

# Relevance of *Chrysanthellum americanum* (L.) Vatke extracts in rat liver protection

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## ABSTRACT

**Aims:** to assess the relevance of *Chrysanthellum americanum* (L.) Vatke extracts in rat liver protection.

**Place and Duration of Study:** Laboratory of Biochemistry and Applied Chemistry (LABIOCA), also in Laboratory of Department of Medicine and Traditional Pharmacopoeia (MEPHATRA-PH) of Institute for Research in Health Sciences (IRSS/CNRST) of Burkina Faso between July 2014 and August 2015.

**Study design:** Polyphenolic extract of *Chrysanthellum americanum*- in vivo liver protection- in vivo liver intoxication- liver necrosis parameters analysis, histopathology analysis, in vivo and in vitro antioxidant assay.

**Background:** *Chrysanthellum americanum* L. (Vatke) is a medicinal plant well known for its flavonoids and saponins richness, but also for its strong antioxidant potential and use traditionally for liver disease treatment.

**Methodology:** In vivo, anti hepatotoxicity effects of *Chrysanthellum americanum* was evaluated using CCl<sub>4</sub> as hepatotoxic agent. Also, acute toxicities were determined using standards methods, serum parameters of liver injury using Cypress Diagnostics kits and histopathology analysis using Mayer's haematoxylin- eosin-phloxine coloration method. For in vitro tests, malondialdehyde and thiobarbituric acid method were used in lipid peroxidation assessment and the ABTS method in Trolox Equivalent Antioxidant Capacity assessing.

**Results:** Result showed that the crude extract of *C. americanum* has a very low oral toxicity but, in intraperitoneal route this extract presented a high toxicity (LD<sub>50</sub>= 175 mg / kg of body weight). Histopathology micrograph indicated reduction in number of necrotic cells induced by CCl<sub>4</sub>. This beneficial action was confirmed by reduction in serum transaminases and malondialdehyde (22.68 ± 0.68 mmol MDA/ g of liver weight). In vitro antioxidant capacities, this plant extract presented a result of 35.01 ± 0.26 % and 42.01 ± 0.26 mg TE/ g respectively in LPO and TEAC.

**Conclusion:** Given our results, our research confirms that *Chrysanthellum americanum* extracts have in vivo physiological impact and benefits in traditional medicine for specific care of liver diseases.

**Keywords:** Hepatotoxicity; Medicinal plant; antioxidant; toxicity

## 1. INTRODUCTION

Diet and good digestion are very important factors for good health and also for life good mood [1]. For this, gastroenterology diseases purpose a negative impact on the functioning of body vital organs but also on the psychology of the human being.

The liver is one of the main gastroenteric organs that has several functions of which the main ones are detoxification, synthesis (carbohydrates, lipids and proteins) and storage (vitamins A, D, E, K and glycogen) [2,3]. Being that the liver a purifying organ, its diseases

are very numerous by passing from alcoholic diseases to toxic diseases and inflammatory diseases as well[4,5].

Causes of liver pathologies are several (alcohol, toxins, hepatitis virus...). However, oxidative stress is a primary factor in the appearance of these diseases with pronounced psychological effects (anxiety)[6]. Oxidative stress defined as a state of imbalance between oxidants (toxic compounds) and antioxidant defense system (molecular and enzyme) of an organism is involved in several diseases especially in metabolic diseases[7,8].

In Burkina Faso, as in most low-income countries, poverty equated with lack of hygiene keeps many people in a state of fairly high stress. In this context, the populations are subject to food and alcoholic poisoning and also viral hepatitis which have the liver as potential target.

In European countries liver diseases remain a problem[9]. Also, for WHO, hepatitis will have to be eliminated by 2030. Research to fight against liver diseases have seen many encouraging results but there are still dissatisfactions. One thing is also the high cost of treatments available for low income populations, so medicinal plants are their alternative.

Since ancient times, in African, Chinese and Ayurvedic medicines, plants have been a very important source of natural chemical compounds with enormous therapeutic potentials. Looking for remedies to establish health, researchers are turning more and more to these medicinal plants[10,11]. *Chrysanthellum americanum* (L.) Vatke is a plant used in Burkina Faso traditional medicine for its extracts antioxidant power but also well-known in herbal medicine research area.

*Chrysanthellum americanum* is a small erect or less prone herbaceous plant with very few leaves and yellow flowers belonging to *Asteraceae* family[12].

This plant extracts are known to possess antioxidant, P-vitamin and antilithiasis remarkable properties<sup>13</sup>. Most of therapeutical properties of *C. americanum* extracts are attributed to saponins (Chrysantheine A & B) and to flavonoids (luteolin 7-O-glucoside, eriodictyol 7-O-glucoside, isookanin 7-O-glucoside or flavonomarein, okanin 4'-O-glucoside or marein, maritimetin 6-O-glucoside or maritimein) of which they are consisted[14,15]. Polyphenolic compounds are well known for their antioxidant capabilities, their capacity to improve hepatoprotection[16,17,18,19].

*C. americanum* is a medicinal plant that its extracts are endowed with very good antioxidant capacity, but also a good candidate for treatment of pathologies related to oxidative stress[20].

Liver pathologies are disorder or diseases exacerbated by oxidative stress and affect the psychology of the patient. Thus, this present study aims to evaluate impact of polyphenolic extract of *Chrysanthellum americanum* (L.) Vatke on carbon tetrachloride hepatotoxicity on rat model.

## 2. MATERIALS AND METHODS

### 2.1 Plant material and extraction

*Chrysanthellum americanum* (L.) Vatke whole species was collected during August 2014 in Loumlila, 15 Km north of Ouagadougou, the capital of Burkina Faso. The plant was identified by Prof. Millogo- Rasolodimby from plants Biology Department of the University of Ouagadougou. A voucher specimen (ID-10474) was deposited at the Herbarium of the University of Ouagadougou.

*Chrysanthellum americanum* (L.) Vatke whole-plant was dried at room temperature and ground to fine powder. Seventy-five gram of this powder was macerated during 48 hours with mechanical stirring using 750 mL of aqueous ethanol (80% v/v) at laboratory conditions. After, extract solutions were concentrated under reduced pressure in a rotary evaporator (BÜCHI, Rotavapor R-200, Switzerland) at approximately 40°C, frozen and lyophilized using a lyophilizer (Telstar-Cryodos 50, Spain). The aqueous ethanol extract (Crude extract) obtained was fractionated by solvents of increasing polarity (dichloromethane, ethyl acetate, butanol and water residual). Crude extract and butanol fraction (polyphenols extract) were

weighted before packed in waterproof plastic flasks and stored at 4°C until use. The yields of crude aqueous ethanol extract and polyphenols extract were 8.00% and 6.22% respectively (Yield of the crude extract was calculated with respect to vegetable powder mass and the yield of the butanol fraction relative to crude extract mass).

## **2.2 Animals**

Thirty female and male Wistar rats weighting respectively  $238.40 \pm 18.70$  g and  $310 \pm 48.60$  g and mice from Naval Medical and Research Institute (NMRI) ( $31.83 \pm 4.77$  g) at the start of the experiment were used. The animals were housed in a temperature and light-controlled room (22 °C, a 12 h cycle starting at 08:00 h) and were fed with industrial pellets with 29% protein and allowed to drink water ad libitum. Rats and mice were treated in accordance with the guidelines of animal bioethics from the Act on Animal Experimentation and Animal Health and Welfare Act from Romania and all procedures were in compliance with the European Council Directive of 24 November 1986 (86/609/EEC). "Principles of laboratory animal care" (NIH publication No. 85-23, revised 1985) were followed, as well as specific national laws where applicable. All experiments have been examined and approved by the appropriate ethics committee".

All evaluations were performed between 9 h and 16 h.

## **2.3 Chemicals**

To carry out our activities, we used analytical grade solvents and various classic reagents. Ethyl acetate and 2-thiobarbituric acid were purchased from Sigma Aldrich chimie (Steinheim, Germany); potassium persulfate, 2,2'-azinobis (3 ethylbenzothiazoline-6-sulphonate) ABTS and trichloroacetic acid were supplied by Fluka chimie (Buchs, Switzerland); dichloromethane, ferric dichloride, Carbon tetrachloride, ethanol were sourced from Probalo (Paris, France); butanol was sourced from sds (Peyin, France).

## **2.4 In vivo experiments**

### **2.4.1. Toxicities evaluations**

The mice were randomized into groups of 6 mice (3 males and 3 females) including a control group for the crude extract of *C. americanum*. Each animal was identified by a different mark (head, back, right flank, left flank, tail and without mark). The animals were fasted for 12 hours, then the weight of each rat was taken, and they received a given dose of extract per group. The route of administrations of the extracts were oral or intraperitoneal[21]. The number of deaths per group was determined after 2h, 24h, 48h, 72h and the animals were kept under observation for a week.

50% lethal dose (LD<sub>50</sub>) determination and its confidence limits is what was described by Ouedraogo[22]. It consists of directly carrying Log Probit paper the percentage of mortality according to the log of the dose. Before going to the tests, pre-tests were carried out on group of three (03) animals allowing to locate the lethal dose 50%.

### **2.4.2. Anti hepatotoxicity activity of *C. americanum* phenolic extract**

The anti hepatotoxicity activity of *C. americanum* was evaluated according to the protocol described by Sanogo[23].

#### **2.4.2.1 Experiment design**

Rats were randomized into four (4) groups of six (6) animals:

**Group I:** normal control group, animals received distilled water (10 mL / kg of body weight per day) for 7 days *per os* and the 7<sup>th</sup> day received olive oil 2mL / kg of body weight intraperitoneally 1 hour after water administration;

121 **Group II:** negative control group, animals received distilled water (10 mL / kg of body weight  
122 per day) for 7 days *per os*; the 7<sup>th</sup> day received 2mL / kg of CCl<sub>4</sub> (50% dissolved in olive oil)  
123 intraperitoneally 1 hour after the administration of the water;  
124 **Group III:** positive control group, animals were treated with silymarin (50 mg / kg of body  
125 weight) for 7 days *per os* then the 7<sup>th</sup> day received 2 mL / kg of CCl<sub>4</sub> (50% dissolved in olive  
126 oil) intraperitoneally 1 hour after administration of silymarin.  
127 **Group IV:** test group, animals were treated with *C. americanum* phenolic extract (100 mg /  
128 kg of body weight) for 7 days *per os* then the 7<sup>th</sup> day received 2 mL/kg of CCl<sub>4</sub> (50%  
129 dissolved in olive oil) intraperitoneally 1 hour after administration of the extract.

#### 130 **2.4.2.2 Anti hepatotoxicity evaluation**

131 On day 8<sup>th</sup>, animals were sacrificed after being anesthetized with ketamine (150 mg / kg  
132 body weight).

##### 133 **2.4.2.2.1 Biochemical analysis**

134 **Transaminases assay:** The animals' blood were collected in dry tubes, centrifuged at 3000  
135 rpm for 5 minutes and the sera were taken to evaluate enzymatic parameters of hepatic  
136 necrosis: Aspartate Amino Transferase (AST) and Alanine Amino Transferase (ALT) using  
137 kits (Cypress Diagnostics).

138 **Lipid peroxidation evaluation:** Animals liver pieces from treated animals were removed,  
139 ground in (10% w/v) Tris-HCl buffer (50Mm, pH 7.40), centrifuged at 6000 rpm for 10  
140 minutes, and Supernatants were used to evaluate lipid peroxidation[24].

##### 141 **2.4.2.2.2 Histopathology analysis**

142 Small fragments (approximately 0.2 x 0.2 cm) of liver were removed and fixed in formalin  
143 solution 10%[25]. They were dehydrated in solutions of increasing concentration of ethanol  
144 (70 to 100%) for 2 hours in each concentration. They were cleaned then in 2 xylene baths,  
145 infiltrated into 2 paraffin baths, and transferred to paraffin-filled molds. The sections of livers  
146 prepared by rotary microtome (Leitz 1512) were placed on clean slides and stained with  
147 Mayer's haematoxylin solution for 15 min, washed with water and alcohol 80% and mounted  
148 in eosin-phloxine solution. Finally, these assemblages of tissue slides were examined under  
149 an optical microscope.

#### 150 **2.5 In vitro experiments**

##### 151 **2.5.1 Trolox Equivalent Antioxidant Capacity (TEAC)**

152 ABTS radical cation decolorization assay was used to evaluate crude and phenolic  
153 extracts TEAC according to Guenné [20] with some modifications. ABTS radical cation  
154 (ABTS<sup>+</sup>) was produced by reacting aqueous ABTS stock solution (7 mM) with 2.45 mM  
155 potassium persulfate. The mixture was put down in dark at room temperature for 16 h before  
156 use. This mixture was diluted with ethanol to give an absorbance of 0.70 ± 0.02 units at 734  
157 nm using microplates **UV/visible light spectrophotometer** reader (Epoch 251465, Biotek  
158 Instruments, U.S.A.). 50 µL of diluted sample (1 g/mL in methanol) were added with 200 µL  
159 of fresh ABTS<sup>+</sup> solution and the absorbance was taken 15 min exactly after initial mixing.  
160 Trolox was used to produce the calibration curve (R<sup>2</sup> = 0.99) and the antioxidant capacity of  
161 extracts were expressed as mg Trolox Equivalent per g of extract.

##### 162 **2.5.2 Liver lipid peroxidation inhibition**

163 Crude and phenolic extracts lipid peroxidation (LPO) inhibitory activities were determined  
164 according to the 2-thiobarbituric acid method[26]. Ferrous dichloride (FeCl<sub>2</sub>) and H<sub>2</sub>O<sub>2</sub> were  
165 used to induce rat liver homogenate fats peroxidation. In this method 0.2 mL of extracts (1.5  
166 mg mL<sup>-1</sup>) was mixed with 1.0 mL of 1% liver homogenate in Tris-HCl buffer, then 50 µL of

167 FeCl<sub>2</sub> (0.5 mM) and 50 µL of H<sub>2</sub>O<sub>2</sub> (0.5 mM) were added. The mixture was incubated at 37°C  
 168 for 60 min, then 1.0 mL of trichloroacetic acid (15%) and 1.0 mL of 2-thiobarbituric acid  
 169 (0.67%) were added and the mixture was heated up in boiled water for 15 min. The  
 170 absorbance was recorded at 532 nm using spectrophotometer. Quercetin was used as the  
 171 positive controls.

172

### 173 3. STATISTICAL ANALYSIS

174 All results were expressed as mean ± standard deviations (SD). Tukey's test (one-way  
 175 ANOVA) was used to determine level of significance of all results obtained on XLSTAT 7.1.  
 176 Results were regarded as significant at p< 0.05.

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## 178 4. RESULTS AND DISCUSSION

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### 180 4.1. Extract toxicities

181 Through oral administration, the plant crude extract showed a lethal dose of 50% (LD<sub>50</sub>)  
 182 greater than 3000 mg / kg of body weight because on groups of six (06) mice no mortality  
 183 was observed after seventy-two hours (72 h) observation following extracts administration.  
 184 Through intraperitoneal administration, *C. americanum* hydro-ethanolic extract toxicities  
 185 values were presented in the following table (table 1).

186 **Table 1:** *C. americanum* toxicity by intraperitoneal voice

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Plant	Doses	Mice numbers used	Death numbers				% of of death at 72 H
			2H	24H	48H	72H	
<i>C. americanum</i>	75 mg/kg	06	00	00	00	00	00
	150 mg/kg	06	00	01	01	00	33,33
	200 mg/kg	06	00	02	01	01	66,66
	250 mg/kg	06	00	03	01	01	83,33
	400 mg/kg	06	00	04	02	00	100

188

189 The Log Probit paper plot of mortalities percentages based on log of dose determined *C.*  
 190 *americanum* LD<sub>50</sub> of 175 mg / kg of body weight. The line obtained has good validity  
 191 because LD<sub>50</sub> / DL<sub>1</sub> (2.18) is substantially equal to DL<sub>99</sub> / LD<sub>50</sub> (2.28) (with LD<sub>1</sub> = 80 mg / kg  
 192 of body weight and DL<sub>99</sub> = 400 mg / kg of body weight). The safety index of the extract is  
 193 DL<sub>99</sub> / DL<sub>1</sub> = 4.98 <5.

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### 196 4.2. Liver protection

#### 197 Enzymatic parameters of liver damage

198 The table 2 showed transaminases and lipids peroxidations values.

199 **Table 2:** Liver necrosis blood parameters

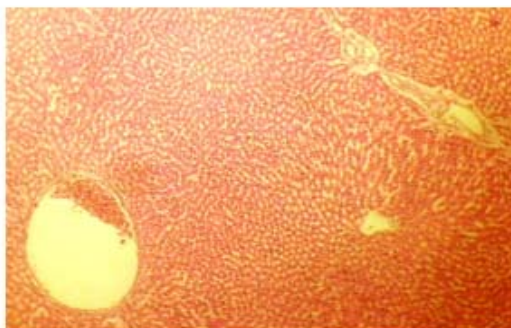
Samples	Liver weight/100 g body weight	ALAT (UI/L)	ASAT (UI/L)	lipid Peroxidation (mmol MDA/g of liver)
Control	2.67± 0.18 <sup>a</sup>	14.63± 5.71 <sup>a</sup>	20.20± 1.51 <sup>a</sup>	14.29± 0.23 <sup>a</sup>
Negative control	3.95± 0.14 <sup>d</sup>	61.96 ±13.50 <sup>c</sup>	98.16± 16.15 <sup>c</sup>	27.73± 4.13 <sup>c</sup>
Positive control	3.29± 0.21 <sup>b,c</sup>	26.80 ±14.79 <sup>a,b</sup>	49.41± 5.25 <sup>a, b</sup>	22.92±0.88 <sup>b,c</sup>
<i>C. americanum</i>	3.36± 0.32 <sup>b,c</sup>	58.67 ±12.62 <sup>b</sup>	60.46± 7.55 <sup>b,c</sup>	22.68±0.68 <sup>b,c</sup>

200 ALAT: Alanine Amino-Transferase; ASAT: Aspartate Amino-Transferase; MDA: MalonDiAldehyde. The results  
 201 presented in the table columns with the letters a- d are significantly different at P <0.05.

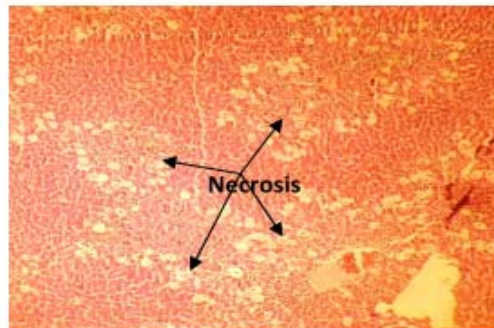
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*C. americanum* extract has a protective effect against the oxidative aggression of carbon tetrachloride on rat livers. This effect was inferior to the beneficial effect of sylimarin, which is the reference compound used in hepatic poisoning.

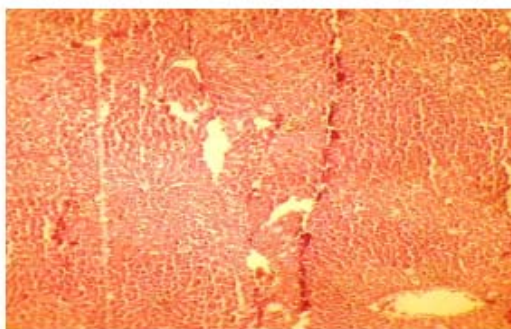
**Histopathology:** The presence of necrotic cells due to CCl<sub>4</sub> (hepatotoxic agent) action and these necrosis reduction by the sylimarin or *C. americanum* polyphenolic extract actions are represented by the **photo 1**.



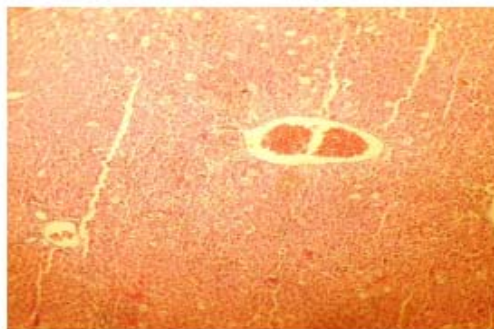
**Photo 1a:** Normal liver cut (X 10)



**Photo 1b:** CCl<sub>4</sub> liver treated cut (X 10)



**Photo 1c:** Sylimarin and CCl<sub>4</sub> liver treated cut (X 10)



**Photo 1d:** *C. americanum* and CCl<sub>4</sub> liver treated cut (X 10)

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**Photo 1:** Livers histopathology's analysis using photonic Microscope

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Carbon tetrachloride has caused hepatic necrosis (Photo 1 b) compared to normal liver (Photo 1a). Sylimarin significantly prevented the hepatic necrosis establishment (Photo 1c). This action was also borrowed by *C. americanum* (Photo 1d) extract but it remains less important than that of sylimarin.

#### **4.3 *In vitro* antioxidant capacities**

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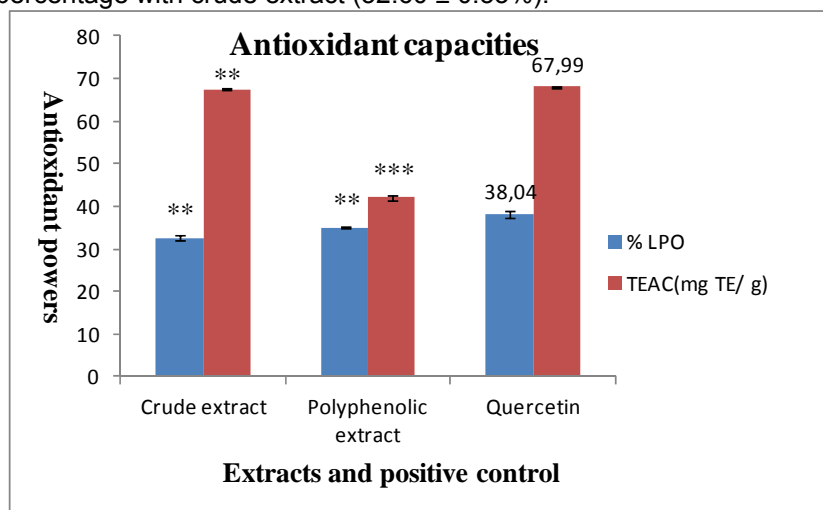
##### **4.3.1 Trolox Equivalent Antioxidant Capacity**

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The radical cation ABTS<sup>•+</sup> reducing power of the crude extract, polyphenol extract and quercetin are shown in Figure 1. This figure shows that the best reducing power was obtained with quercetin (67.99 ± 0.79 mg TE/ g) followed to the crude extract (67.53 ± 0.05 mg ET/ g) and the polyphenolic extract (42.01 ± 0.26 mg ET/ g).

### 223 4.3.2 Lipid peroxidation inhibition

224 Lipid peroxidation inhibition percentages of crude extract, polyphenolic extract and  
 225 quercetin are shown in Figure 1. The best inhibition percentage was obtained with quercetin  
 226 and the lowest percentage with crude extract ( $32.60 \pm 0.53\%$ ).

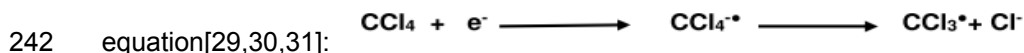


227 **Figure 1:** Crude and polyphenolic extracts *in vitro* antioxidants powers.  
 228 The values are mean ± S.E.M. (n=3 per test). \*\*p < 0.001 vs. quercetin and \*\*\*p<0.0001 vs. quercetin.  
 229  
 230

### 231 4.4. Discussion

232 The polyphenolic extract had low antioxidant capacity in TEAC and a high capacity in  
 233 lipid peroxidation compared to the crude extract of *Chrysanthellum americanum*. Our  
 234 previous studies have shown that this butanol fraction was richer in total phenolic ( $85.65 \pm$   
 235  $1.77$  against  $79.09 \pm 0.80$  GAE / 100 mg of extract), in flavonoids ( $24.03 \pm 0.88$  against  
 236  $13.54 \pm 0.44$  QE) compared to the crude extract[27]. Flavonoids are well known for their  
 237 electron receptor hence their capacity to break the chain of free radical oxidation  
 238 process[28].

239 It is well known and documented that a single dose of  $\text{CCl}_4$  administration to a rat produces  
 240 centrilobular necrosis and fatty degeneration of the liver. This action begins with an  
 241 activation of  $\text{CCl}_4$  and a production of  $\text{CCl}_3^\cdot$  radical compound according to the following



243 In our study, this condition was obtained and represented with the photo 1a with necrotic  
 244 cells. The richness of *C. americanum* polyphenolic extract<sup>14</sup> in phenolic compounds would  
 245 have the advantage of protecting the liver of rats from the oxidizing action of  $\text{CCl}_3^\cdot$ . These  
 246 polyphenols would inhibit  $\text{CCl}_4$  activation or reduce the  $\text{CCl}_3^\cdot$  radical to non-free radical  
 247 compounds. Luteolin 7-O-glucoside, a flavonoid from this plant extracts was well  
 248 known for its antioxidant ability. So, this molecule would has contributed to break  
 249 the radical reaction or activated liver antioxidant enzymes [32, 33]. Also, luteolin has  
 250 a strong anti-inflammatory effects could protect rat's liver against inflammation  
 251 induced by  $\text{CCl}_4$  [34].

252 The result observed in photo 1d (reduction of necrosis cells number) is explained by this  
 253 plant *in vitro* antioxidant activity ( $42.01 \pm 0.26$  mg ET / g and  $32.60 \pm 0.53\%$  inhibition of lipid  
 254 peroxidation) and also by the plant extract *in vivo* activities by reducing the blood level of  
 255 transaminases and malondialdehyde ( $22.68 \pm 0.68$  mmol MDA / g of liver weight) produced  
 256 by  $\text{CCl}_4$  injection action.



257 Some authors have cited *Chrysanthellum americanum* extract for the treatment of kidney  
258 calculi, cholelithiasis and also as a food additive because of its richness in  
259 protein[13,35,36,18]. These properties additional to his hepatoprotection confirm this plant  
260 massive use around the world.

261 In addition to primary usage of this plant extract in health care with metabolic origin, Mevy  
262 group[37] found that essential oils of this plant (caryophyllene oxide, hexa-2,4-dienol,  $\beta$ -  
263 caryophyllene,  $\alpha$ -pinene and verbenol) have antifungal potentials.

264 Our study showed that the hydroalcoholic extract of *C. americericum* has a very low oral  
265 route toxicity[38] and a high intraperitoneal route toxicity ( $LD_{50}\%$  = 175 mg / kg of body  
266 weight). This intraperitoneal toxicity can be explained by saponins (Chrysantheline A & B)  
267 presence in the plant extracts[39,40,41,42]. Fortunately, this plant is used traditionally by  
268 decoction and drink. Nevertheless, precautions are to be taken for people who would  
269 present lesions in their digestive tract.

270 The polyphenolic extract of *C. americanum* has a protective effect against intoxication  
271 through its antioxidant potential and has a beneficial effect on health. Also, the traditional  
272 use of this plant extract orally has virtually no toxicity.

273

#### 274 **4. CONCLUSION**

275 Our literature review on *Chrysanthellum americanum* showed that this species has  
276 flavonoids and saponins high content and strong antioxidant capacity.

277 This research has made a screening of *Chrysanthellum americanum* polyphenol extract  
278 effect on rats anti hepatotoxicity using  $CCl_4$  as hepatotoxicity agent.

279 The polyphenolic extract of *C. americanum* significantly prevented the oxidative aggression  
280 of carbon tetrachloride on rat liver. This beneficial action was manifested by the considerable  
281 necrotic cells number reduction and the decrease of transaminases and malondialdehyde  
282 serum levels. Our preview surveys near traditional phytotherapists of Burkina Faso central  
283 region had shown that this plant is used traditionally by decoction and per orally. This  
284 present study found also that this plant extract had a very low oral acute toxicity.

285 In short, our research confirms the benefits of *Chrysanthellum americanum* extracts used in  
286 traditional medicine for specific care of liver diseases.

287

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290 (IRSS/CNRST), Department of Medicine and Traditional Pharmacopoeia (MEPHATRA-PH)  
291 for these assistance and supervision during the experimentation. Rest In Peace Dr Tibiri.

292

#### 293 **COMPETING INTERESTS**

294



The authors declare no conflict of interest.

## AUTHORS' CONTRIBUTIONS

**“Conceptualization**, Samson Guenné, Nabère Ouattara, Noufou Ouédraogo and Adama Hilou; **Data curation**, Samson Guenné, Prosper T. Kinda, Nâg-Tiero Roland Meda, Alin Ciobica and Martin Kiendrebéogo; **Formal analysis**, Samson Guenné and Adama Hilou; **Investigation**, Samson Guenné, Nabère Ouattara, Noufou Ouédraogo and Adama Hilou; **Methodology**, Samson Guenné, Nabère Ouattara, André Tibiri and Adama Hilou; **Project administration**, Odile G. Nacoulma; **Resources**, Prosper T. Kinda; **Software**, Samson Guenné; **Supervision**, Adama Hilou and André Tibiri; **Validation**, Noufou Ouédraogo; **Writing – original draft**, Samson Guenné; **Writing – review & editing**, Samson Guenné, Nabère Ouattara, Nâg-Tiero Roland Meda, Noufou Ouédraogo, Prosper T. Kinda, Alin Ciobica, Adama Hilou, and Martin Kiendrebéogo. All authors read and approved the final manuscript.”

## ETHICAL APPROVAL

“All authors hereby declare that "Principles of laboratory animal care" (NIH publication No. 85-23, revised 1985) were followed, as well as specific national laws where applicable. All experiments have been examined and approved by the appropriate ethics committee”

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#### ABBREVIATIONS

- |   |   |
|---|---|
| 425 <b>ABTS</b> : 2,2'-azino bis (3                     | 438 <b>LD</b> : Lethal Dose                     |
| 426 ethylbenzothiazoline-6-sulphonate                   | 439 <b>LD<sub>50</sub></b> : Lethal Dose of 50% |
| 427 <b>ALT</b> : Alanine Amino Transferase              | 440 <b>LPO</b> : Lipid Peroxidation             |
| 428 <b>AST</b> : Aspartate Amino Transferase            | 441 <b>MDA</b> : malonedialdehyde               |
| 429 <b>C. amercericum</b> : <i>Chrysanthellum</i>       | 442 <b>MEPHATRA-PH</b> : Department of Medicine |
| 430 <i>amercericum</i>                                  | 443 and Traditional Pharmacopoeia               |
| 431 <b>CCl<sub>3</sub></b> : Carbon trichloride radical | 444 <b>NMRI</b> : Naval Medical and Research    |
| 432 <b>CCl<sub>4</sub></b> : Carbon tetrachloride       | 445 Institute                                   |
| 433 <b>FeCl<sub>2</sub></b> : Ferrous dichloride        | 446 <b>SD</b> : standard deviations             |
| 434 <b>GAE</b> : Gallic acid Equivalent                 | 447 <b>TE</b> : Trolox Equivalent               |
| 435 <b>HCl</b> : Hydrochloric acid                      | 448 <b>TEAC</b> : Trolox Equivalent Antioxidant |
| 436 <b>IRSS/CNRST</b> : Institute for Research in       | 449 Capacity                                    |
| 437 Health Sciences                                     | 450 <b>WHO</b> : World Health Organization      |

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