

# *In vitro* larvicidal effect of a hydroalcoholic extract from *Acacia cochliacantha* leaf against ruminant parasitic nematodes

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**Abstract** The aim of this study was to evaluate the *in vitro* lethal effect of a hydroalcoholic extract (HAE) from *Acacia cochliacantha* leaf against three gastrointestinal nematodes species (*Haemonchus contortus*, *H. placei* and *Cooperia punctata*) of domestic ruminants. The HAE was assessed using five concentrations: 100, 125, 175, 150 and 200 mg/ml; 0.5% Ivermectin was used as a positive control and distilled water, as negative control. The data were normalized using the square root and analysed with a completely randomized design through ANOVA analysis using the general lineal model (GLM) of the SAS program. The HAE tannin content was determined through spectrophotometry (UV-visible) and the other major phenols, were identified by chromatographic processes. The results showed an *in vitro* larvicidal activity of

the HAE against the three assessed nematode species with all assessed concentrations. A clear HAE increased concentration dependence effect was observed. The highest activity of the HAE was obtained at the highest concentration (close to 100%,  $P < 0.05$ ). This result was similar to the one obtained with Ivermectin. On the other hand, the chemical analysis of HAE showed the presence of tannins, caffeoyls and coumaroyl derivates and quercetin as the main compounds. The results suggest that the HAE from this plant species possess *in vitro* anthelmintic properties. The identified compounds in this study would good candidates for further *in vivo* researches.

**Keywords** *Haemonchus* · *Cooperia* · Tannins · Flavonoids · Nematodes · *Acacia cochliacantha*

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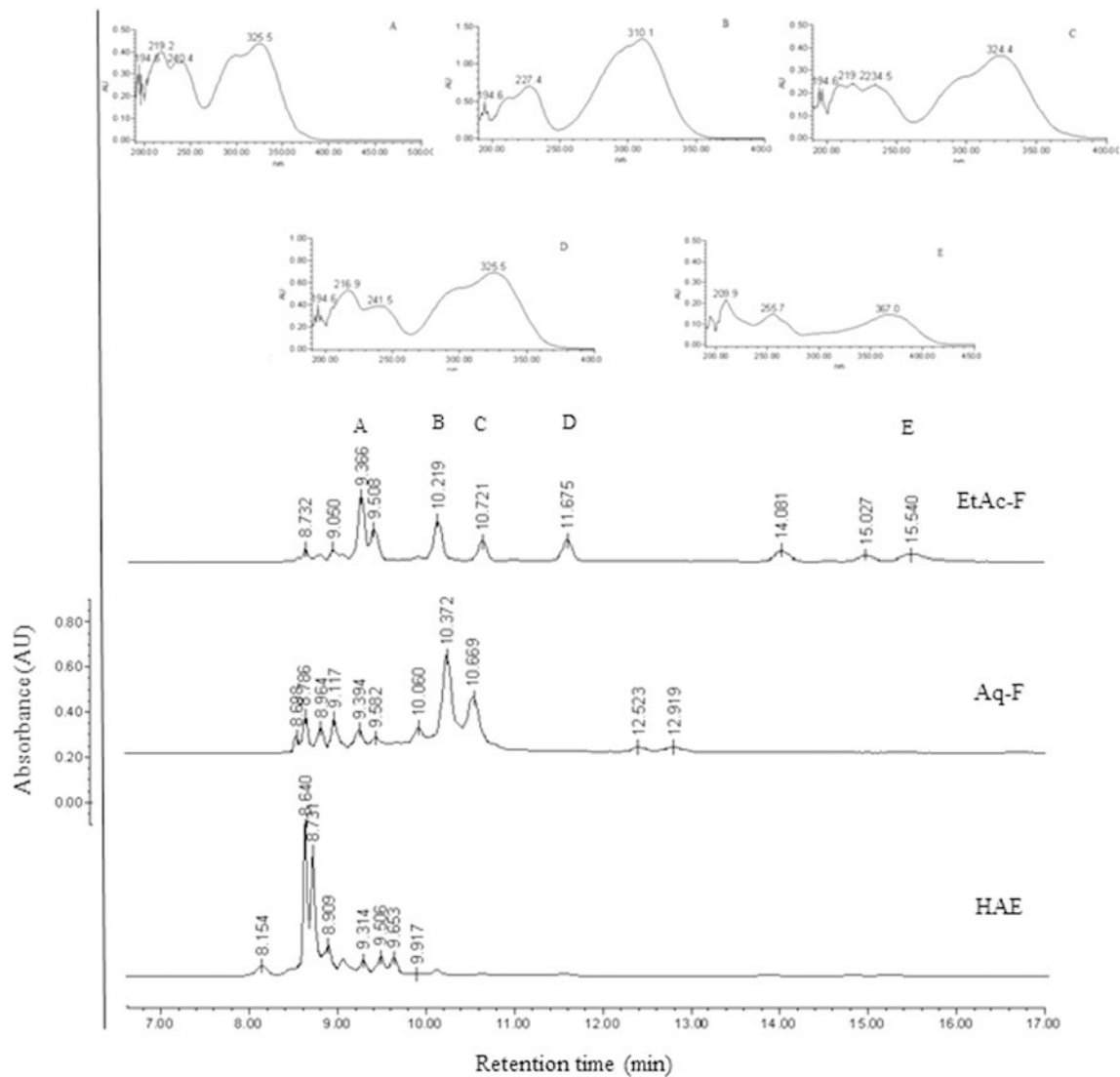
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## Introduction

Gastrointestinal nematode (GINs) parasitic infection is one the major health concern in the ruminant production. The excessive use of chemical anthelmintic drugs is a widespread practice in livestock production worldwide; although their continuous and frequent use triggers a serious problems of anthelmintic resistance (Jabbar et al. 2006; Muñoz-Lagunes et al. 2015). The use of plants with anthelmintic (AH) properties is considered as one possible method for controlling GINs in ruminants. A number of *in vitro* and *in vivo* studies, using plant extracts from Leguminosae family, have provided information of phenolic compounds such as tannins and flavonoids with AH activity (Olmedo-Juárez et al. 2014; Vargas-Magaña et al. 2014; von Son-de Fernex et al. 2015). *Acacia* is a large genus of the Fabaceae family, with about 1350 species. Most of the species belonging to the *Acacia* genus are rich in

57	secondary metabolites containing mainly condensed tannins	the protein- (PCT) and fiber- (FCT2) bound CT analyses were	102
58	and flavonoids (Seigler 2003; León-Castro et al. 2015). In	conducted following the technique reported by Porter et al.	103
59	some Mexican tropical areas, the leaves and fruits from	(1986). Purification was performed using a Shepadex LH-20	104
60	<i>Acacia cochliacantha</i> are found scattered in pastures and liv-	column, as described by Hedqvist et al. (2000).	105
61	ing fences, where ruminants harvest the leaves and fruits to		
62	feed themselves during the dry season. The secondary metab-		
63	olites identified in this plant species are condensed tannins as		
64	main compounds (Olivares-Pérez et al. 2011). <i>Acacia</i>		
65	<i>cochliacantha</i> showed an <i>in vivo</i> anthelmintic effect on		
Q2 66	<i>H. contortus</i> (León-Castro et al. 2016) but more information		
67	regarding the metabolites involved and the effect on other		
68	parasitic stages is needed. Thus, the objective of this study		
69	was to evaluate the <i>in vitro</i> effect of a hydroalcoholic extract		
70	of <i>A. cochliacantha</i> leaves against infective larvae (L <sub>3</sub> ) of		
71	three gastrointestinal parasite species ( <i>Haemonchus contortus</i> ,		
72	<i>Cooperia punctata</i> and <i>Haemonchus placei</i> ).		
73	<b>Materials and methods</b>		
74	<b>Plant material</b>		
75	<i>Acacia cochliacantha</i> leaves Humb. & Bonpl. (Cubata) were		
76	collected from a Salitre Palmarillos village, Amatepec		
77	Municipality, in the State of Mexico, Mexico (18°43'28.4" N,		
78	100°17'03.5" W). Plants were collected between March and		
79	April 2016. The plant was taxonomically identified by Prof.		
80	Rafael Torres-Colin and deposited at the Herbario Nacional de		
81	México at Universidad Nacional Autónoma de México, México,		
82	City (Voucher code number OD07042016). Fresh material was		
83	washed and dried at room temperature in the dark for one week.		
84	Plant leaves were milled using an electrical miller (Wiley mill,		
85	TS3375E15 model), so as to reach a size of 4–6 mm.		
86	<b>Preparation of the hydroalcoholic extract</b>		
87	One kg of dried and ground leaves were used to obtain the		
88	extract by maceration with an aqueous methanol solution		
89	(70%, 1:10 ratio, <i>w/v</i> ) at room temperature during 24 h. The		
90	liquid extract was paper-filtered and the residual solvent was		
91	evaporated using a rotary evaporator (Heidolph Laborota		
92	4000, Germany) under reduced pressure at 50–60 °C to obtain		
93	a semisolid extract, which was finally freeze-dried to get 120 g		
94	(12%). The dry extract was stored at –40 °C until bioassays		
95	and phytochemical analysis.		
96	<b>Condensed tannin content</b>		
97	The hydroalcoholic extract (HAE) was analysed to determine		
98	the total condensed tannin content (TCT) by using of the		
99	butanol-HCL method (López et al. 2004); the <i>Lysiloma</i>		
100	<i>acapulcensis</i> free condensed tannins (FCT) were used as in-		
101	ternal standards (Olmedo-Juárez et al. 2014). The free (FCT1),		
		The hydroalcoholic extract (HAE, 60 g) was processed for bi-	107
		partition via liquid-liquid chromatography using water/ethyl	108
		acetate solvents (600 mL each, Merck, Germany). Two frac-	109
		tions, an aqueous fraction (Aq-F) and an organic fraction	110
		(EtAc-F) were obtained. The solvents in both fractions were	111
		eliminated using low-pressure distillation. Fraction yields	112
		were as follows: Aq-F = 58.1 g and EtAc-F = 1.92 g.	113
		Chromatographic analysis was developed by HPLC using a	114
		Waters 2695 separation module HPLC system equipped with	115
		a Waters 996 photodiode array detector and Empower Pro	116
		software (Waters Corporation, USA). Chemical separation	117
		was achieved in a supelcosil LC-F column (4.6 mm × 250 mm	118
		i.d., 5-µm particle size) (Sigma-Aldrich, Bellefonte, USA).	119
		The mobile phase consisted of 0.5% trifluoroacetic acid aque-	120
		ous solution (solvent A) and acetonitrile (solvent B). The gra-	121
		dient system was obtained as follows: 0–1 min, 0% B; 2–	122
		3 min, 5% B, 4–20 min, 30% B; 21–23 min, 50% B 14–	123
		15 min; 24–25 min, 80% B; 26–27,100% B; 28–30 min, 0%	124
		B. The flow rate was maintained at 0.9 mL/min and the injec-	125
		tion volume was 10 µL. The absorbance was measured at	126
		330 nm. Caffeic acid and coumaric acid were identified by	127
		comparison of the retention times and UV spectra with the	128
		reference standards (Sigma-Aldrich, St Louis Mo, USA).	129
		Other caffeoyl and coumaroyl derivatives were established	130
		based on their UV spectra (Wagner and Bladt 2001).	131
		<b>Biological material</b>	132
		<i>Haemonchus contortus</i> infective larvae (L <sub>3</sub> ) (strain,	133
		INIFAP), were obtained from a donor sheep artificially in-	134
		fectured with 350 L <sub>3</sub> larvae per kg BW. Likewise, infective	135
		larvae from <i>H. placei</i> (wild strain) and <i>C. punctata</i> (Cp de	136
		Fernex-MEX strain) were obtained from two young cattle.	137
		Faecal cultures were prepared by mixing faeces with poly-	138
		styrene particles in plastic bowls. Water was added to the	139
		faecal cultures and mixed with a wooden spoon for	140
		obtaining an adequate oxygenation to promote a better	141
		egg hatching. The faecal cultures were covered with foil	142
		and incubated for 7 days at room temperature (25–31 °C).	143
		The infective larvae were extracted from faecal material	144
		using the Baermann funnel technique (Liebano-Hernández	145Q3
		2004). The L <sub>3</sub> were cleaned by density gradient (40%	146
		Sacharose) and centrifugation; the larvae were later	147
		exsheathed with sodium hypochlorite at 0.187%. Finally,	148
		the exsheathed larvae were used for the mortality assay.	149



**Q4** **Fig. 1** HPLC chromatogram of a hydroalcoholic extract (HAE), an aqueous fraction (Aq-F) and an ethyl acetate fraction (EtAc-F) indicating the presence of phenols (showing UV-spectral); as caffeoyl

derivatives displayed  $\lambda_{max} = 325$  nm (peaks A, C, D); coumaroyl derivatives gave  $\lambda_{max} = 310$  nm (peaks B) and quercetin displayed  $\lambda_{max} = 360$  nm (peak E)

**150 Larval mortality assay**

151 The assay was carried out using 96-well micro-titration plates  
 152 ( $n = 12$ ) for each treatment. Treatments were designed with the  
 153 HAE concentration at 100, 125, 150, 175 and 200 mg/ml, re-  
 154 spectively. Each treatment was tested using a negative control  
 155 (water) and anthelmintic (0.5% ivermectin) as the positive  
 156 control. Fifty microliters of an aqueous suspension containing  
 157 150 nematode (*H. contortus*, *H. placei*, *C. punctata*) larvae  
 158 were distributed in each well. Then, 50- $\mu$ l aliquots of the  
 159 extract and controls were added to each well. The plates were  
 160 incubated at room temperature (18–25 °C) during 48 h. Ten  
 161 aliquots of 10  $\mu$ l were taken from each well to count dead or  
 162 living larvae; the larval mortality was assessed if mobility was  
 163 not observed during 20 s. When larvae remained motionless

but their aspect caused confusion about if they were death or  
 164 alive; a physical stimulus was applied touching their coat with  
 165 a metal needle and the final decision was based on their moti-  
 166 lity. Finally, the larval mortality percentage was determined  
 167 using the following formula: % mortality = [(number of living  
 168 larvae)/ (number of dead larvae + number of living  
 169 larvae)]\* 100.  
 170

**Statistical analysis**

171  
 172 The data of larval mortality were normalized using the square  
 173 root transformation and it was analysed through a completely  
 174 randomized design through ANOVA analysis using the gen-  
 175 eral lineal model (GLM) of the SAS program. Differences  
 176 among means were assessed by the Tukey's test. Likewise,  
 177

**Q5** t1.1 **Table 1** Mortality percentages of  
t1.2 infective larvae (L<sub>3</sub>) of three  
t1.3 different ruminant parasitic  
t1.4 nematodes exposed to an *Acacia*  
t1.5 *cochliacantha* hydroalcoholic  
t1.6 extract at different concentrations

Mortality percentage of infective larvae (%)				
Treatment	<i>Haemonchus contortus</i> (INIFAP strain)	<i>Cooperia punctata</i> (de Fernex-MEX strain)	<i>Haemonchus placei</i> (wild strain)	
Distilled water (C <sup>-</sup> )	1.00 <sup>f</sup>	0.75 <sup>d</sup>	0.00 <sup>c</sup>	
Ivermectin (C <sup>+</sup> )	100.00 <sup>a</sup>	100.00 <sup>a</sup>	100.00 <sup>a</sup>	
<i>A. cochliacantha</i> hydro-alcoholic extract (mg/ mL)				
200	97.75 <sup>ab</sup>	99.25 <sup>a</sup>	97.00 <sup>a</sup>	
175	89.50 <sup>b</sup>	77.50 <sup>b</sup>	92.75 <sup>a</sup>	
150	73.00 <sup>c</sup>	37.00 <sup>b</sup>	76.00 <sup>b</sup>	
125	46.25 <sup>d</sup>	10.00 <sup>c</sup>	39.00 <sup>c</sup>	
100	25.00 <sup>e</sup>	8.50 <sup>c</sup>	16.00 <sup>d</sup>	
SEM	1.75	2.90	2.20	

Means with different letters in the same column represent statistical differences  $P < 0.05$   
SEM standard error of mean

177 the lethal concentrations (LC<sub>50</sub> and LC<sub>90</sub>), were estimated  
178 through a Probit analysis (SAS 2006).

100% were achieved at the HAE highest concentration 193  
(200 mg/ml). On the other hand, the HAE mean lethal concen- 194  
trations (LC<sub>50</sub> and LC<sub>90</sub>) for the three nematode assessed spe- 195  
cies are show in Table 2. The HA extract LC<sub>50</sub> and LC<sub>90</sub> against 196  
*H. placei*, were: 126.53 and 172.59 mg/ml, respectively; mean- 197  
while, these values were 129.39 and 177.88 mg/ml, for 198  
*H. contortus*, respectively and 136.90 and 174.7 mg/ml for 199  
*C. punctata*, respectively. 200

179 **Results**

180 **Condensed tannin content and other main compounds**

181 The TCT, PCT and FCT2 bound resulted in 140.0, 26.0 and  
182 36.6 g/kg of dry matter, respectively. On the other hand, the  
183 chromatographic analysis in the HAE revealed the presence of  
184 caffeoyl derivates (Fig. 1 ACD) and coumaroyl derivatives  
185 (Fig. 1B) as well as some flavonoids (Fig. 1E) such as quer-  
186 cetin as the main compounds.

**Discussion** 201

The use of plants with medicinal properties represents a sus- 202  
tainable alternative for controlling diseases with important 203  
repercussions on livestock health, such as internal parasitic 204  
infections. The leaves of some leguminous trees like 205  
*Lysiloma acapulcensis* and *Leucaena leucocephala* have 206  
shown possessing anthelmintic activity against ruminant par- 207  
asitic nematodes in a number of *in vitro* and *in vivo* studies 208  
(Mejía-Hernández et al. 2014; Olmedo-Juárez et al. 2014; 209  
von Son-de Fernex et al. 2015; García-Hernández et al. 210  
2017). It is common to find a miscellaneous GIN fauna 211

187 **Infective larvae (L<sub>3</sub>) mortality test**

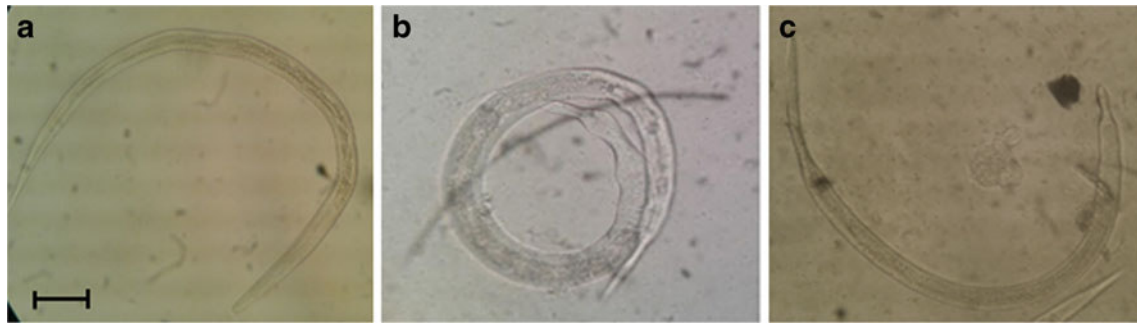
188 Table 1 shows the results of the GIN mortality percentages from  
189 cattle and sheep exposed to the extract at the different assessed  
190 concentrations and at their proper controls. A larvicidal effect  
191 ( $P < 0.05$ ) was observed in all the nematode species as well as a  
192 concentration/dependence. Mortality percentages close to

t2.1 **Table 2** Fifty and ninety lethal  
t2.2 concentrations of a hydroalcoholic  
extract from *Acacia cochliacantha*  
leaves against *Haemonchus*  
*contortus*, *H. placei* and *Cooperia*  
*punctata* infective larvae after 48 h  
t2.5 *in vitro* exposure

Nematode specie	LC <sub>50</sub>	95% CI limits		LC <sub>90</sub>	95% CI limits	
		Lower	Upper		Lower	Upper
<i>Haemonchus contortus</i>	127.39	123.99	130.33	177.88	172.90	183.77
<i>Haemonchus placei</i>	126.53	121.26	131.17	172.59	167.53	178.33
<i>Cooperia punctata</i>	136.90	134.61	139.06	174.07	170.79	177.84

Values are expressed as mg/ml  
CI confidence interval

t2.3



**Fig. 2** Photographies taken through an optical microscope showing the aspect of *Haemonchus contortus* infective larvae (L<sub>3</sub>) (40 x): **a** Normal larvae (control), **b** and **c** infective larvae after 48 h exposure to an *Acacia cochliacantha* hydroalcoholic extract. Bar scale (40 μM, **—**)

212 infecting grazing animals simultaneously. However, some  
 213 genera/species are more pathogenic than others. The GIN  
 214 *Haemonchus contortus*, *H. placei* and *C. punctata*, are con-  
 215 sidered as the main genera of parasitic nematodes affecting  
 216 ruminants under tropical grazing conditions (Howell et al.  
 217 2008; Vlaminck et al. 2015). The present research demon-  
 218 strated that the HAE from *A. cochliacantha* leaves had an  
 219 important larvicidal effect against the infecting larvae L<sub>3</sub> of  
 220 three different nematode species. Such effect is likely related  
 221 with a secondary metabolite profile, especially associated  
 222 with condensed tannins (Brunet and Hoste 2006; Martínez-  
 223 Ortiz-de-Montellano et al. 2013; Williams et al. 2014).  
 224 Nevertheless, Klongsiriwet et al. (2015) demonstrated that  
 225 tannins are not the only plant secondary metabolites respon-  
 226 sible for affecting the gastrointestinal nematodes of rumi-  
 227 nants; these authors reported a synergism of tannins with  
 228 other compounds, such as flavonoids, which enhance their  
 229 nematocidal effect. In the present study, some phenols such  
 230 as flavonoids and coumaroyl and caffeoyl derivates were  
 231 identified through chromatographic techniques (Fig. 1).  
 232 These compounds could also be related to the biological ac-  
 233 tivity of this plant. In another study, an anthelmintic effect of  
 234 quercetin and caffeic acid obtained from *L. leucocephala*  
 235 leaves was found through a bio-guided egg hatching inhibi-  
 236 tion assay (von Son-de Fernex et al. 2015). On the other  
 237 hand, significant structural changes on the larvae bodies were  
 238 observed (Fig. 2). Such morphological changes were ob-  
 239 served in the larvae exposed to the two highest HAE concen-  
 240 trations (175 and 200 mg/ml). A slimming of either the an-  
 241 terior and posterior parts of the larvae bodies was observed in  
 242 most of the HAE exposed larvae at these concentrations. The  
 243 slimmed extremes of the larval body looked like finger-shape  
 244 (Fig. 2b, c). Unfortunately, in our study was not possible to  
 245 identify the metabolite responsible of this structural change.  
 246 In another study, some phenols such as caffeoyl and  
 247 coumaroyl derivates as well as the flavonoid quercetin were  
 248 identified as responsible for inhibiting the *H. contortus* egg  
 249 hatching (Castillo-Mitre et al. 2016).  
 250 According to the above-explained facts, the larvicidal ef-  
 251 fects of the HAE in our study could be related to those

identified metabolites; although this will need to be demon- 252  
 strated in future studies. 253

**Conclusion** 254

The results of this research show that the HAE of 255  
*A. cochliacantha* leaves possess larvicidal properties 256  
 against *H. contortus*, *H. placei* and *C. punctata* infective 257  
 larvae. Thus, this plant species could be an option for the 258  
 control of nematode infestations in ruminants under an 259  
 environment-sustainable approach. Nevertheless, *in vivo* 260  
 studies with experimental cattle infected with GINs are 261  
 required in order to evaluate the effect. 262

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**Compliance with ethical standards** 266  
 267

**Competing interests** The authors declare that they have no competing 268  
 interests. 269

270

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