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## **Analysis of the contrast between natural occurrence of toxigenic Aspergilli of the Flavi section and aflatoxin B1 in cassava**

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### **► To cite this version:**

Yann Adjovi, Sylviane Bally, B. J. G. Gnonlonfin, Souria Tadrict, Arlette Querin, et al.. Analysis of the contrast between natural occurrence of toxigenic Aspergilli of the Flavi section and aflatoxin B1 in cassava. *Food Microbiology*, 2014, 38, pp.151-159. 10.1016/j.fm.2013.08.005 . hal-02632483

**HAL Id: hal-02632483**

**<https://hal.inrae.fr/hal-02632483>**

Submitted on 7 Jun 2023

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1     **Analysis of the contrast between natural occurrence of toxigenic *Aspergillii***  
2                     **of the *Flavi* section and aflatoxin B1 in cassava**

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15

16    **Key words**

17    *Aspergillus flavus*, *Aspergillus parvisclerotigenus*, *Aspergillus novoparasiticus*, cassava,  
18    *Aflatoxin B1*, inhibition, Benin

19

20

## 21 **Abstract**

22 Aflatoxin B1 (AFB1) is a carcinogenic mycotoxin produced by *Aspergillii* of the section  
23 *Flavi* that may contaminate food, in the field or during storage. Cassava represents an  
24 important staple food in sub-saharian Africa. The analysis of aflatoxigenic fungi in 36  
25 cassava samples obtained from producers in Benin indicated that 40% were  
26 contaminated by *Aspergillii* of the section *Flavi*. Upon morphological and molecular  
27 characterization of the 20 isolates, 16 belonged to *A. flavus*, 2 to *A. parvisclerotigenus* and  
28 2 to *A. novoparasiticus*. This is the first time that this latter species is isolated from food.  
29 Although most of these isolates were toxigenic on synthetic media, no AFB1  
30 contamination was observed in these cassava samples. In order to determine the action  
31 of cassava on AFB1 synthesis, a highly toxigenic strain of *A. flavus*, was inoculated onto  
32 fresh cassava and despite a rapid development, no AFB1 was produced. The anti-  
33 aflatoxin property was observed with cassava from different geographical origins and on  
34 other aflatoxigenic strains of the section *Flavi*, but it was lost after heating, sun drying  
35 and freezing.

36 Our data suggest that fresh cassava is safe regarding AFB1 contamination, however,  
37 processing may alter its ability to block toxinogenesis leading to secondary  
38 contamination.

## 39 1. Introduction

40 Mycotoxins are toxic secondary metabolites produced by fungi and are common  
41 contaminants of food and feed commodities worldwide. Over 400 mycotoxins are  
42 known today; due to their occurrence and toxicity, aflatoxins, fumonisins, ochratoxin,  
43 zearalenone and trichotecenes are usually considered as the most important for human  
44 health (Leslie et al., 2008).

45 The most toxic and dangerous mycotoxins are aflatoxins. Indeed, aflatoxin B1 (AFB1) is  
46 the most potent hepatic carcinogen known in mammals and has been classified by the  
47 International Agency for Cancer Research in the group I of molecules that are  
48 carcinogenic for both humans and animals (IARC, 1993). AFB1 also possesses  
49 immunosuppressive properties (Meissonnier et al., 2008) and is involved in growth  
50 impairment observed in children (Gong et al., 2004; Khlangwiset et al., 2011). Exposure  
51 to aflatoxins in sub-Saharan Africa is very frequent. In some areas, 99% of children  
52 tested have aflatoxins in their blood (Gong et al., 2002) and this high exposure  
53 contributes to appearance of chronic hepatomegaly in children (Gong et al. 2012).  
54 Contamination of food by very high levels of aflatoxins can lead to fatal consequences  
55 such as the reported death of 125 people in Kenya (Lewis et al., 2005; Probst et al.,  
56 2007). The contamination of food by AFB1 may therefore have important consequences  
57 for human health and more than 100 countries have established maximal tolerable  
58 limits for this molecule (FAO, 2003; EU, 2006).

59 In the tropical countries highly exposed to aflatoxin, cassava (*Manihot esculenta Crantz*)  
60 belongs to the short list of most consumed foodstuffs. It is ranked as the third most  
61 important food crop in tropical regions after rice and maize. Due to its high energetic  
62 value, it represents an important part of the diet of almost one billion people, mostly in  
63 sub-Saharan Africa. In Africa, cassava is mostly devoted to human food, livestock  
64 feedstuff, but it is also used as a raw material in various industries that produce flour,  
65 starch, adhesives, etc. As human food, cassava may be consumed as fresh roots usually  
66 as a snack, but also after being peeled, grated or soak in water to ferment and then  
67 processed into a wet mash (fufu) or varied dry products (gari, chips/crumbs/chunks, or  
68 milled into flour) (Akoroda, 2007). The conditions of production and storage of cassava  
69 and its traditional derivatives may be favorable to the contamination and development  
70 of molds (Westby and Twiddy, 1991). Generally such uncontrolled mold developments

71 may not only result in alterations in the organoleptic properties of the foodstuffs but  
72 may also lead to the accumulation of toxic secondary metabolites.

73 Due to its nutritional and economic importance in many tropical countries, it is  
74 necessary to assess the contamination of these foods by mycotoxins. Previous surveys  
75 performed in order to evaluate fungal and mycotoxin contaminations showed that some  
76 mycotoxins such as patulin, cyclopiazonic, ochratoxin A, zearalenone could be found in  
77 cassava (Wareing et al., 2001). However, no trace of aflatoxins has been mentioned,  
78 although the warm and humid climate that prevails in tropical areas can promote the  
79 development of aflatoxin-producing *Aspergillus* species (Muzanila et al, 2000; Jimoh and  
80 Kolapo, 2008). Recently, we did a preliminary survey on cassava chips sold in Beninese  
81 markets and found no aflatoxin contamination of cassava chips, despite a high presence  
82 of *A. flavus* spores (Gnonlonfin et al., 2012).

83 Within this context, the objective of this current study was to understand the absence of  
84 AFB1 in cassava products. Consequently, we firstly did a small survey to analyze fungal  
85 flora and aflatoxins in cassava chip samples taken directly from producers in order to  
86 evaluate the fungal contamination at this critical point in the food chain. We then  
87 characterized the isolates of *Aspergillus* of the section *Flavi*, isolated during this survey  
88 and the previous one, regarding their toxigenic potential. Finally, we investigated the  
89 action of cassava on AFB1 synthesis and demonstrated that fresh cassava was able to  
90 block AFB1 synthesis by toxigenic strains of *A. flavus*.

91

## 92 **2. Material and methods**

93

### 94 **2.1 Chemicals and reagents**

95

96 All reagents including solvents were purchased from Fisher Scientific (Fontenay sous  
97 Bois, France) and were of analytical grade. Aflatoxin B1 (AFB1), aflatoxin G1 (AFG1),  
98 scopoletin and ergosterol standards were purchased from Sigma-Aldrich (Saint-Quentin  
99 Fallavier, France). AFB1 and scopoletin were dissolved in methanol and ergosterol in  
100 ethanol. Taq DNA polymerase used for molecular identification was purchased from  
101 Invitrogen (Carlsbad, CA, USA). Agarose gel and primers used for gene determination  
102 were purchased from Eurobio (Courtaboeuf, France).

103

### 104 **2.2 Cassava samples**

105

106 For the survey, thirty six samples of cassava of about 1kg each were purchased from  
107 local producers in the north of Benin during the dry season (December 2010) (Figure 1).  
108 Samples were randomly chosen and checked for the absence of macroscopically visible  
109 alteration or dryness.

110 For the experiments on *Aspergillus* development and toxinogenesis, two varieties of  
111 cassava were used. The first one, named *Logoguesse Kotorou* in Bariba dialect was  
112 grown in the north of Benin. The second variety, grown in Costa Rica, was purchased on  
113 the market in Toulouse, France. These roots were protected from contamination and  
114 alteration by a thin pellicle of wax.

115 These two varieties were intended for human consumption and corresponded to sweet  
116 varieties with concentrations of cyanogenic compounds below 50 mg/Kg as evaluated  
117 according to Essers et al. (1993).

118

### 119 **2.3 Isolation and Identification of fungi**

120

#### 121 *2.3.1 Origin*

122 Fungal analysis of cassava samples was done according to Gnolongin et al. (2012). In  
123 brief, 1 ml of decimal dilutions were plated on Petri dishes containing dichloran glycerol  
124 agar (DG18). Petri dishes were incubated at 25°C in alternating 12h periods of

125 fluorescence light and dark for 5 days. Colony were isolated by several plating out and  
126 further identified as described below.

127 Twelve isolates coming from a survey done in 2009-2010 on cassava chips collected in  
128 markets in Benin were added to the study (Gnonlonfin et al., 2012). Other strains of  
129 *Aspergillus* of the section *Flavi* used in this study were obtained from the Agricultural  
130 Research Service collection (NRRL) and CBS-KNAW Fungal Biodiversity Centre collection  
131 (CBS): *A. flavus* NRRL 1957, *A. parasiticus* NRRL 5862, *A. parasiticus* NRRL 4123, *A.*  
132 *tamarii* NRRL 20818, *A. bombycis* NRRL 26010, *A. pseudatomarii* NRRL 25517,  
133 *Aspergillus parvisclerotigenus* CBS 121.62, *Aspergillus minisclerotigenes* NRRL 29000,  
134 *Aspergillus minisclerotigenes* CBS117635.

135

### 136 2.3.2 Morphological and molecular identification

137 The 32 strains isolated from cassava were cultivated on Potato Dextrose Agar (PDA) (24  
138 g/l potato-dextrose broth, 15% (w/v) technical agar (Becton, Dickinson and Co.,  
139 Franklin Lakes, NJ, USA)) at 25°C for seven days. They were identified as *Aspergillus*  
140 section *Flavi* isolates by both macroscopic and microscopic examination according to  
141 Pitt and Hocking (2009) and Varga et al. (2011).

142 Identification of *Aspergillus flavus* isolates was also performed by PCR amplification of  
143 the *cypA/norB* region as described in Probst et al. (2012). The culture conditions and  
144 genomic DNA isolation were described by El Mahgubi et al. (2013). PCR reactions were  
145 carried out in a GenAmp PCR 2700 thermocycler (Applied Biosystems, Forster City, CA,  
146 USA). This identification was completed for 10 randomly selected isolates which showed  
147 0.3 or 0.9 kb fragments by internal transcribed spacers (ITS) rRNA and 5.8S rRNA gene  
148 amplification and sequencing. For the other isolates displaying no gap, added to ITS, the  
149 molecular identification was performed by calmodulin, beta-tubulin or acetamidase  
150 (*Amds12*) and o-methyltransferase (*Omt12*) genes amplification and sequencing.  
151 Primers used in this study are listed in Table 1. For ITS amplification, PCR conditions  
152 were those described by White et al. (1990). Beta-tubulin, calmodulin were amplified  
153 under conditions previously described by Peterson (2008) whereas (*Amds12*) and o-  
154 methyltransferase (*Omt12*) genes were amplified according to Geiser et al. (1998).

155 PCR products were purified with GenElute PCR clean-up kit (Sigma-Aldrich) and  
156 sequenced by the dye terminator technology on an ABI3130XL sequencer (Applied  
157 Biosystems). PCR products were sequenced in both directions.

158 Nucleotide sequence accession numbers for internal transcribed spacers (ITS) rRNA and  
159 5.8S rRNA, beta-tubulin, calmodulin, acetamidase and O-methyltransferase of tested  
160 isolates are reported to Table 2.

161

### 162 2.3.3 Analysis of toxin pathway genes

163 The toxigenic status of the *A. flavus* isolates was determined by analyzing the presence  
164 of six genes involved in the aflatoxin biosynthesis pathway: *AflD* (*Nor-1*), *AflO* (*OmtB*),  
165 *AflP* (*OmtA*), *AflQ* (*OrdA*), *AflR*, and *AflS*. The primers used were designed using the  
166 sequences of the *A. flavus* strain NRRL 3357 deposited in GenBank and are reported in  
167 Table 1. The PCR amplification was carried out in 50  $\mu$ L reaction mixture (5  $\mu$ L of Taq  
168 polymerase buffer 10X, 1.5  $\mu$ L of 50 mM  $MgCl_2$ , 1  $\mu$ L of 10 mM dNTP (Promega, Madison,  
169 WI, USA), 1.5  $\mu$ M of each primer, 1.5 units of Taq DNA polymerase (Invitrogen, Carlsbad,  
170 CA, USA), 5  $\mu$ L of genomic DNA (100 ng/ $\mu$ L), and 31  $\mu$ L of pure water (Laboratoire  
171 Aguetant, Lyon, France)). Reaction conditions were: 94°C for 5 min, 40 cycles (1 min at  
172 94°C, 1 min at 58°C, and 1 min at 72°C) followed by 10 min at 72°C. The amplified  
173 products were examined by 1.2% w/v agarose gel electrophoresis.

174

## 175 2.4 Aflatoxin B1 quantification

176

### 177 2.4.1 In cassava chip samples

178 AFB1 quantification in cassava chip samples was done on site according to the method  
179 developed and validated by Gnonlonfin et al. (2010). Briefly, each sample (10 g) was  
180 extracted with 1 g of sodium chloride and 25 mL of extraction solution [methanol/water  
181 (80/20, v/v)] and purified on an Aflatest®immunoaffinity column (VICAM, Watertown,  
182 MA, USA). The extract was dried under nitrogen gas at 60°C, dissolved in 200  $\mu$ L of  
183 methanol and analyzed by HPLC with a post-column derivatization using a  
184 photochemical reactor for enhanced detection (PHRED). The limit of detection (LOD) of  
185 the method was 0.1  $\mu$ g/kg.

186

### 187 2.4.2 In culture medium

188 AFB1 production by *A. flavus* strains was determined after culture on PDA for 7 days at  
189 25°C. Toxins were extracted from culture medium by mechanical agitation with 70 ml of  
190 chloroform. The organic phase was evaporated in a rotary evaporator at 50°C. The



191 residue was taken up in 400 µl of methanol, and this suspension was filtered-through a  
192 0.45 mm pore-size filter before analysis.

193 The mycotoxin was quantified by high performance liquid chromatography with  
194 photodiode array detector (Thermo-Finnigan Suveyor HPLC-PDA system,  
195 ThermoElectron Corporation, Waltham, MA, USA). The liquid chromatography  
196 separation was performed using a 150 mm x 2.00 mm Luna 5 µm C18 column  
197 (Phenomenex, Torrance, CA, USA). HPLC was done with a linear gradient elution using  
198 33 mM acetic acid (solvent A) and acetonitrile (solvent B) with a flow rate of 200 µl/min.  
199 The compounds were eluted starting from 80% of solvent A for 5 min, followed by a  
200 50% solvent B step, then a linear gradient up to 90% solvent B within 5 min. After  
201 isocratic elution for 5 min, the gradient was decreased to its initial value within 2 min,  
202 and maintained at it for the last 5 min.

203 Calibration curves for AFB1 were constructed using standard solutions in methanol  
204 (from 0.1 to 100 µg/ml) at 365nm. The LOD was 2 ng/ml.

205

## 206 **2.5** *A. flavus* growth and toxinogenesis on cassava

207

### 208 *2.5.1 Culture conditions*

#### 209 *Inoculum preparation*

210 Strains were cultured on PDA at 25°C in the dark for 7 days. After this period, the plates  
211 were washed with 10 ml of 0.05% Tween and a spore suspension was harvested and  
212 adjusted to 10<sup>5</sup> spores/ml after counting in a Malassez cell.

#### 213 *Culture on cassava*

214 Before inoculation, the water activity of cassava was measured with an HC2-AW device  
215 (Rotronik AG, Basserdorf, Switzerland) and adjusted to 0.98 by addition of the required  
216 quantity of sterile water.

217 After peeling, cassava was cut into thin slices and inoculated immediately (fresh) or  
218 after heat treatment (120°C for 20 min), freezing (-20°C for 2 months) or drying. For this  
219 later treatment, cassava chips were processed as previously described (Gnonlonfin et al.,  
220 2010). Everyday, the evolution of the water content was followed according to AOAC  
221 934.01 (AOAC, 2000) and 200 g of cassava were used and inoculated as follow.

222 Fifteen grams of fresh or treated cassava were inoculated with 100 µl of a spore  
223 suspension ( $10^4$  spores) and incubated at 25°C for 3 and 6 days in dark and high  
224 humidity conditions. Each experiment was done in triplicate.

225

### 226 *2.5.2 Fungal development measurement*

#### 227 *Conidia formation:*

228 Spore numbers were determined after grinding cultures in phosphate buffer [(0.1M,  
229 pH7.4); 50 ml for 15 g cassava]. Ten-fold dilutions were prepared with tween 0.05% and  
230 the numeration of spores was done both by counting in a Malassez cell and by plating  
231 dilutions on malt agar and determining Colony Forming Units per gram (CFU/g) after 5  
232 days at 25°C.

#### 233 *Ergosterol concentration*

234 Ergosterol measurement was done according to Bailly et al. (1999). In brief, after  
235 saponification in methanol, ethanol, KOH and pyrogallol and extraction with petroleum  
236 ether, ergosterol was quantified by fluorodensitometry with a Shimadzu CS930  
237 densitometer (Shimadzu, Kyoto, Japan) after separation of the extracts by thin layer  
238 chromatography on silica plates (Macherey-Nagel, Düren, Germany) in toluene-  
239 acetonitrile (70:30). Quantification was done by comparison with known amounts of  
240 standard spotted on the same plate.

241

### 242 *2.5.3 Aflatoxin B1 measurement in cassava*

243 The aflatoxin B1 was extracted from cassava with chloroform (50 mg/ml) and filtered-  
244 through a phase separator PS Whatman filter (GE Healthcare Biosciences, Pittsburgh,  
245 PA, USA). The extract was evaporated to dryness in a TurboVap Lv Evaporator (Zymark,  
246 Hopkinton, MA, USA) under N<sub>2</sub> atmosphere. Dry extracts were re-suspended and  
247 analyzed by HPLC-PDA as described above.

248 For spiking experiments, pure AFB1 was added on fresh or autoclaved cassava (5 ng/g  
249 cassava). Extraction and quantification of AFB1 was then done as described above.

250

## 251 **2.6 Statistical analysis**

252

253 Differences for ergosterol production and sporulation were evaluated for statistical  
254 significance using an analysis of variance (two-way ANOVA).  $P \leq 0.05$  was required for  
255 significance.

256

### 257 **3. Results**

258

#### 259 **3.1 Contamination of cassava chips with *Aspergillus* section *Flavi***

260

261 A survey was done to evaluate contamination with *Aspergilli* and the presence of AFB1  
262 on 36 samples randomly collected from producers located in the North of Benin (Figure  
263 1).

264

##### 265 *3.1.1 Morphological and molecular characterization of fungal isolates*

266 Identification of fungal isolates from cassava samples was firstly done by morphologic  
267 examination after growth on PDA medium. Nearly forty percent of samples collected  
268 (38.9%) were found to be contaminated with strains that were morphologically  
269 identified as *Aspergillus* of the section *Flavi* and a total of 20 different strains were  
270 isolated, two samples being contaminated with 2 different strains. The presence of  
271 sclerotes was observed in half of the strains.

272 Identification of these isolates to the species level was performed using molecular  
273 methodologies. Twelve additional isolates previously isolated from marketed cassava  
274 chips (Gnolonfin et al., 2012) were also submitted to such identification. As a first step,  
275 identification of these isolates to the species level was performed by the determination  
276 of the presence of a gap in *cypA/norB* region. The amplification of *cypA/norB* of 28  
277 isolates revealed 0.9 or 1.5 kb deletion. Among the 32 isolates, 18 displayed 0.9 kb gap  
278 and 10 displayed a 1.5 kb deletion (Table 3). The presence of these deletions strongly  
279 suggested that the isolates belonged to *Aspergillus flavus* species. Indeed, the  
280 cytochrome P450 monooxygenase encoded by *cypA* is required for aflatoxins G<sub>1</sub> and G<sub>2</sub>  
281 production and the partial deletions of *cypA* explain that *A. flavus* lacks capacity to  
282 synthesize aflatoxins G (Ehrlich et al., 2004). Molecular identification based on ITS

283 sequences was also performed on 10 strains randomly selected from these isolates. This  
284 confirmed that all belonged to *A. flavus* species (Table 2).

285 The amplification of *cypA/norB* region (1.8 kb) of the 4 remaining isolates (AFc31,  
286 AFc32, AFc35 and AFc36) revealed an absence of gap. For these isolates, the alignment  
287 of *cypA/norB* region sequences (KC990464-KC990468) to the full length sequence of  
288 *Aspergillus parasiticus* NRRL 5862 (SRRC143) strain suggested that they could be able to  
289 produce aflatoxins G.

290 For AFc31 and AFc32 isolates, the sequences obtained after ITS sequencing matched  
291 100% with those of *A. novoparasiticus* species. In order to confirm that these isolates  
292 belong to *A. novoparasiticus*, fragments of *AmdS12* and *Omt12* genes were amplified and  
293 sequenced. The obtained sequences (Table 2) matched 100% with the *A. novoparasiticus*  
294 sequences previously deposited in GenBank (Gonçalves et al., 2011).

295 AFc35 and AFc36 share identical ITS sequences with *A. flavus*. Since ITS sequence is not  
296 discriminatory to differentiate *A. flavus* from *A. parvisclerotigenus*, the sequences of  
297 parts of beta-tubulin and calmodulin gene were determined and displayed 100%  
298 identity with those of *A. parvisclerotigenus* (Table 2).

299

### 300 3.1.2 Toxin pathway gene profile of fungal strains isolated from cassava

301 To evaluate the toxigenic potential of *A. flavus* isolates, the presence of 6 different genes  
302 (*AflD*, *AflO*, *AflP*, *AflQ*, *AflR* and *AflS*) involved in the aflatoxin biosynthetic pathway was  
303 investigated by PCR. All the strains possessed these six genes in their genome (Table 3).

304 The ability of the strains to produce mycotoxins was then confirmed by AFB1 and AFG1  
305 analysis after culturing in conditions known to favor mycotoxin synthesis. Table 3 shows  
306 that most of the isolates were able to produce AFB1. Although the biosynthetic pathway  
307 genes detection was positive, the culture extracts from two strains do not showed any  
308 AFB1 trace. Four isolates identified as *A. novoparasiticus* (AFc31 and AFc32) and *A.*  
309 *parvisclerotigenus* (AFc35 and AFc36) were also found capable to produce AFG1. These  
310 results are in agreement with results of sequences analysis of *cypA/norB* region  
311 reported above.

312

### 313 3.1.3 Detection of aflatoxin B1 in cassava chips

314 As shown in Table 2, despite the aflatoxigenic character of the *A. flavus* strains isolated  
315 from cassava chips collected in Benin, no trace of AFB1 was detected neither in the chips  
316 collected in the present survey nor in the samples obtained from the previous survey.

317

### 318 **3.2 Effect of cassava on *A. flavus* growth and toxinogenesis**

319

320 In order to understand the absence of contamination of cassava by AFB1 despite the  
321 presence of toxigenic strains of *A. flavus*, the ability of cassava to inhibit AFB1 synthesis  
322 was investigated.

323

#### 324 *3.2.1 Effect of fresh cassava on aflatoxinogenesis*

325 One highly toxigenic strain of *A. flavus* isolated from cassava chips (strain AFc5) was  
326 used to inoculate fresh cassava pieces of two varieties from different geographic origins.  
327 As shown on Figure 2, fungal development was macroscopically visible after 3 and 6  
328 days of incubation, whatever the variety of cassava was used. The measurement of the  
329 ergosterol content in the cultures and spore formation confirmed that the *A. flavus* strain  
330 AFc5 was able to develop on both cassava varieties (Table 4).

331 As expected from the survey data, the fungal development and sporulation was not  
332 associated with AFB1 synthesis, regardless of the time and the cassava variety tested  
333 (Table 4). A similar inhibition of AFB1 production was observed when toxigenic strains  
334 belonging to three other *Aspergillus* section *Flavi* species (*A. parasiticus*, *A. bombycis* and  
335 *A. pseudotamarii*) were grown on cassava (Table 5), although these strains produced  
336 AFB1 when cultured on PDA medium.

337 The spiking of both fresh and autoclaved cassava with known amount of pure AFB1  
338 demonstrated that the lack of AFB1 after *A. flavus* development on cassava was not  
339 linked to a matrix interference. Indeed, 71 and 74% of the added toxin could be  
340 recovered from fresh and autoclaved spiked samples respectively.

341

#### 342 *3.2.2. Effect of cassava processing on AFB1 synthesis*

343 The impact of different processing used during the production or the storage of cassava  
344 products were then tested for their effect on AFB1 synthesis.

345 Cassava pieces were submitted to different processing before inoculation with toxigenic  
346 *A. flavus* strain AFc5. Figure 3 shows the effect of heat treatment on both mold

347 development and AFB1 synthesis. A small but significant increase in spore numbers  
348 after development on heat-treated cassava was noted (Figure 3a). By contrast, no  
349 difference was observed for ergosterol content with the exception of an increased value  
350 after development of AFc5 strain on fresh cassava from Costa Rica for 6 days (Figure  
351 3b). Regarding toxinogenesis, when the toxigenic strain was inoculated on heat-treated  
352 cassava, AFB1 was detected after 3 (data not shown) and 6 days of incubation (Figure  
353 3c) whereas no toxin was detectable after mold development on fresh cassava.

354 The effect of other process methods was also tested. Table 6 shows that sun drying of  
355 cassava pieces may also interfere with the ability of this raw material to prevent AFB1  
356 synthesis by a toxigenic strain of *A. flavus*. Indeed, after 7 days of drying, a re-moistening  
357 of cassava chips allowed AFB1 production ( $5.5 \pm 0.5$  ng/g).

358 Similarly, we observed that a 2-month freezing period at  $-20^{\circ}\text{C}$  inhibited the anti-AFB1  
359 ability of cassava, the mycotoxin becoming detectable after inoculation with the  
360 toxigenic strain AFc5 and incubation for 7 days at  $25^{\circ}\text{C}$  ( $42 \pm 18$  ng/g).

361

### 362 3.2.3. Role of scopoletin in the inhibition of AFB1 synthesis

363 Among constituents of cassava, scopoletin is known to have anti-fungal properties. Even  
364 if no antifungal activity was noted, we nevertheless investigated if this compound could  
365 explain the inhibition of AFB1 biosynthesis in fresh cassava.

366 Table 7 shows that there were no significant differences in the scopoletin concentration  
367 in heat treated cassava compared to fresh one ( $p=0.159$ ) whereas in the former  
368 condition, AFB1 was detected after *A. flavus* development. The same results were  
369 obtained using the two varieties of cassava that contained comparable concentrations of  
370 scopoletin (Table 7).

371

#### 372 4. Discussion

373

374 Cassava is a major crop in many regions of the world where it is used as human food and  
375 animal feed. The aim of this study was to assess the fungal and mycotoxin contamination  
376 that could occur after harvest and during processing of cassava.

377 In order to characterize fungal contamination of cassava at the first step of the  
378 processing chain, cassava chips were collected from several local producers in the north  
379 of Benin. Almost 40% of the samples were contaminated with *Aspergillus* of the section  
380 *Flavi* demonstrating that these fungal species may contaminate cassava before  
381 processing. Among the 20 isolates from these samples, sixteen were identified as  
382 *Aspergillus flavus*. All isolates previously collected from cassava on Benin markets were  
383 also identified as *A. flavus*. This confirmed the data published by Gnonlonfin et al. (2008;  
384 2012) which have identified *A. flavus* as a frequent contaminant of cassava chips  
385 marketed in Benin as well as other studies done elsewhere in Africa (Manjula et al.,  
386 2009; Westby et al., 1995; Essono et al., 2007). This is also in agreement with the  
387 observation that *A. flavus* is more prevalent in foods and feeds than *A. parasiticus*  
388 (Essono et al., 2009).

389 Most of the isolates produced aflatoxin B1 when cultured on PDA. This proportion of  
390 toxigenic isolates appeared quite high compared to what is usually reported. Indeed, the  
391 proportion of toxigenic *A. flavus* often ranges between 40% and 65% in raw materials  
392 such as maize (Saleemi et al., 2012; Giorni et al., 2007), cocoa beans (Sanchez-Hervas et  
393 al., 2008), groundnut (Bankole et al., 2004) or spices (El Mahgubi et al., 2013; Elshafie et  
394 al., 2002). Four isolates were found able to produce aflatoxin G<sub>1</sub>. These strains were  
395 further identified as *A. parvisclerotigenus* and *A. novoparasiticus*.

396 This last species was recently discovered and characterized (Gonçalves et al., 2012).  
397 With one exception, all strains have been isolated from patients and hospital  
398 environment samples. So far, its geographical distribution was limited to South America  
399 (Brazil and Colombia). This is the first report which mentions the presence of *Aspergillus*  
400 *novoparasiticus* in another continent other South America. It is also the first time that  
401 this species is isolated from food products.

402 Unlike *A. flavus*, *A. parvisclerotigenus* is a poorly documented species. Although the first  
403 mention regards atypical isolates from tropical soils in Thailand (Saito and Tsurata,  
404 1993) all isolates which were subjected to molecular characterization came from Guinea

405 Gulf area (Nigeria, Benin). Recently, five isolates were identified from dried edible  
406 mushrooms in Nigeria (Ezekiel et al., 2013) while the Type strain (CBS 121.62) was  
407 isolated from peanut, also in Nigeria (Frisvad et al., 2005). Blast analysis of *cypA/norB*  
408 region sequences of AFc35 and Afc36 isolates displayed a 100% identity with those of  
409 the type strain but also with those of BN008R (ATCC MYA-379) strain, an atypical *S<sub>BG</sub>*  
410 strain collected from agricultural soil in Benin (Cotty and Cardwell, 1999). *A.*  
411 *parvisclerotigenus* occurrence observed in our study is very low (<6%) as already  
412 described (Ezekiel et al., 2013).

413

414 Despite the high prevalence of contamination with toxigenic isolates, the analysis of  
415 AFB1 contamination of cassava samples revealed no trace of this toxin in the chip  
416 samples. Similar results were obtained on cassava chips sampled at the market stage in  
417 Benin (Gnonlonfin et al., 2012). This confirms that, in spite of a high occurrence of *A.*  
418 *flavus* spores, cassava chips are not contaminated with aflatoxins.

419 This observation suggests that cassava may interfere with AFB1 production. To  
420 characterize the interaction between cassava and aflatoxinogenesis, fresh cassava chips  
421 were inoculated with spores of a highly toxigenic *A. flavus* strain. The fungus was able to  
422 rapidly grow on cassava, and was macroscopically visible after 3 days of incubation. This  
423 growth was confirmed by both ergosterol measurement and spore numeration.  
424 Nevertheless, the fungal development on fresh cassava was not associated with AFB1  
425 production. So it demonstrates that fresh cassava is able to block AFB1 production when  
426 colonized by a toxinogenic strain of *A. flavus*. This inhibition of AFB1 production was  
427 observed using 2 different varieties of cassava and with several AFB1 producing species  
428 of the *Flavi* section. To the best of our knowledge, this is the first report indicating that a  
429 raw material, which allows fungal development, may interfere with aflatoxin B1  
430 synthesis. Cassava can therefore be considered as resistant to the contamination with  
431 this mycotoxin. Plant extract/products have been described that inhibit AFB1  
432 production. For instance, Yoshinari et al. demonstrated that German Chamomille blocks  
433 aflatoxin G production (Yoshinari et al., 2008). Razzaghi-Abyaneh et al. (2009) also  
434 showed that *Carum carvi* L. inhibits AF production without any obvious effect on fungal  
435 growth. More recently, Gorran (2013) showed that an aqueous extract of *Thymus*  
436 *daenensis* was able, at a concentration of 2000 mg/l, to reduce the production of AFB1  
437 by 97%.



438 For cassava, our data suggest that cassava is a substrate non-permissive for secondary  
439 metabolism. Indeed, HPLC analysis of extracts revealed that no peak corresponding to  
440 secondary metabolites was observed on a chromatogram when the strain AFc5 was  
441 grown on cassava in contrast to what was observed when the *A. flavus* strain was grown  
442 on the usual media (PDA, MEA or YES) (data not shown). The absence of *A. flavus*  
443 secondary metabolites despite normal growth suggests that cassava acts on the global  
444 regulation of secondary metabolism. In the last decade, several transcription factors  
445 have been found to regulate secondary metabolism in *A. flavus*. The alteration of  
446 transcription factors is often linked to fungal development anomalies. An inhibition of  
447 the recently described transcription factors *nsdC* and *nsdD* is very unlikely. Indeed, the  
448  $\Delta nsdC$  and  $\Delta nsdD$  mutants exhibit columnar arrangement of the conidial chains on the  
449 conidiophores and shorter conidiophore stipes (Cary et al., 2012). In the present study,  
450 no macroscopic or microscopic morphological alteration was observed when *A. flavus*  
451 grew on fresh cassava. A heterotrimeric nuclear complex composed of three proteins  
452 *LaeA*, *VeA* and *VelB* regulates the secondary metabolism (Bayram et al., 2008). *LaeA* has  
453 a major effect on aflatoxin B1 synthesis in *A. flavus* as well as on the whole secondary  
454 metabolism in other *Aspergillus* species (Kale et al., 2008). The disruption of *VeA* leads to  
455 decreased activation of the aflatoxin biosynthesis pathway in *A. parasiticus* and *A. flavus*  
456 (Calvo et al., 2004; Amaike and Keller, 2009). In *A. flavus*, *VeA* is also necessary for the  
457 synthesis of cyclopiazonic acid and aflatrem (Duran et al., 2007). *VelB* regulates the  
458 same biosynthetic pathway (aflatoxin/sterigmatocystin) in *A. nidulans* (Bayram et al.,  
459 2008). In contrast to *nsdC* or *nsdD* genes, the deletion of *LaeA*, *VeA* or *VelB* only  
460 marginally modifies the morphological aspect of the fungi. For instance, disruption of  
461 *VelB* leads to a considerable decrease in conidia production whereas deletion of *VeA* or  
462 *LaeA* only induces a slight decrease in spore numbers (Park et al., 2012). This reduction  
463 of conidiogenesis is accompanied by a decrease in the growth diameter when the  $\Delta laeA$   
464 or  $\Delta veA$  mutants grow on a media plate. The deletion of both *LaeA* and *VeA* genes is  
465 characterized by an inability to produce sclerotia. In the present study, when the *A.*  
466 *flavus* strain AFc5 was grown on fresh cassava, we observed a small decrease in  
467 sporulation and the absence of sclerote formation as well as an inhibition of whole  
468 secondary metabolism. Despite the absence of an effect on fungal growth, these  
469 observations suggest that fresh cassava may act on the *velB/VeA/LaeA* complex.

470 However, a further molecular study should be carried out in order to confirm this  
471 hypothesis.

472 In order to better understand the mechanisms underlying the inhibition of AFB1  
473 synthesis by cassava, the impact of several processes was studied. We observed that the  
474 ability of cassava to block AFB1 production by a toxigenic strain of *A. flavus* was  
475 inhibited by heat treatment, sun drying or freezing of cassava samples. When each of  
476 these processes was applied, the growth of a toxinogenic strain of *A. flavus* on treated  
477 cassava was associated with the production of AFB1. These assays demonstrated that  
478 the molecule responsible for the inhibition of toxin production is quite sensitive and  
479 could correspond to a peptide or small protein.

480 Among candidate compounds that may participate to AFB1 inhibition by cassava, the  
481 role of scopoletin was investigated. This molecule is a thermostable coumarin  
482 phytoalexin with medical properties. It is accumulated in some plants such as carrot  
483 (Coxon et al., 1973), cotton (Zeringue, 1984), sunflower (Gutierrez et al, 1995), noni or  
484 *Morinda Citrifolia* (Deng et al., 2010) and cassava (Buschmann et al., 2000) in response  
485 to both biotic and abiotic stress (Edwards et al., 1997). It can also accumulate in roots  
486 and tubers as a result of post-harvest physiological deterioration. This compound has  
487 been reported to be a potent anti-microbial molecule with fungicidal properties  
488 (Rodriguez et al., 2000; Gomez-Vasquez et al., 2004). In the present study, scopoletin  
489 was not responsible for AFB1 inhibition. Indeed, thermal treatment (120°C, 20 min) did  
490 not modify the scopoletin content of cassava chips whereas it restored AFB1 production  
491 by toxinogenic *A. flavus* strain.

492 Similarly, cyanogenic compounds present in fresh cassava may not be involved in the  
493 anti-aflatoxigenic ability of the product since the varieties used in our study were sweet  
494 ones characterized by a low content of cyanogenic compounds. Moreover, many fungi  
495 display natural linamarase activity and are therefore able to break down cyanogenic  
496 glucosides present in cassava (Birk et al., 1996; Amoa-Awua et al., 1997).

497

498 In conclusion, this study demonstrated for the first time that fresh cassava can be  
499 considered as a crop that is resistant to aflatoxin B1 contamination. This report is very  
500 important as a food safety issue due to the importance of cassava in the diet of millions  
501 of people. Indeed, it appears that cassava is naturally protected from AFB1  
502 contamination during the pre-harvest period. This is usually a critical step regarding

503 mycotoxin synthesis due to the high water activity of plants. After that, since this ability  
504 will be lost by cassava during drying, it is necessary to ensure good storage procedure  
505 and avoid remoistening of chips. It is now important to identify the active compound(s)  
506 and the precise mechanism of action in order to be able to inhibit toxin production  
507 during processing and storage and to possibly use it in other applications.

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509

510

### 511 **Acknowledgment**

512

513 Adjovi Y. C. S was supported by the Service of Cooperation and Cultural Action (SCAC) of  
514 the Embassy of France in Benin, the European project MYCORED (FP7-KBBE-2007-2A)  
515 and the International Union of Biochemistry and Molecular Biology (IUBMB). This study  
516 was supported by the projects CAPES/COFECUB (Inhibitox), ANR Alid (Aflafree) and IRD  
517 (Aflared). The authors are grateful to Dr. D Dufour from CIRAD/CIAT, Cali, Colombia, for  
518 the quantification of cyanogenic compounds in cassava. The authors thank S. Peterson,  
519 former curator of the ARS culture collection for providing cultures of reference strains.  
520 They also thank I. Alassane-Kpembi for statistical analysis, Dr L Lagnika and A Lokman  
521 for technical assistance and all the cassava chips producers. The authors are grateful to  
522 Dr J Woodley for his correction of the original manuscript.

523

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796 **Figure legends**

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798 **Figure 1:** Benin map with sampling sites. ■ producer sites. ★ market sites.

799 For the strains isolated at the cassava producers, the survey was performed during the  
800 dry season in 2010. One sample was collected from 6 different producers in 6 villages  
801 (total number of 36 samples).

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803 **Figure 2:** Visual aspect of cassava after inoculation with a toxigenic *A. flavus*.

804 Sterile cassava from Benin and from Costa Rica were contaminated with  $10^4$  spores of *A.*  
805 *flavus* strain AFc 5, and incubated at 25°C for 3 or 6 days.

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807 **Figure 3:** *A. flavus* development and toxinogenesis on non-autoclaved (NA) and  
808 autoclaved (A) cassava from Benin and Costa Rica after 6 days of incubation.

809 Autoclaved and non-autoclaved cassava were contaminated with  $10^4$  spores of *A. flavus*  
810 strain AFc5 and incubated at 25°C for 6 days.

811 a) Spore formation, b) Quantification of ergosterol, c) AFB1 production.

812 Two-way ANOVA indicated a significant effect of heat treatment on spore numbers and  
813 on ergosterol concentration on cassava from Costa Rica after 6 days of culture ( $p < 0.05$ )

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830 **Table 1:** Primers used for molecular characterization of fungal strains isolated from  
 831 cassava samples

Genes	Primers	Nucleotides 5'→3'	References
<i>ITS</i>	ITS5	GGAAGTAAAAGTCGTAACAAGG	White et al, 1990
	ITS4	TCCTCCGCTTATTGATATGC	
<i>β-tubulin</i>	Bt2a	GGTAACCAAATCGGTGCTGCTTTC	Glass et al, 1995
	Bt2b	ACCCTCAGTGTAGTGACCCTTGGC	
<i>Calmodulin</i>	cmd5	CCGAGTACAAGGAGGCCTTC	Hong et al, 2006
	cmd6	CCGATAGAGGTCATAACGTGG	
<i>Acetamidase AmdS12</i>	amdS1	CCATCGGTATAGGAACTGA	Geiser et al, 1999
	amdS2	AGGGTGCCACGGTATGTC	
<i>O-methyltransferase Omt12</i>	omt1	GGAGTATCAGAGGATTTA	Geiser et al, 1999
	omt2	AGTGCTGTAATAGTCAA	
<i>NorB-CypA region</i>	AP1729	GTGCCCAGCATCTTGGTCCACC	Ehrlich et al, 2004
	AP3551	AAGGACTTGATGATTCCTC	
<i>AflD</i>	AflDF	CGGTGTATTTGGTCACCGGGGC	This study
	AflDR	CGGCTGCCTGGGCATCAGTTTC	
<i>AflQ</i>	AflQF	CGTTATGGGAGGATCGGACACG	This study
	AflQR	CCCAGATCTGATCCTCCTGCG	
<i>AflP</i>	AflPF	GGGCATTCATGCCTTGGTTG	This study
	AflPR	CCCATACCTAGATCAAAGCGG	
<i>AflO</i>	AflOF	CTCTGGCGAAGGTCGGCATTG	This study
	AflOR	CTCTCGGCCAGGAAGTCAGG	
<i>AflR</i>	AflRF	GTCGATTTCTTGGCCGAGTC	This study
	AflRR	CTCAGCAAGTAGCCATCCTG	
<i>AflS</i>	AflSF	CAATTGATGCCGGGTGGAG	This study
	AflSR	CAAGTGATGCGTGCGCGTAG	

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840 **Table 2.** Accession number of the sequences deposited at Genbank

<i>Aspergillus</i> species	Gene sequences				
	ITS	Acetamidase	O-methyltransferase	Beta-tubulin	Calmodulin
Isolate number		<i>Amds12</i>	<i>Omt12</i>		
<i>Aspergillus flavus</i>					
AFc4	JX456207	---	---	---	---
AFc5	KC153995	---	---	---	---
AFc6	KC153996	---	---	---	---
AFc21	KC990469	---	---	---	---
AFc22	JX456209	---	---	---	---
AFc25	KC994648	---	---	---	---
AFc33	KC990470	---	---	---	---
AFc34	KC990471	---	---	---	---
AFc37	JX456208	---	---	---	---
AFc40	KC994649	---	---	---	---
<i>Aspergillus novoparasiticus</i>					
AFc31	KC964099	KC921994	KC964097	---	---
AFc32	KC964100	KC921995	KC964098	---	---
<i>Aspergillus parvisclerotigenus</i>					
AFc35	KC964101	---	---	KC954603	KC954605
AFc36	KC964102	---	---	KC954604	KC954606

841 --- : Not tested

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844 **Table 3:** Aflatoxin contamination of cassava samples and toxigenic profile of *Aspergillus*  
 845 section *Flavi* isolated from cassava chips

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Origin of chips	Chip sample (city/N°sample)	AFB1 concentration in chips	Characterization of <i>Aspergillus</i> section <i>Flavi</i> isolates			
			Isolates number	Presence of AF biosynthetic pathway genes	<i>NorB-CypA</i> gap (bp)	Aflatoxins production
PRODUCER	<b>Gobé 1</b>	ND	AFc 21	yes	900	B
	<b>Gobé 2</b>	ND	AFc22	yes	1500	B
	<b>Gobé 6</b>	ND	AFc 23	yes	900	B
	<b>Savè 2</b>	ND	AFc 24	yes	1500	B
	<b>Tchaourou 1</b>	ND	AFc 25	yes	900	B
	<b>Tchaourou 1</b>	ND	AFc 26	yes	900	B
	<b>Tchaourou 2</b>	ND	AFc 27	yes	1500	B
	<b>Tchaourou 3</b>	ND	AFc 28	yes	900	B
	<b>Tchaourou 5</b>	ND	AFc 29	yes	900	B
	<b>Tchaourou 6</b>	ND	AFc 30	yes	900	B
	<b>Goro 1</b>	ND	AFc31	yes	-	B+G
	<b>Goro 2</b>	ND	AFc32	yes	-	B+G
	<b>Ina 1</b>	ND	AFc 33	yes	900	B
	<b>Ina4</b>	ND	AFc 34	yes	1500	B
	<b>Sinissou 1</b>	ND	AFc 35	yes	-	B+G
	<b>Sinissou 2</b>	ND	AFc36	yes	-	B+G
	<b>Sinissou 8</b>	ND	AFc 37	yes	1500	B
	<b>Ina 3</b>	ND	AFc 38	yes	900	B
	<b>Ina 2</b>	ND	AFc 39	yes	900	B
	<b>Ina 2</b>	ND	AFc 40	yes	1500	B
MARKET	<b>Parakou/Depot 1</b>	ND	AFc 1	yes	900	B
	<b>Save 5 S1</b>	ND	AFc 2	yes	900	B
	<b>Save 5 S2</b>	ND	AFc 3	yes	900	-
	<b>Cocotomey</b>	ND	AFc 4	yes	1500	B
	<b>Gobe 2</b>	ND	AFc 5	yes	900	B
	<b>Bohicon 3</b>	ND	AFc 6	yes	1500	-
	<b>Gobe 3</b>	ND	AFc 7	yes	900	B
	<b>Parakou/Arzeke 2</b>	ND	AFc 10	yes	900	B
	<b>Cocotomey 4</b>	ND	AFc 11	yes	900	B
	<b>Save 3</b>	ND	AFc 12	yes	900	B
	<b>Bougou 5</b>	ND	AFc 13	yes	1500	B
	<b>Kandi</b>	ND	AFc 14	yes	1500	B

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848 ND: not detected (&lt; 0.1 µg/kg)

849 yes: presence of the six biosynthesis genes tested (*AflD*, *AflO*, *AflP*, *AflQ*, *AflR*, *AflS*);

850 All isolates were identified as *Aspergillus flavus* except AFc31, Afc32 (*A. novoparasiticus*)  
 851 and AFc35, Afc36 (*A. parvisclerotigenus*).

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855 **Table 4:** Effect of fresh cassava on *A. flavus* growth and toxinogenesis

Culture duration (days)	Origin of cassava			
	Benin		Costa Rica	
	3	6	3	6
Ergosterol ( $\mu\text{g/g}$ )	13.82 $\pm$ 0.63	26.47 $\pm$ 4.37	18.09 $\pm$ 1.49	50.67 $\pm$ 2.77
Spores ( $\times 10^7$ )	2.33 $\pm$ 0.38	133.00 $\pm$ 11.50	7.43 $\pm$ 2.24	310.00 $\pm$ 26.50
AFB1 (ng/g)	ND	ND	ND	ND

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857 ND: Not detected

858 Results are expressed as mean  $\pm$  SD of three independent experiments

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879 **Table 5:** Effect of fresh cassava on aflatoxin B1 production by different *Aspergillus*  
 880 species of the section *Flavi*

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	Fungal species		
	<i>A. parasiticus</i>	<i>A. bombycis</i>	<i>A. pseudotamarii</i>
	NRRL 4123	NRRL 26010	NRRL 25517
AFB1 production on MEA (ng/g)	477.14 ± 56.88	9.43 ± 3.10	190.53 ± 77.26
AFB1 production on fresh cassava (ng/g)	ND	ND	ND
Ergosterol on cassava (µg/g)	1.18 ± 0.05	1.301 ± 0.05	1.456 ± 0.15

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883 AFB1 and ergosterol measurement were done after 6 days of incubation at 25°C;

884 ND: Not detected.

885 Results are expressed as mean ± SD of three independent experiments

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904 **Table 6:** Impact of drying on *A. flavus* development and toxinogenesis on cassava chips

	Drying time (days)			
	2	3	4	7
Ergosterol ( $\mu\text{g/g}$ )	$36.63 \pm 4.23$	$40.81 \pm 7.03$	$56.51 \pm 2.97$	$43.62 \pm 2.0$
Spores ( $\times 10^7/\text{g}$ )	$6.18 \pm 0.44$	$9.03 \pm 2.68$	$14.9 \pm 0.95$	$35.1 \pm 9.95$
AFB1 ( $\text{ng/g}$ )	ND	ND	ND	$5.5 \pm 0.5$

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906 ND: Not detected

907 Results are expressed as mean  $\pm$  SD of three independent experiments.

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930 **Table 7:** Scopoletin and aflatoxin concentrations in non-autoclaved and autoclaved  
 931 cassava samples from Benin and Costa Rica after 3 days of incubation

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	Origin of cassava			
	Benin Cassava		Costa Rica Cassava	
	Non autoclaved	Autoclaved	Non autoclaved	Autoclaved
Scopoletin ( $\mu\text{g/g}$ )	$13.72 \pm 0.30$	$12.62 \pm 0.30$	$13.45 \pm 4.40$	$10.53 \pm 3.80$
AFB1 ( $\mu\text{g/g}$ )	ND	$1.20 \pm 0.4$	ND	$1.69 \pm 0.5$

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934 ND: Not detected



935 Results are expressed as mean  $\pm$  SD of three independent experiments

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Figure 1



Figure 1 *Adjovi et al.*

Benin	Costa Rica	
		<b>3 days of incubation</b>
		<b>6 days of incubation</b>

**Figure 2** *Adjovi et al.*

Figure 3

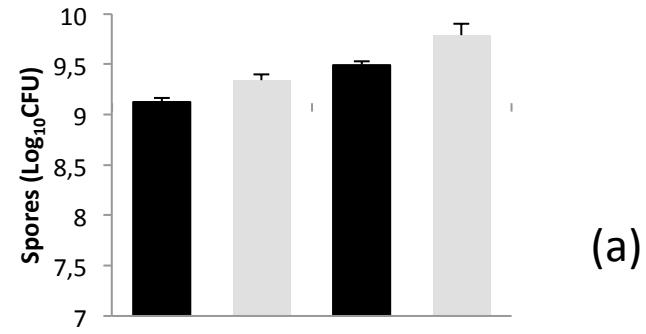
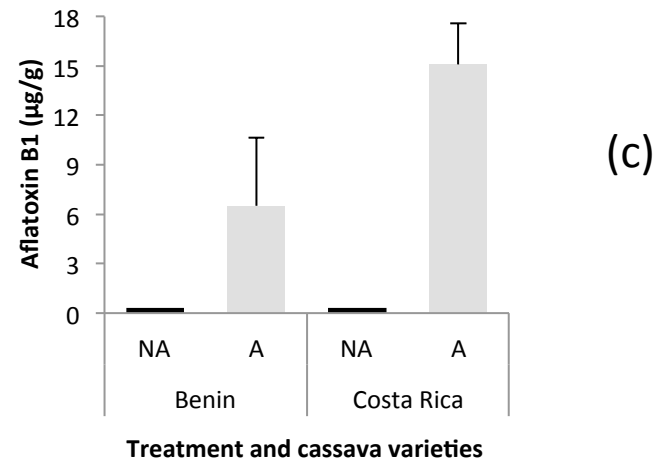
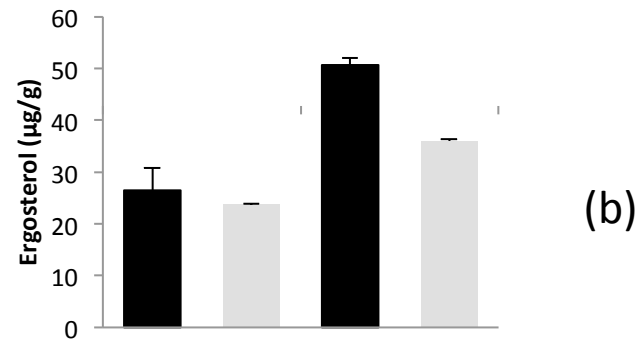


Figure 3 *Adjovi et al.*



### Research Highlights

- Cassava is contaminated with toxigenic *Aspergillus flavus* but not by aflatoxin B1
- Cassava can be contaminated with *A. novoparasiticus*.
- Fresh cassava is able to block AFB1 synthesis by *Aspergillus flavus*
- But cassava does not modify *Aspergillus flavus* development
- Thermal treatment of cassava leads to a loss of the anti-aflatoxin ability