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**Original Research Article**  
***In vivo antifibrotic potential of extracts of  
Acanthospermum hispidum DC. evaluated in  
wistar rats using diethylnitrosamine***

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**ABSTRACT**

**Aims:** Liver fibrosis is a chronic disease of the liver. This disease is a stage of passage to liver cancer. The objective of this work was to evaluate the ability of the ethanolic extract of *Acanthospermum hispidum* to block the progression of hepatic fibrosis induced in rats using diethylnitrosamine (DEN).

**Study design:** Study of the antifibrotic potential of extracts of *Acanthospermum hispidum*.

**Place and Duration of Study:** *In vivo* tests were performed from September 2018 to January 2019. The animal model tests were carried out in the pet shop of the Institute for Health Sciences Research (IRSS) of Burkina Faso and in the Cytogenetics Laboratory (FSS/ISBA) of the Republic of Benin.

**Methodology:** The evaluation of the antifibrotic activity consisted in treating in *wistar* rats a liver fibrosis induced with the DEN which is a chemical agent whose effect on the liver has already been confirmed. As a result of the treatment, all animals were removed from the liver and blood. The livers were used for macroscopic and microscopic observations. Blood has been used for the evaluation of biochemical parameters in relation to fibrosis.

**Results:** The analysis of the results of the biochemical parameters in relation to the fibrosis showed that the ethanolic extract of *Acanthospermum hispidum* at the dose of 250 mg / kg made it possible to obtain an improvement of these parameters compared to the other batches of animals. These results have been confirmed by those of the anatomopathological studies.

**Conclusion:** The results of biochemical and histological analyzes revealed a capacity of *Acanthospermum hispidum* extracts to block the evolution of hepatic fibrosis in the rat. These results confirm the hepatoprotective potential of this medicinal plant used in traditional medicine in Burkina Faso.

**Keywords:** *Acanthospermum hispidum*, Diethylnitrosamine, liver, *invivo* antifibrotic

**1. INTRODUCTION**

Hepatic fibrosis is due to the excessive accumulation of matrix components in the liver. In addition to the quantitative increase in collagen and other matrix proteins, it is characterized by qualitative changes in the nature of the matrix components deposited and their distribution in the liver [1]. Hepatic fibrosis complicates all chronic liver diseases, whether due to chronic alcoholism, viral B or C infection, or autoimmune, biliary, parasitic or medicinal. It is now accepted that hepatic fibrosis is a dynamic process, causing not only excessive production of matrix components (fibrogenesis) [2], but also a decrease in their degradation (fibrolysis) [3].

The WHO estimates that 2 billion people are infected with the hepatitis B virus and 400 million have chronic carriers, including 60 million in Africa [4]. Burkina Faso has an estimated

27 prevalence of 14.4% of the hepatitis B virus [5]. Viral hepatitis, in particular those caused by  
28 hepatitis B and C viruses, cause respectively 1300 and 900 deaths from liver cancer each  
29 year, which they can cause [6]. The latter constitutes in Burkina Faso the first cause of  
30 health evacuations out of the country and the third cause of mortality after infectious  
31 diseases and cardiovascular diseases [7][8]. Treatment of chronic hepatitis in Burkina Faso  
32 could cost \$ 909 per month per patient [5]. The fibrosing diseases represent 45% of the  
33 causes of mortality in the world. The lack of therapeutic alternatives for the management of  
34 patients with chronic hepatitis makes liver fibrosis a very poor prognosis [9].

35 *Acanthospermum hispidum* is an herb that has had a good hepatoprotective capacity. The  
36 objective of this study was to evaluate the ability of the ethanolic extract of *Acanthospermum*  
37 *hispidum* to block the progression of hepatic fibrosis induced in experimental animals. For  
38 this purpose, diethylnitrosamine (DEN), a toxic substance known to induce hepatic fibrosis in  
39 laboratory animals, has been used as a hepatotoxin.

## 40 **2. MATERIAL AND METHODS**

### 41 **1.1 Material**

#### 42 **2.1.1 Plant material**

43 The plant material consists of the whole plant of *Acanthospermum hispidum* harvested in  
44 2018 in Loubila (12 ° 19'35.84 N, 1 ° 35'13.5 W). The plant has been identified at the  
45 Laboratory of Plant Ecology and Botany of University Ouaga 1 Pr Joseph KI-ZERBO.

#### 46 **2.1.2 Consumables**

47 Aluminum foil, Kit surgery, 1cc and 5cc Syringes, porcelain mortar, blades and microscope  
48 slides, gloves, bleach, blotting paper, micropipettes, Eppendorff tubes, Alcohol 90. They are  
49 all analytical grade.

#### 50 **2.1.3 Reagents**

51 Diphenylboryloxyethylamine, sodium phosphate monobasic ( $\text{NaH}_2\text{PO}_4$ ), dibasic sodium  
52 phosphate ( $\text{Na}_2\text{HPO}_4$ ), EDTA (Ethylenediaminetetraacetic acid), Diethylnitrosamine (DEN),  
53 Silymarin.

#### 54 **2.1.4 Physiological Solutions**

55 Phosphate buffer, tris buffer, dimethylsulfoxide (DMSO), sodium hydroxide, sodium chloride  
56 (9 ‰), potassium chloride (9 ‰), formalin buffer (10%).

### 57 **1.2 Methods**

#### 58 **2.2.1 Extraction by ethanol maceration**

59 Fifty grams (50 g) of the whole plant powder were extracted with stirring for 24 hours with  
60 1000 mL of ethanol. After filtration under reduced pressure, the filtrate was frozen and  
61 freeze-dried.

#### 62 **2.2.2 Animal treatment**

63 Pre-test allowed to identify doses of DEN to be administered to rats, as well as the duration  
64 of treatment required to obtain the liver fibrosis.

65 The antifibrotic activity in curative mode was evaluated according to the following protocol  
66 [10] with some modifications:

67 Male *Wistar* rats were randomly assigned to batches of eight (8) rats after a two-week  
68 acclimation period. The rats used were free of pathogenic organisms and healthy status. The  
69 experiments met the requirements of the Code of Ethics: The Institutional Animal Ethics  
70 Committee (Directive 2010/63 / EU on the protection of animals used for scientific purposes).  
71 Ethical approval code: 2010/63 / EU, Date of approval: 20 October 2010.

72 Group I (Normal Group): The rats received standard treatment during the eight weeks.

73 Group II (Negative control group): The rats in the group received water in place of the extract  
74 after administering the DEN intraperitoneally (75 mg/kg body weight) once a week during the  
75 4 first weeks.

76 Group III (Positive Control Group): Rats received a daily dose of 100 mg/kg silymarin for 4  
77 weeks after intraperitoneal injection of DEN (75 mg/kg body weight) once a week during the  
78 first four weeks.

80 Test Groups IV and V: The rats received intraperitoneally DEN (75 mg/kg of body weight per  
81 week) during the first four weeks and during the last four weeks these animals received daily  
82 doses (100 and 250 mg/kg body weight) of ethanolic extract of *Acanthospermum hispidum*.

### 83 **2.2.3 Registration of body weight of animals**

84 The body weight of the treated animals was recorded using a scale at 1<sup>st</sup> day, week 2, week  
85 4, week 6 and week 8, and compared to animals from normal group (group II).

### 86 **2.2.4 Biological analyzes**

#### 87 *2.2.4.1 Collection of blood and liver*

88 Animal blood was collected by cardiac puncture using a 5 mL syringe. To collect, it was first  
89 necessary to stabilize the heart using a pair of pliers. The sample was taken from the left  
90 ventricle. The collected blood had to reach at least a volume of 3 mL so that after  
91 centrifugation we can collect a sufficient volume of serum for the various analyzes. The  
92 collected blood was centrifuged at 3000 g for 10 minutes. After centrifugation, the clear  
93 (supernatant) serum was recovered using 1 mL syringes and placed in the cryotubes for  
94 biochemical markers analysis.

95 The livers of the animals were removed by getting rid of the stomach, diaphragm and  
96 adhesions. The livers were kept in formalin (10%) for the pathological study.

#### 97 *2.2.4.2 Biochemical analyzes*

98 Blood samples were taken for biochemistry. These blood samples in the tubes without  
99 anticoagulant were centrifuged for 10 minutes to obtain serum. Serum has been used for the  
100 evaluation of biochemical parameters that are indirect markers of liver fibrosis such as  
101 aspartate aminotransferase (ASAT), alanine aminotransferase (ALAT), total bilirubin  
102 (bilirubin T), albumin and alkaline phosphatase (PAL). All these parameters were determined  
103 using kits (Selectra XL Vital Scientific Elitech Group Company) according to the instructions  
104 of the manufacturer.

### 105 **2.2.5 Histopathological analyzes**

106 The livers of the treated animals were removed, weighed and used for histological analysis.  
107 The methodology used was that of Hould[11]. Liver sections (about 0.2 × 0.2 cm) were made  
108 with the rotating microtome (Leitz 1512). These sections were fixed in 10% formalin and then  
109 placed in a paraffin bath. The liver slices were then labeled with hematoxylin-eosin. Finally,  
110 these labeled liver slices were subjected to microscopic examination for histological analysis.

### 111 **2.2.6 Statistical analysis**

112 The data were expressed as mean ± standard deviation. Graphics were drawn and statistical  
113 analysis was performed using GraphPad Prism software version 5.0 for Mac OS X  
114 (GraphPad software, San Diego, California, USA).

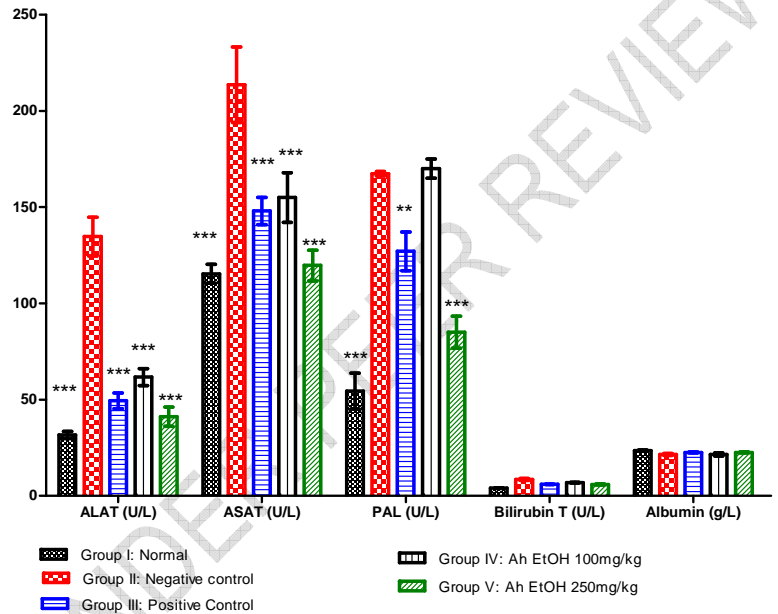
## 116 **3. RESULTS AND DISCUSSION**

### 117 **3.1. Evolution of biochemical parameters**

118 For transaminase values, a very significant difference ( $p < 0.001$ ) exists between ALAT and  
119 ASAT values in DEN-only and 250 mg-treated animals. kg of body weight. This finding was  
120 also made for the ALAT and ASAT values of animals treated with silymarin and those  
121 treated with the extract at a dose of 100 mg / kg body weight compared to the negative  
122 control (DEN). In addition, the ASAT value in animals treated with silymarin and in animals  
123 treated with the ethanolic extract of *Acanthospermum hispidum* at a dose of 250 mg / kg  
124 body weight, showed a significant difference ( $0.05 > p > 0.01$ ) with respectively  $148.00 \pm$   
125  $12.17$  U/L and  $119.67 \pm 13.80$  U/L.

126 Compared to alkaline phosphatase values, a very significant difference ( $p < 0.001$ ) was  
 127 observed between the mean value of DEN alone and those treated with ethanolic extract at  
 128 250 mg/kg body weight as well as for controls. Animals treated with the ethanolic extract at  
 129 the dose of 250 mg/kg also showed a mean value of alkaline phosphatase which is not  
 130 statistically different from that of the controls; which on the other hand is statistically very  
 131 different ( $p < 0.001$ ) from the value of animals treated with silymarin (100 mg/kg) with  
 132 respectively  $127.00 \pm 17.52$  U/L and  $85.00 \pm 14.42$  U/L.

133 The values of albumin and bilirubin did not differ significantly between those treated with  
 134 DEN alone and animals from other lots. In contrast, a low mean value of albumin was  
 135 recorded in the animals that received only DEN ( $21.43 \pm 0.76$  g/L). The highest mean value  
 136 of bilirubin was also observed in animals treated with DEN alone ( $8.47 \pm 0.76$  U/L) (Figure  
 137 1).



138 **Fig. 1. Results of Biochemical Parameters of Treated Animals**

139 ALAT: Alanine Amino-transferase; ASAT: Aspartate Amino-Transferase; MDA: MalonDiAldehyde;  
 140 PAL: Alkaline phosphatase: significant from positive control, \*  $P < 0.05$ ; \*\*  $P < 0.005$ ; \*\*\*  $P <$   
 141  $0.001$

142  $Mean \pm S.E.M = Mean values \pm Standard error of means of eight experiments.$

143 **3.2 Variation in animal weight and relative weight of livers**

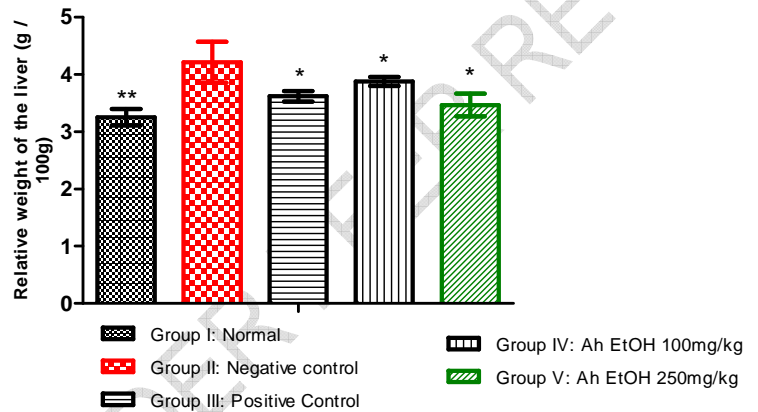
144  
 145  
 146 Animal weight analysis showed a highly significant difference ( $p > 0.001$ ) between animals  
 147 treated with ethanolic extract at 100 mg / kg and 250 mg / kg compared to those in group 2  
 148 (DEN alone) at the fourth week. By the eighth week, it appears that the difference between  
 149 the average weight of the animals treated with the extract at the dose of 250 mg/kg and that  
 150 of the animals treated with the DEN alone is very highly significant ( $p < 0.001$ ). In addition,

151 there was a statistical difference ( $p > 0.01$ ) between the mean weight of animals treated with  
 152 silymarin and those treated with the extract at a dose of 250 mg/kg (Table 1).  
 153

| Groups    | 1st day      | 2nd week       | 4th week       | 6th week        | 8th week      |
|-----------|--------------|----------------|----------------|-----------------|---------------|
| Group I   | 165.33±3.78* | 189.33±13.69** | 209±12.43***   | 229.33±16.93*** | 235.67±14.87  |
| Group II  | 170.17±4.12  | 165.33±3.62    | 162.67±10.48   | 160.67±11.18    | 168.00±15.72  |
| Group III | 180.85±9.5*  | 178.82±7.25**  | 170.48±6.48**  | 168.42±6.98*    | 173.64±7.01** |
| Group IV  | 174.83±3.60  | 1693.00±6.26*  | 167.00±13.24** | 161.33±9.18     | 165.67±5.96*  |
| Group V   | 178.67±3.78  | 172.67±9.18*   | 170.30±11.54** | 169.00±5.59*    | 175.67±8.12*  |

154 *Group I: control, Group II: DEN, Group III: DEN + Silymarin, Group IV: DEN + Ah 100mg, Group V:*  
 155 *DEN + Ah 250mg, significant from positive control, \*  $P < 0.05$ ; \*\*  $P < 0.005$ ; \*\*\*  $P < 0.001$  Mean ± S.E.M*  
 156 *= Mean values ± Standard error of means of eight experiments.)*

157 The relative weight values of the livers of the treated animals did not show a statistical  
 158 difference between the batches. On the other hand, the average values of the relative weight  
 159 of the organs show that the relative weight of the livers of the negative control lot is relatively  
 160 high compared to the other lots (Figure 2).



161

162 **Fig. 2. Effect of the extracts on the variation of the relative weight of the livers of the**  
 163 **treated animals**

164 *significant from positive control, \*  $P < 0.05$ ; \*\*  $P < 0.005$ ;*  
 165 *Mean ± S.E.M = Mean values ± Standard error of means of eight experiments.*  
 166

### 167 3.3 Histopathological studies

168 Macroscopic observation of the liver in treated animals showed that, compared to the liver of  
 169 the animals in the control group (photos 1.a), that of the negative control group (photos 1.b)  
 170 had a brownish or whitish coloration on its surface (nodules). In addition, compared to the  
 171 negative control group, the animals treated with the extract at the dose of 250 mg/kg (photo  
 172 1.e) have livers whose state is significantly improved. Finally, compared to the liver of the

173 animals in the positive control group (silymarin 100 mg/kg of body weight), the ethanolic  
174 extract at the dose of 250 mg/kg presented a liver with a more regular appearance.

175 Microscopic observation of liver sections in normal control animals shows normal liver cells  
176 with a well preserved cytoplasm and a visible central vein. This shows the absence of  
177 collagen deposition on hepatocytes (photos 2.a). In contrast, rats treated with DEN alone  
178 showed liver cuts with damaged structures and characterized by necrosis around the central  
179 vein, inflammatory cell infiltration, hot air balloon degeneration and sinusoidal dilatation  
180 (photo 2.b). However, the liver sections of the animals that received the 250 mg/kg dose  
181 extract (photo 2.d) showed a moderate degree of damage to the liver and inflammatory cells.  
182 Extracts at this dose protected the liver against hepatocyte degradation and centrilobular  
183 necrosis (photos 2.d). Histopathological examination of hepatic sections of animals treated  
184 with ethanolic extract at a dose of 250 mg/kg also showed normal hepatocytes and lacked  
185 collagen accumulation comparable to the positive control group (photos 2.e).



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187 *a. Liver (Group I)*      *b. Liver (Group II)*

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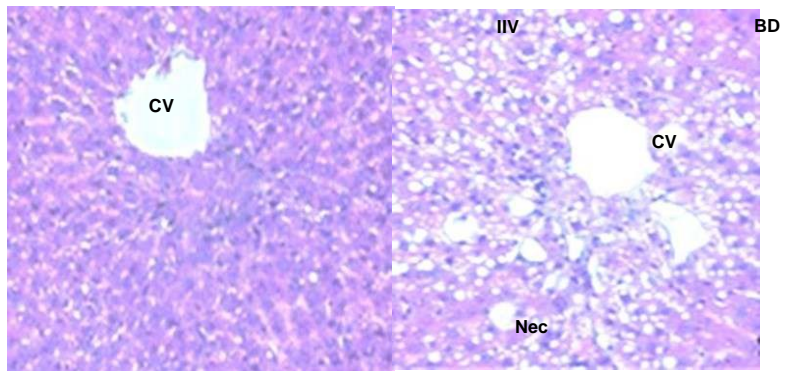


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190 *c. Liver (Group III)*      *d. Liver (Group IV)*      *e. Liver (Group V)*

191 **Photo 1. Macroscopic appearance of the livers of treated animals**

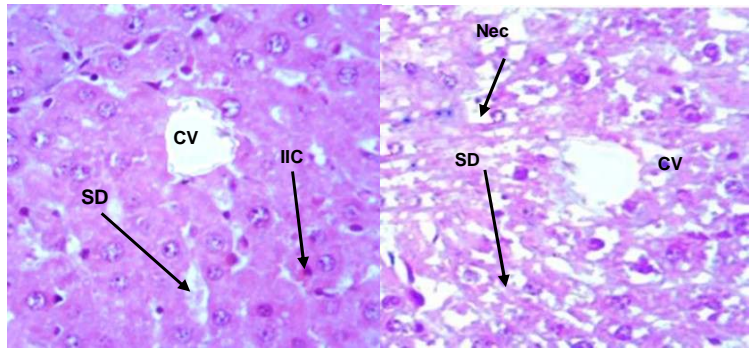
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a. Group I: liver cup: Normal Group

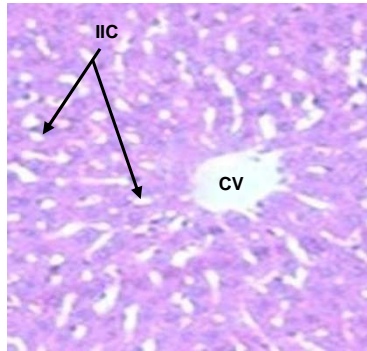
b. Group II: liver cup: negative control



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c. Group III Liver section, DEN + Silymarin.

d. Group IV liver section, DEN + Ah (100mg/kg)



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e. Group V liver section, DEN + Ah (250mg/kg)

201 **Photo 2. Structures of histopathological sections of the livers of treated rats**

202 *CV: central vein; IIC: Infiltration of inflammatory cells; BD: Bloating degenerations, SD: Sinusoidal*  
203 *dilation; Nec: Necroses (The cuts were stained with H and E, x 400)*

204

#### 205 **4. DISCUSSION**

206 Fibrosis usually presents with signs and symptoms of chronic liver disease such as portal  
207 hypertension, fatigue, weight loss, hepatosplenomegaly, ascites, varicose veins and muscle  
208 atrophy [12]. Registration of the weight of animals in the negative control lot confirmed a  
209 significant loss of weight, which was improved in the test animals (100 mg/kg and 250 mg/kg  
210 body weight extract). Elevated ASAT and ALT values in animals in the negative control lot  
211 ( $134.67 \pm 17.47$  U/L and  $213.67 \pm 33.97$  U/L) were also identified in a liver fibrosis study.  
212 human [13]. Moreover, according to Edouardo et al. [14], the ratio greater than 1, obtained in  
213 lot II (negative control), would show advanced liver fibrosis in these animals. The serum  
214 activity of alkaline phosphatase (ALP) comes mainly from the liver [15]. The serum PAL of  
215 the test lots showed hepatic function regulation which would prevent the establishment of  
216 fibrosis in animals receiving the dose of 250 mg/kg body weight of ethanolic extract of  
217 *Acanthospermum hispidum*. On the other hand, in animals in Lot II (negative control), high  
218 PAL values could explain a shift to cirrhosis or liver failure [16]. Compared with bilirubin and  
219 albumin values, the low levels observed in animals in lots III and V show liver synthesis  
220 capacity in these animals after the aggression [17]. The results of the histopathological  
221 analyzes were confirmed those of the histopathological studies.

222 In the present study, evidence of hepatotoxicity under the effect of DEN was confirmed. This  
223 hepatotoxin is likely to cause profound damage to the liver following the intensive production  
224 of free radicals causing an imbalance in the cellular redox status in favor of pro-oxidants.  
225 Indeed, it was found during the pre-test that the antioxidant defense system decreased  
226 significantly in the liver homogenates of animals of the negative control (DEN alone), leaving  
227 room for the pro-oxidants responsible for lipoperoxidation and destruction. membrane  
228 structures. The ethanolic extract of *Acanthospermum hispidum* plays a chemoprotective role  
229 against the oxidative stress produced in the cytosol and mitochondria of hepatocytes,  
230 following the administration of DEN to laboratory animals in the evaluation of the antifibrotic  
231 capacity of the extracts of *Acanthospermum hispidum*. By its ability to neutralize the reactive  
232 species produced through the metabolism of DEN [4], the ethanolic extract has shown that it  
233 has an ability to block the progression of liver fibrosis.

234

#### 235 **5. CONCLUSION**

236 It is clear from this study that the ethanolic extract of *Acanthospermum hispidum* has  
237 antifibrotic properties. It is an interesting extract, rich in therapeutics, by its power to prevent  
238 the progression of liver fibrosis. The ethanolic extract at a dose of 250 mg/kg yielded  
239 interesting results in the relative weights of the animals and livers of the treated animals.  
240 Mean values for transaminase, alkaline phosphatase, total protein and total bilirubin levels  
241 observed in the animals treated with the extract were significantly improved compared to  
242 animals in the negative and positive control groups. The results of the histological studies  
243 performed on the livers of the treated animals also showed aspects of liver tissue with  
244 improved structure for group V. All of these results militate in favor of the use of the ethanolic  
245 extract of *Acanthospermum hispidum* against chronic liver infections such as fibrosis.

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247

#### 248 **COMPETING INTERESTS**

249 Authors have declared that no competing interests exist.

250

#### 251 **CONSENT (WHEREEVER APPLICABLE)**

252 All authors declare that "written informed consent was obtained from the patient (or other  
253 approved parties) for publication of this case report and accompanying images. A copy of



254 the written consent is available for review by the Editorial office/Chief Editor/Editorial Board  
255 members of this journal."

256

257

#### ETHICAL APPROVAL (WHEREEVER APPLICABLE)

258 ALL AUTHORS HEREBY DECLARE THAT "PRINCIPLES OF LABORATORY ANIMAL  
259 CARE" (ETHICAL APPROVAL CODE: 2010/63/EU, DATE OF APPROVAL: 20.10.2010)  
260 WERE FOLLOWED, AS WELL AS SPECIFIC NATIONAL LAWS WHERE APPLICABLE.  
261 ALL EXPERIMENTS HAVE BEEN EXAMINED AND APPROVED BY THE APPROPRIATE  
262 ETHICS COMMITTEE.

263

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#### REFERENCES

- 265 1. Poole LG, Arteel GE. Transitional Remodeling of the Hepatic Extracellular Matrix in  
266 Alcohol-Induced Liver Injury. *Biomed Res Int.* 2016;2016.
- 267 2. Chang H, Meng HY, Liu SM, Wang Y, Yang XX, et al. Identification of key metabolic  
268 changes during liver fibrosis progression in rats using a urine and serum  
269 metabolomics approach. *Sci Rep.* 2017;7(1):1–12.
- 270 3. Younossi Z, Loomba R, Rinella M, Bugianesi E, Marchesini B. Risk scores for HCC in  
271 CHB. *Hepatology.* 2017;77(5):1–36.
- 272 4. OMS. Hépatite: Améliorer la santé des personnes atteintes d'hépatite virale. 2014.  
273 French
- 274 5. Lingani M, Akita T, Ouoba S, Sanou AM, Sugiyama A, et al. High prevalence of  
275 hepatitis B infections in Burkina Faso (1996–2017): a systematic review with meta-  
276 analysis of epidemiological studies. *BMC Public Health.* 2018;18(551):1–11.
- 277 6. Zeba MTA. Co-infection des virus des hépatites B et C au Burkina Faso : Prévalence,  
278 marqueurs viraux et caractérisation moléculaire. Université de Ouagadougou; 2011.
- 279 7. N'do JY-P, Hilou A, Pare D, Sombie EN, Traore TK, et al. Protective Effect of  
280 *Acanthospermum hispidum* DC (Asteraceae) Extracts against Diethylnitrosamine  
281 Induced Hepatocellular Damage. *J Complement Altern Med Res.* 2019;4(6):1–13.
- 282 8. Plan stratégique de lutte contre le cancer 2013-2017MS (Ministère de la Santé). Plan  
283 stratégique de lutte contre le cancer 2013 - 2017. 2013. French
- 284 9. Sebastiani G, Gkouvatsos K, Pantopoulos K. Chronic hepatitis C and liver fibrosis.  
285 *World J Gastroenterol.* 2014;20(32):11033–53.
- 286 10. Hu Z, Wang W. Effect of *Carthamus tinctorius* L Extract on Diethylnitrosamine-  
287 Induced Liver Cirrhosis in Rats. *Trop J Pharm Res.* 2015;14(July):1213–6.
- 288 11. Hould R. Techniques d'histopathologie et de cytopathologie. Maloine. 1984. French
- 289 12. Laouar A, Klibet F, Bourogaa E, Benamara A, Boumendjel A, et al. Potential  
290 antioxidant properties and hepatoprotective effects of *Juniperus phoenicea* berries  
291 against CCl<sub>4</sub> induced hepatic damage in rats. *Asian Pac J Trop Med [Internet].*  
292 2017;10(3):263–9. Available from: <http://dx.doi.org/10.1016/j.apjtm.2017.03.005>
- 293 13. Robert C, Thomas R, Syed MA, Mattew WP. Mildly Elevated Liver Transaminase  
294 Levels: Causes and Evaluation. *Am Fam Physician.* 2017;11(96):709–15.
- 295 14. Giannini EG, Testa R, Savarino V. Liver enzyme alteration: A guide for clinicians.  
296 *Cmaj.* 2005;172(3):367–79.
- 297 15. Behera S, Ray S, Jena I, Sundar Ray C, Singh B. Low Alkaline Phosphatase (ALP)  
298 In Adult Population an Indicator of Zinc (Zn) and Magnesium (Mg) Deficiency. *Curr*  
299 *Res Nutr Food Sci J.* 2017;5(3):347–52.
- 300 16. Thapa BR, Walia A. Liver Function Tests and their Interpretation. *Indian J Paediatr.*  
301 2007;74(7):67–75.
- 302 17. Adak M, Shivapuri JN. Research Journal of Pharmaceutical , Biological and  
303 Chemical Sciences Enzymatic and Non-enzymatic Liver Function Test : A Review. *Res J*  
304 *Pharm Biol Chem Sci.* 2010;1(593):593–605.
- 305