Ve ^C ; [0.41 ; 0.51 ; 0.6],J ^C ; [0.69 ; 0.71],P ^C ; [0.8 ; 0.9],J ^C Anthocyanins : [0.22 ; 0.86],B ^f Tannins : [0.08 ; 0.19 ; 0.22 ; 0.3 ; 0.71],Gr ^g ; [0.29 ; 0.36],Ro ^g ; [0.4],B ^g ; [0.5 ; 0.6]Ro ^g
n-butanol	Pul	Flavonoïds : [0.1],B ^{b,f} ; [0.15 ; 0.2],Ve ^b ; [0.6],B ^b ; [0.75],J ^f ; [0.89],Vi ^f Anthocyanins : [0.12],B ⁱ ; [0.89],Vi ^f Coumarins : [0.07 ; 0.29 ; 0.4],Ve ^C ; [0.2 ; 0.47 ; 0.62],Be ^C
	Noy	Flavonoïds : [0.06 ; 0.08 ; 0.14],P ^f ; [0.17],J ^b ; [0.19],Ve ^f ; [0.27],P ^f ; [0.3],J ^b ; [0.41],Ve ^b , J ^f ; [0.49],J ^b ; [0.5],Ve ^f ; [0.6],J ^f Anthocyanins : [0.06 ; 0.08 ; 0.14 ; 0.27],Vi ^f Coumarins : [0.08],P ^f ; [0.1],J ^{C,f} ; [0.14],P ^f ; [0.2],J ^C , Ve ^f ; [0.27],J ^C , P ^f ; [0.4],Ve ^f , J-Ve ^f ; [0.47],J ^C ; [0.61],Ve ^C ; [0.81],J ^f Tannins : [0.27 ; 0.4],Gr ^g ; [0.47],B ^g ; [0.55],Or ^g
	Coq	Flavonoïds : [0.06],J ^b , B ^f ; [0.12],B ^{b,f} ; [0.16],Ve ^{b,f} ; [0.2],B ^{b,f} ; [0.31 ; 0.5],J ^f ; [0.25 ; 0.31 ; 0.45 ; 0.59],Ve ^b ; [0.67],J ^b ; [0.75],Ve ^{b,f} ; [0.8],B ^b Anthocyanins : [0.07 ; 0.12 ; 0.21],B ^f Coumarins : [0.07],Ve ^C ; [0.2],B ^C ; [0.29 ; 0.4],Ve ^C ; [0.47 ; 0.62],B ^C ; [0.76],J ^C Tannins : [0.16 ; 0.25],Gr ^g ; [0.69],Be ^g

210 b: compounds developed with aluminium chloride (AlCl₃); c: compounds developed with 5% potassium
211 hydroxide (KOH); f: compounds developed with ammonia vapour (NH₃); g: compounds developed with 2%
212 iron chloride (FeCl₃).
213 B: blue, Gr: grey, J: yellow, Or: orange, P: purple, Ro: pink, Ve: green, Vi: violet or purplish.
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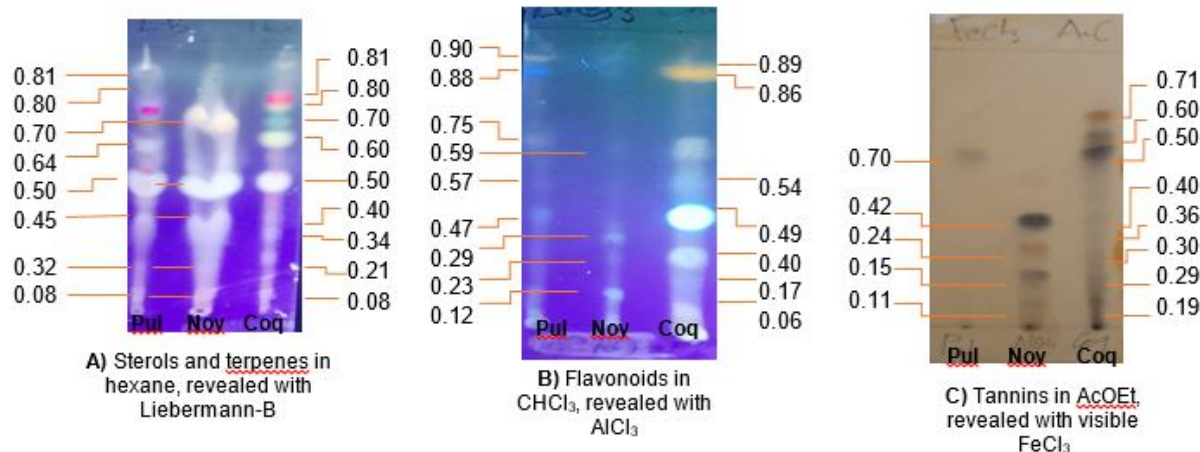


Figure 2: Some chromatographic profiles of the different parts of the *D. guineense* fruit.

215
 216
 217
 218
 219 The presence of various phytochemicals was discovered through phytochemical screening,
 220 such as sterols, terpenes, flavonoids, anthocyanins, coumarins and tannins, in the various parts
 221 of the *D. guineense* fruit. These results show the wide diversity of bioactive compounds present
 222 in this fruit. The presence of flavonoids, anthocyanins, coumarins, and tannins has been
 223 observed in other *Dialium* species, including *D. gossweileri* and *D. corbisieri*[34].

224 **3.3. Quantitative analysis**

225 **3.3.1. Total phenol content**

226 The levels of total phenols in the different parts of the *D. guineense* fruit (pulp, kernel and shell)
 227 (Figure 3) were measured using a calibration curve with gallic acid, the equation of which is $y =$
 228 $0.0232x + 0.0002$.

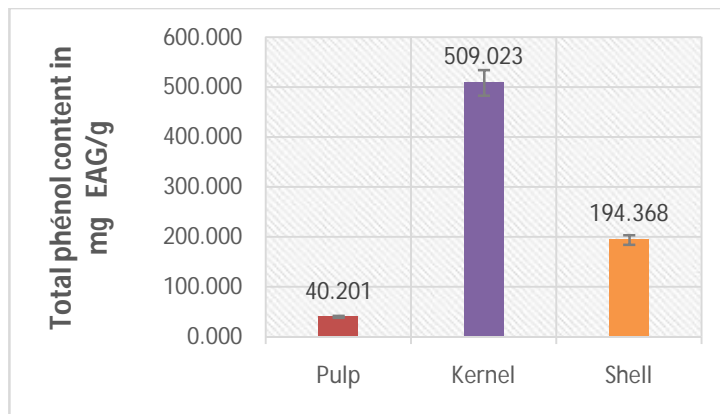


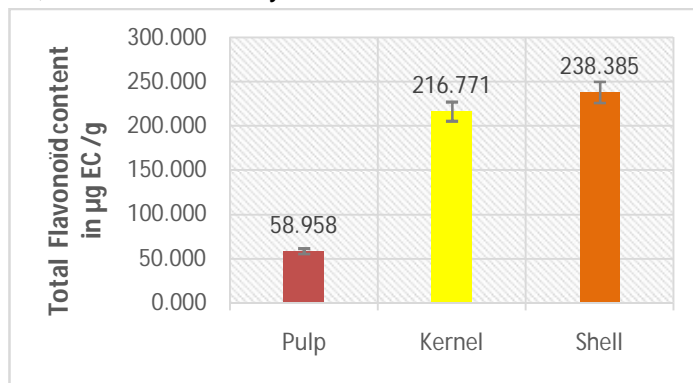
Figure 3: Total phenol content of the different parts of *D. guineense* fruit

229
 230
 231 The concentrations of phenols obtained varied from 40.201 ± 0.718 to $509.022 \pm 5.912 \mu\text{g}$
 232 EAG/g in samples. The kernel ($509.022 \pm 5.912 \mu\text{g}$ EAG/g) had the highest content, while the
 233 pulp showed the lowest content ($40.201 \pm 0.718 \mu\text{g}$ EAG/g). The total phenol content of the
 234 different parts of the *D. Guineense* fruit was significantly different ($p < 0.05$) in the analysis of
 235 variance at the 5% level. Due to the variability of the variances, the Tamhane test was used as a
 236 post-hoc test. This indicated that the total phenol content of the different parts of the fruit differed

237 significantly from one organ to another. Consequently, they are not comparable. This
238 observation would suggest an uneven distribution of natural compounds in the different organs
239 and physiological tissues of plants. This phenomenon has also been observed by other authors
240 [22, 35, 36]. In addition, the phenol content of the kernel is 12.7 times higher than that of the
241 pulp and around 2.6 times higher than that of the shell. This high concentration of phenols in the
242 kernel suggests that it could be considered a reservoir of phenolic compounds, which could be
243 attributed to the presence of flavonoids, coumarins and tannins detected during phytochemical
244 screening. Similar studies conducted on the different parts of the fruit of *Tamarindus indica*
245 (Fabaceae) revealed levels of $152 \pm 2.2 \mu\text{g EAG/g}$ for the pulp, $94 \pm 2.1 \mu\text{g EAG/g}$ for the shell
246 and $66 \pm 0.7 \mu\text{g EAG/g}$ for the kernel [37]. These results contrast with those obtained for *D.*
247 *guineensis*, where the tamarind pulp contains more phenolic compounds than the shell, which is itself
248 richer than the kernel. These observations suggest that there is an unequal distribution of
249 phenolic compounds in the organs of a plant species, despite their belonging to the same family.

250 3.3.2. Total flavonoid content

251 The figure 4 shows that the flavonoid levels in different parts of the *D. Guineensis* fruit varied
252 from 58.958 ± 2.476 to $238.385 \pm 2.533 \text{ EC/g}$ dry matter. The catechin calibration curve was
253 used to obtain the latter, with the formula $y = 0.0004x - 0.0026$.



254

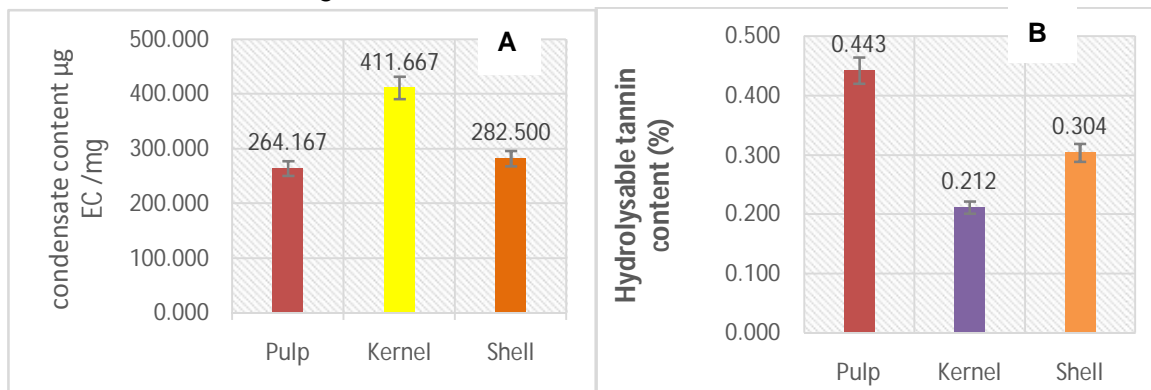
255

Figure 4: Total flavonoid content of different parts of *D. guineensis* fruit

256 The total flavonoid content of different parts of *D. Guineensis* fruit was significantly different ($p <$
257 0.05) according to analysis of variance at the 5% level. According to the Tamhane test, the
258 flavonoids content in the shell ($238.385 \pm 2.533 \mu\text{g EC/g}$) and kernel ($217.71 \pm 8.863 \mu\text{g EC/g}$)
259 of *D. Guineensis* is comparable. *D. Guineensis* shell and kernel may have similar
260 pharmacological properties, including antioxidant potential. Flavonoids are recognized in the
261 literature as powerful antioxidants, beneficial for the body [38]. It would therefore be wise to
262 consume all parts of the fruit in order to maximize its many benefits. A Comparison of the
263 proportions of total flavonoids of black tamarind and *Tamarindus indica* reveals that the latter
264 contains fewer flavonoids (pulp ($24 \pm 1.4 \mu\text{g EC/g}$); shell ($27 \pm 1.0 \mu\text{g EC/g}$) and kernel (21 ± 0.7
265 $\mu\text{g EC/g}$)) [37]. Furthermore, Pearson's test indicated a mean positive correlation ($r = 0.672$)
266 between total phenol and total flavonoid levels. This could indicate an average contribution of
267 flavonoids to the estimation of phenolic compound levels.

268 **3.3.3. Condensed and hydrolysable tannin content**

269 The catechin calibration line ($y = 0.004x + 0.006$) was utilized to calculate condensed tannin
270 levels, and hydrolysable tannin levels were measured using a 660 nm spectrometer. The results
271 obtained are shown in figure 5.



272
273 **Figure 5: Condensed (A) and hydrolysable (B) tannin content of the various parts of the**
274 **fruit of *D. guineense***

275 The percentage of condensed tannins in the different parts of *D. Guineense* fruit was
276 significantly different ($p < 0.05$) in the analysis of variance at the 5% level. The Tamhane test
277 revealed that the shell ($282.5 \pm 2.141 \mu\text{g EC/g}$) and pulp ($264.167 \pm 5.689 \mu\text{g EC/g}$) of *D.*
278 *guineense* have comparable condensed tannin contents (Figure 5A) but are different from that of
279 the kernel. The kernel ($411.667 \pm 10.541 \mu\text{g EC/mg}$) thus remains the part richest in condensed
280 tannins. In addition, a strong positive correlation ($r = 0.942$) was observed between condensed
281 tannin content and total phenol content, indicating the significant contribution of condensed
282 tannins to the quantification of phenolic compounds. In general, the analysis of variance applied
283 to the hydrolysable tannin content showed a significant difference ($p < 0.05$) between the
284 different averages. The Tukey test showed that the hydrolysable tannin contents of the different
285 parts were not comparable (Figure 5B). The presence of tannins in the fruit of *D. guineense* fruit
286 could explain several properties attributed to it, in particular the astringent flavor felt when the
287 pulp is eaten. This sensation arises from the interaction between the tannins of the consumed
288 matrix and salivary proteins, particularly those rich in proline [39].

289 **3.4. Antioxidant activity**

290 **3.4.1. Antioxidant capacity by DPPH test**

291 The figure 6 shows the results of the DPPH test.

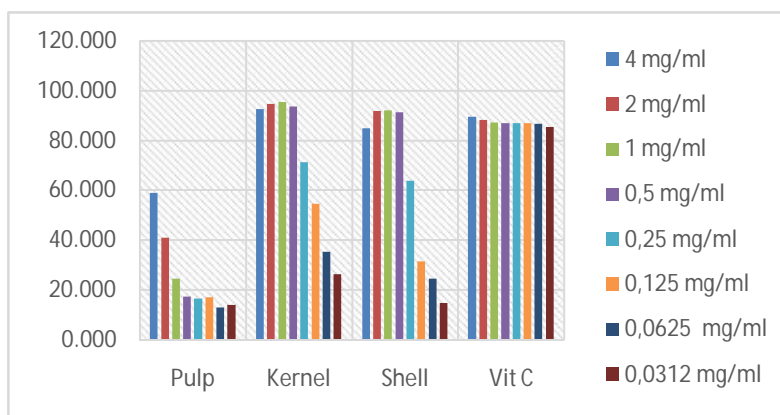


Figure 6: Antioxidant profiles of different *D. guineense* extracts

292
293
294 Overall, the extracts tested had an anti-radical effect on the DPPH. Extracts from the kernel and
295 shell showed significantly greater reduction capacity than those from the pulp. For
296 concentrations of 4 to 0.0312 mg/mL, the reduction percentages (PR) for the kernel range from
297 92.728 to 26.380% and for the shell, it ranges from 84.873 to 14.756%. They are greater than
298 50% for concentrations between 0.125 and 4 mg/mL (Figure 6). In addition, for concentrations
299 between 0.5 and 4 mg/mL, PR above 90% were observed for most extracts, except for the shell
300 at 4 mg/mL with a PR of 84.873%. These results suggest that the kernel and shell are potential
301 sources of antioxidant compounds. For the pulp, the highest PR (59.023%) was observed at 4
302 mg/mL, indicating a reduced capacity to neutralize DPPH compared to other organs. The Table
303 IV presents the different CR₅₀ values, designating the concentrations that cause the loss of 50%
304 of the DPPH. These values were determined graphically.

Table IV: CR₅₀ (mg/mL) of crude extracts of *D. guineense* and vitamin C

Extract	Pulp	Kernel	Shell	Vitamin C
CR ₅₀	3.102	0.110	0.193	0.008

306
307 According to the literature, the lower the CR₅₀, the greater the antioxidant activity and vice versa.
308 On the basis of this principle, it can be stated that the kernel extract has a greater antioxidant
309 capacity, followed by the shell. On the other hand, compared to vitamin C, the different parts of
310 *D. guineense* exhibit much lower activity. The antioxidant properties of the kernel and shell of *D.*
311 *guineense* are much stronger than those of the fruits of *D. gossweileri*. (CR₅₀ = 500 µg/mL).
312 Conversely, their activity is less than that of *D. corbisieri*(CR₅₀ = 14.44 µg/mL) [34].
313 Consumption of these three parts of the fruit could therefore be beneficial for the body.

3.4.2. Antioxidant activity by FRAP test

314
315 The concentrations of ferric ions (Fe³⁺) to ferrous ions (Fe²⁺) were varied from 2115.91 to
316 51101.52 µg EFS/g DM in the different parts of *D. Guineense* fruit. (Figure 7). These values
317 were obtained after projection onto the calibration line of equation $y = 0.0197x - 0.0187$ with $R^2 =$
318 0.9991, performed with iron (II) sulfate, prepared under the same conditions as the samples.

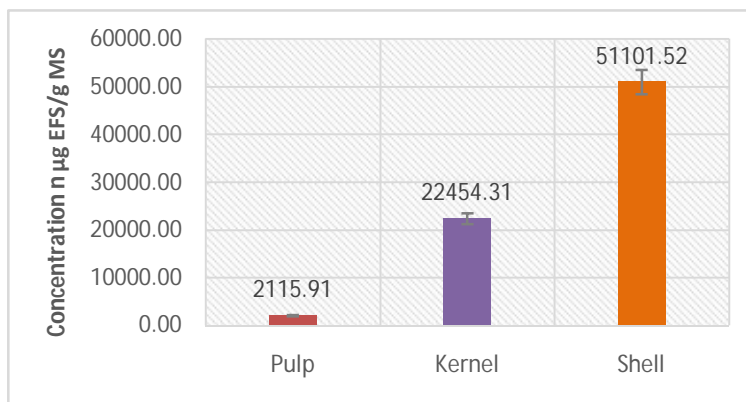


Figure 7: Reduction concentrations of Fe³⁺ to Fe²⁺ in the FRAP test

319
 320
 321
 322 The shell signed the highest reduction concentration (51101.52 µg EFS/g DM), while the pulp
 323 presented the lowest (2115.91 µg EFS/g DM). This could be explained by the presence of
 324 phytochemicals such as flavonoids in the shell of *D. Guineense*, which are capable of providing
 325 electrons to reduce the ferric ion. This hypothesis was confirmed by a strong positive correlation
 326 ($r = 0.999$) between the FRAP test and the total flavonoid content ($238.385 \pm 6.2 \mu\text{g EC/g}$). [40]
 327 report that most flavonoids possess the ferric ion's reducing properties. However, those which
 328 have in their structure, the O-dihydroxy group in the ring B and the 3-hydroxy group and a
 329 double bond at the 2,3 positions in the ring C are the most active during the FRAP test [40]. The
 330 pulp was the least active part of *D. Guineense* fruit, regardless of the method used to evaluate
 331 its antioxidant potential. Consequently, incorporating the other parts of the fruit (shell and kernel)
 332 into dietary habits and other uses could be beneficial.

333 4. CONCLUSION

334 This study focused on improving the value of *D. Guineense* fruit from Côte d'Ivoire by analyzing
 335 its chemical composition and evaluating the antioxidant potential of its various components
 336 (shell, pulp, and kernel). Phytochemical screening revealed the coexistence of several
 337 phytoconstituents (flavonoids, coumarins, tannins, sterols, terpenes, alkaloids and saponins) and
 338 the absence of cardiotoxic glycosides in all parts of the fruit. Furthermore, quinones and
 339 alkaloids were not visible in the pulp. Quantitative analysis revealed significant levels of total
 340 phenols ($509.023 \pm 14.5 \mu\text{g EAG/g}$) and condensed tannins ($411.667 \pm 8.7 \mu\text{g EC/mg}$) in the
 341 kernel. Flavonoids are most abundant in the shell ($238.385 \pm 25.8 \mu\text{g EC/g}$). The study of
 342 antioxidant potential using DPPH and FRAP methods revealed interesting antioxidant
 343 compounds in the various parts of the *D. guineense* fruit. These compounds could help to
 344 promote *D. guineense* fruit as a health food, offering an effective response to certain chronic
 345 diseases. We therefore encourage people to consume the whole fruit of *D. guineense* to reap its
 346 full benefits.

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