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Dhakal, Suraj; Meyling, Nicolai Vitt; Williams, Andrew Richard; Mueller-Harvey, Irene; Fryganas, Christos; Kapel, Christian Moliin Outzen; Fredensborg, Brian Lund

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Author: Suraj Dhakal Nicolai V. Meyling Andrew R.
Williams Irene Mueller-Harvey Christos Fryganas Christian
M.O. Kapel Brian L. Fredensborg

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apply to the journal pertain.
Highlights:

- There was a concentration dependent inhibitory effect of condensed tannins on cysticercoid excystation tested in vitro.

- Condensed tannins from hazel nut extract were the most potent to inhibit cysticercoid excystation, followed by pinebark and white clover.

- Anti-excystation activity appeared to be positively linked to the presence of procyanidin tannins.

- The in vivo treatment with hazelnut reduced cysticercoid establishment.

- Host-parasite model employing the flour beetle Tenebrio molitor (Coleoptera) and the rat tapeworm Hymenolepis diminuta (Cestoda) invertebrate model could be useful for a first screening of potentially interesting compounds.
Efficacy of condensed tannins against larval *Hymenolepis diminuta* (Cestoda) *in vitro* and in the intermediate host *Tenebrio molitor* (Coleoptera) *in vivo*

Suraj Dhakal\(^a\), Nicolai V. Meyling\(^a\), Andrew R. Williams\(^b\), Irene Mueller-Harvey\(^c\), Christos Fryganas\(^c\), Christian M. O. Kapel\(^a\), Brian L. Fredensborg\(^a\)*

\(^a\) Department of Plant and Environmental Sciences, Faculty of Science, University of Copenhagen, Frederiksberg C, Denmark.

\(^b\) Department of Veterinary Disease Biology, Faculty of Health and Medical Sciences, University of Copenhagen, Frederiksberg C, Denmark.

\(^c\) Chemistry and Biochemistry Laboratory, School of Agriculture, Policy and Development, University of Reading, Reading, United Kingdom.

* Corresponding author at: Department of Plant and Environmental Sciences, Faculty of Science, University of Copenhagen, Thorvaldsensvej 40, Room 70-3-B324, Frederiksberg C, Copenhagen, Denmark. Tel.: +45 35 33 26 76, E-mail address: blf@plen.ku.dk (Brian L. Fredensborg).
Abstract

Natural anti-parasitic compounds in plants such as condensed tannins (CT) have anthelmintic properties against a range of gastrointestinal nematodes, but for other helminths such effects are unexplored. The aim of this study was to assess the effects of CT from three different plant extracts in a model system employing the rat tapeworm, *Hymenolepis diminuta*, in its intermediate host, *Tenebrio molitor*. An in vitro study examined infectivity of *H. diminuta* cysticercoids (excystation success) isolated from infected beetles exposed to different concentrations of CT extracts from pine bark (PB) (*Pinus sps*), hazelnut pericarp (HN) (*Corylus avellana*) or white clover flowers (WC) (*Trifolium repens*), in comparison with the anthelmintic drug praziquantel (positive control). In the in vitro study, praziquantel and CT from all three plant extracts had dose-dependent inhibitory effects on cysticercoid excystation. The HN extract was most effective at inhibiting excystation, followed by PB and WC. An in vivo study was carried out on infected beetles (measured as cysticercoid establishment) fed different doses of PB, HN and praziquantel. There was a highly significant inhibitory effect of HN on cysticercoid development (p=0.0002). Overall, CT showed a promising anti-cestodal effect against the metacestode stage of *H. diminuta*.

Keywords:

Condensed tannins, praziquantel, cestodes, invertebrate-parasite model.
1. Introduction

For decades, parasite control in livestock has relied intensively on prophylactic treatment with synthetic anthelmintics, but increasing resistance to such drugs and consumer requests for organic animal products increases the need for alternative control strategies. Bioactive plants may offer potential alternatives for parasite control in vertebrates (Waller and Thamsborg, 2004). Condensed tannins (CT) are a group of secondary metabolites commonly found in tropical and temperate plants (Jansman, 1993). They vary widely in their molecular weights and the identity of the monomeric flavan-3-ol units, which make up the tannin polymers. Procyanidins (PC) consist of catechin or epicatechin, whereas prodelphinidins (PD) are comprised of gallocatechin or epigallocatechin flavan-3-ols (Williams et al., 2014). More complex CT structures may occur as heteropolymers (Mueller-Harvey and McAllan, 1992; Molan et al., 2003).

Several laboratory and field experiments have shown that plant CT may control gastrointestinal nematodes (Hoste et al., 2006; Novobilský et al., 2011; Novobilský et al., 2013). Besides anthelminthic properties, these bioactive plant products can also have beneficial effects on animal health and production (Hoskin et al., 2000; Ramírez-Restrepo et al., 2004; Hoste et al., 2005; Hoste et al., 2006), and reduce the level of host infection (Hoste et al., 2012). Although, in vitro anthelminthic efficacy of natural plant cysteine proteinases has also been reported against excysted scolices and adult worms of the rodent cestodes *Hymenolepis diminuta* and *Hymenolepis microstoma* (Mansur et al., 2014), the effect of CT against helminth taxa other than nematodes has not yet been investigated.
Due to close resemblance of drug effects between animals and humans (Lin, 1995), mammals are often used in pre-clinical pharmacological and toxicological assessment of new compounds (Baumans, 2004). International awareness on animal experimentation has enforced the focus on the “3Rs” to Reduce, Replace and Refine (nc3rs, 2014). An insect model could present an alternative to a range of experimental studies, e.g. as a model for human microbiology (Tan, 2002; Kavanagh and Reeves, 2004) and immunology (Pursall and Rolff, 2011). Further, invertebrate models may simplify and reduce costs of laboratory maintenance (Scully and Bidochka, 2006; Vokřál et al., 2012) and ease concerns associated with animal experimentation (Kemp and Massey, 2007).

In the present study, a host-parasite model employing the flour beetle *Tenebrio molitor* (Coleoptera) and the rat tapeworm *Hymenolepis diminuta* (Cestoda) was used to investigate the anti-cestodal effects of CT in three different plant extracts. *Tenebrio molitor-H. diminuta* is a well-known host-parasite model for studies on ecological and evolutionary host-parasite relationships (Shostak, 2014), and Woolsey, (2012) studied its potential for pre-clinical screening of anthelmintics (praziquantel, levamisole hydrochloride and mebendazole).

In the natural lifecycle, rats excrete infective *H. diminuta* eggs which are ingested by the flour beetles where they encyst as larvae (cysticercoid) in the hemocoel (Burt, 1980). The life cycle completes when an infected beetle is eaten by a rat, in which the cysticercoid excysts, attaches to the intestinal wall and develops into an egg producing tapeworm (Chappell et al., 1970). Since the development of egg into cysticercoid and its excystation plays a crucial role in maintaining the life-cycle of *H. diminuta,*
interruption of these processes with CT would indicate that CT contains anti-cestodal
properties.

The objective of the present study was to assess the anti-cestodal effects of three
different CT types against cysticercoids of *H. diminuta* both freely exposed (*in vitro*)
and within their intermediate host (*in vivo*), at a range of concentrations.

2. Materials and Methods

The *in vitro* experiments were performed with cysticercoids dissected from
experimentally infected beetles. The effect of CT was measured as a reduction of
excystation of cysticercoids, which serves as an important measure (proxy) for
infectivity to rats. The *in vivo* study was conducted in live beetles in order to measure
the establishment of cysticercoids in the presence of CT.

2.1. Condensed tannins and praziquantel

Condensed tannins were extracted and purified from three different plant sources. These
were pine bark (PB) (*Pinus sp*), hazelnut pericarp (HN) (*Corylus avellana*) and white
clover flowers (WC) (*Trifolium repens*). Most tannin-rich plants contain complex mixtures
of procyanidins (PC) and prodelphinidins (PD), however our previous work has
demonstrated that these three plants contain narrower tannin profiles, i.e. mainly PC or PD
(Williams et al. 2014). Therefore, we used these plants as a source of well-defined model
tannins that would allow us to investigate whether the molecular structure of the tannins
influenced possible anti-parasitic activity. Tannins were extracted and analyzed as
previously described (Williams et al., 2014). Briefly, 50 g of plant material was extracted
with acetone/water (7:3; v/v) at room temperature, concentrated and freeze-dried. Tannin analysis was carried out by thiolytic degradation of the polymers and subsequent HPLC analysis of the reaction products, providing information on CT content in the extract, the mean degree of polymerization (mDP, i.e. average CT polymer size) and the PC/PD ratio (Williams et al., 2014). Pine bark contained 50.8 g CT/100 g extract, with an mDP value of 2.5 and a PC/PD ratio of 64.2. Hazelnut pericarp contained 73.8 g CT/100 g extract with an mDP value of 9.6 and a PC/PD ratio of 79.5. White clover contained 33.8 g CT/100 g extract with an mDP value of 4.4 and, in contrast to the other extracts, its tannins were almost exclusively comprised of PD, i.e. the PC/PD ratio was 0.8/99.2 (Williams et al., 2014). The well-known anti-cestodal drug praziquantel (99.7 %, VETRANAL™) was used as a positive control for both in vitro and in vivo studies.

2.2. Management of the beetles

_Tenebrio molitor_ larvae (obtained from Avifauna ApS, Denmark) were propagated in plastic containers (30×21×20 cm), placed in a dark incubator (26 °C), and provided with fresh oatmeal and fresh slices of potato. The potato slices were changed twice a week. After 2 weeks of incubation, pupae started to develop and these were then transferred into another plastic container (30×21×20 cm) and kept in a dark incubator (26 °C) until emergence of adults. Newly emerged adults were transferred to new plastic containers (30×21×20 cm) twice a week and held under the conditions described above.
2.3. Infection of beetles

Feces was collected from *H. diminuta* infected rats (*Rattus norvegicus* – Wistar strain) stabled at the Veterinary Institute, Technical University of Denmark (Animal permission no. 2010/561-1914 –section C10) and stored at 10 °C until use (two weeks maximum). Ten g of fecal pellets were soaked 1 h in 25 ml of tap water and then stirred with a wooden stick to make a uniform paste. The fecal paste was poured through a double layer of cotton gauze (1×1 mm pore size) into a 200 ml plastic cup, and the gauze was rinsed thoroughly with approximately 75 ml of tap water to increase egg recovery. The resulting suspension was equally transferred into two 50 ml centrifuge tubes. The tubes were centrifuged (Universal 16R) at 1148 g for 7 min. The supernatant was removed and the sediment was again stirred with a wooden stick. This fecal paste containing the *H. diminuta* eggs was used to infect the beetles.

Before administration of the paste, a group of 50 beetles were left without feed for 72 h in plastic containers (30×21×20 cm) with filter paper at the base, and stored in a dark incubator (26 °C). For infection, a 10 µl fecal suspension was deposited on a coverslip (1.5 × 1.5 cm) placed on filter paper in a series of petri dishes (5.5 cm diameter, 1.42 cm depth). A starved beetle was placed inside each Petri dish covered by a lid, and placed in a dark room for an hour. For assessment of eventual evaporation of the fecal suspension, one petri dish setup (without a beetle) was left an hour. Only beetles that had consumed the entire 10 µl of fecal suspension after 1 h were considered successfully inoculated and were included in the experiments.

2.4. *In vitro* study with praziquantel and pine bark extract
At 15 days post inoculation, beetles were dissected and cysticercoids were recovered using a Pasteur pipette under a dissection microscope (40×). A maximum of 10 cysticercoids (first observed) from each beetle were transferred to a watch-glass (33 mm diameter, 7 mm deep) containing phosphate buffered saline (PBS) and a total of 80 cysticercoids were collected. From these, 10 cysticercoids (first observed) were placed in each of 8 wells (2 wells from 4 different 48 multi-well plastic plates). Then each well was treated with 150 µl of either praziquantel dissolved in 2 % dimethyl sulfoxide (DMSO) with a final concentration of 10^{-2} (high concentration), 10^{-3} (medium concentration) or 10^{-4} mg/ml (low concentration), or CT extracts from PB dissolved in Milli-Q™ water at a final concentration of 0.1, 0.5 or 2.5 mg CT/ml. Control consisted of 2 % DMSO or Milli-Q™ water. All plates were subsequently kept in an incubator at 37 °C for 1 h.

After 1 h of incubation, the 10 cysticercoids along with the respective treatment solution from each well were transferred separately to a watch glass and the treatment solution was then removed with a Pasteur pipette under a dissection microscope. One ml of HCl-pepsin solution [2 ml 37 % HCl, 20 ml warm 0.9 % saline, 0.8 g pepsin powder from porcine gastric mucosa (1:2500, Sigma Life Science)] was added and placed in an incubator (37 °C). After 10 min of incubation, all the HCl-pepsin solution was removed. The cysticercoids were washed three times with 1 ml warm (37 °C) PBS and 1 ml of trypsin-taurocholate solution [0.1 g sodium taurocholate hydrate powder, 0.1 g trypsin powder from porcine pancreas, (97 %, Sigma Life Science), 10 ml warm PBS] was added to the watch glass and placed in the incubator at 37 °C for 2.5 h. The cysticercoids were then observed under the dissection microscope (40×) and recorded.
as excystated (complete evagination and emergence of scolex and body part from the
cyst) or non excystated (absence of the above) (Roberts and Janovy, 2008). This
experiment was repeated five times.

2.5. *In vitro* study with pine bark, hazelnut and white clover extracts

Concentrations of PB, HN, and WC extracts were adjusted such that each extract
contained equal final concentrations of CT in the assay. Three different concentrations:
2.5 (high concentration), 0.25 (medium concentration) and 0.025 mg CT/ml (low
concentration) in Milli-Q™ water from each were prepared. Pure Milli-Q™ water was
used as a control. The procedure was as described above (see: *In vitro* study of
praziquantel and PB) and was repeated five times.

2.6. *In vitro* condensed tannin depletion assay

As the CT extracts used in this experiment were not 100 % pure, CT depletion
experiments were performed, to investigate whether inhibition of cysticercoid
excystation was exclusively due to the effect of CT. A total volume of 250 µl solution
with concentration 2.5 mg CT/ml of Milli-Q™ water was prepared separately from
three types of CT extracts (PB, HN, and WC). For each solution, 12.5 mg of
polyvinylpolypyrrolidone (PVPP) (at a dose rate of 50 mg PVPP/ml of solution) was
added to precipitate CT, and was incubated (4°C) overnight. After centrifugation at
3000 g for 5 min, supernatant (CT depleted extract) was removed and used in the test
assay (Novobilský et al., 2011). As a control, each CT solution was also incubated
(4°C) overnight. The above procedure (see: *In vitro* study of praziquantel and PB) was then followed and was repeated three times.

2.7. *In vivo* study with praziquantel, pine bark and hazelnut extracts

Eighty uninfected beetles (7-14 days after eclosion) were randomly selected and depleted feed for 72 h as described above. Starved beetles were then randomly allocated into 8 groups, each with 10 beetles. The beetles of each group were presented individually to a droplet of 5 µl containing one of the following treatments: praziquantel (25, 50 and 100 mg/kg body weight of beetle), or PB (125, 250 and 500 mg CT/kg body weight of beetle), or 2 % DMSO, or Milli-Q™ water for 15 min. Doses were formulated by measuring the average weight (+SE) of the beetles (103 ± 4.4 mg), which was calculated by weighing randomly allocated 25 beetles in 6 different groups. After 15 min, beetles that consumed the entire treatment solution were infected and maintained individually as described earlier (see: *In vitro* study).

After 15 days of incubation, 4 beetles were selected randomly from each treatment group for quantification of the establishment of cysticercoids. All cysticercoids in the haemocoele of the beetle were counted and recorded. The experiment for praziquantel and PB was repeated five and three times, respectively.

A separate study tested the effects of HN on cysticercoid establishment. A single dose of HN (500 mg CT/kg body weight) or a control (Milli-Q™ water) were fed to infected beetles and all the procedures were done as described above, and was repeated three times.
2.8. Data analysis

All statistical analyses were performed using SAS® version 9.3 (SAS institute Inc, Cary, North Carolina). Data from the in vitro study (except the CT depletion assay) fulfilled all three assumptions of ANOVA. So, the proportions of cysticercoid excystation in treatment groups were analyzed using PROC GLM fitting repetitions of experiments as a random variable. When an overall significant effect was seen, pairwise comparisons were done using a post-hoc Tukey test. The data from the CT depletion assay were analyzed by using a non-parametric Wilcoxon rank sum test. In the in vivo study, the numbers of cysticercoids established in the treatment groups were analyzed using PROC GENMOD, fitting negative binomial distributions. When overall differences were observed individual comparisons were done using least square means.

3. Results

3.1. In vitro study with praziquantel and pine bark extract

All concentrations of praziquantel and PB significantly reduced the mean percentage of cysticercoid excystation compared to their respective controls (Figs. 1A & B) and a significant concentration dependent effect was observed for both treatments with the highest concentrations having the strongest inhibitory effect on excystation ($F_{2,4} = 26.87$, $p = 0.0003$ for praziquantel and $F_{2,4} = 25.57$, $p = 0.0003$ for PB). The cysticercoid excystation inhibitory effect was the same for the praziquantel and PB treatments ($F_{1,4} = 0.49$, $p = 0.4887$).
3.2. *In vitro* study with pine bark, hazelnut and white clover extracts

The mean percentages of excystation of cysticercoids after exposure to the three CT were significantly different among the treatment groups ($F_{3,4} = 28.37$, $p < 0.0001$) and concentrations ($F_{2,4} = 110.58$, $p < 0.0001$; Fig. 2) and there was an interaction between treatment groups and concentrations ($F_{4,4} = 5.48$, $p = 0.0015$). Concentration was found to be a significant parameter for all three CT with the highest concentrations having the strongest negative effect on excystation ($F_{2,4} = 66.48$, $p < 0.0001$ for PB, $F_{2,4} = 9.97$, $p = 0.0067$ for HN and WC).

The mean percentage of cysticercoid excystation with all three CT depleted solutions showed significantly more cysticercoid excystation compared to their respective controls ($\chi^2 = 4.09$, df = 1, $p = 0.043$ for PB and WC, $\chi^2 = 3.97$, df = 1, $p = 0.043$ for HN; Fig. 3).

3.3. *In vivo* study with praziquantel, pine bark and hazelnut extracts

There was a significant effect of treatment (praziquantel and PB) on cysticercoid establishment ($\chi^2 = 133.1$, df=3, $p<0.0001$) but the effect was not dose dependent ($\chi^2 = 1.92$, df=2, $p=0.382$). All three doses of praziquantel reduced cysticercoid establishment to almost zero while each beetle in the control treatment had $+SE$ 27.1 ± 6.65 cysticercoids (Fig. 4A), whereas the effect of CT from PB at all doses did not differ from the control, although there was a trend of reduced establishment ($p=0.841$, 0.374 and 0.098 for low, medium and high doses respectively; Fig. 4B).

The separate experiment with HN at 500 mg CT/kg beetle body weight revealed significantly lower cysticercoid establishment (mean cysticercoids per beetle $+SE$: 25.5
± 2.54) in comparison to the control group (mean cysticercoids per beetle ±SE: 36.3 ± 2.33), \( \chi^2 = 10.48, \text{df} = 1, p = 0.0012 \).

4. Discussion

The results from the *in vitro* study suggest that treatment with CT from three different plants (PB, HN and WC) and praziquantel can substantially reduce the excystation of *H. diminuta* cysticercoids in a concentration-dependent manner. A similar concentration-dependent inhibitory effect of praziquantel has been previously shown using the same model (Woolsey, 2012). Our *in vivo* results showed that PB did not significantly inhibit cysticercoid establishment, but praziquantel and HN were associated with a reduction in the number of established cysticercoids. The cysticercoid excystation inhibitory effect of CT from the three plant extracts disappeared in the presence of tannin-inhibitor polyvinylpolypyrrolidone (PVPP) (Hagerman and Butler, 1981), confirming that CT are the major active compounds for the observed inhibition.

The observed effects of CT in this model are most likely due to their direct antiparasitic activity, although additional, indirect effects by increasing host resistance may occur in mammals (Hoste et al., 2006).

The cysticercoid capsule, scolex and other cellular structures contain protein with polysaccharides and lipids (Burt, 1980). As CT are able to bind to proteins (Hoste et al., 2006), they might interact with the protein portion of the cystic capsule and alter its physical and chemical properties, as reported previously for *Trichostrongylus colubriformis*, where direct damage to the cuticle was observed after incubation with CT (Hoste et al., 2006). Condensed tannins might also interfere with enzyme activities,
which are involved in metabolic pathways responsible for the development and functioning of parasites (Athanasiadou et al., 2001). Furthermore, due to the presence of pores and vesicles in the cystic capsule (Burt, 1980), bioactive compounds may reach the internal structures of the scolex and other cellular proteins. Taken together, all of these changes might interfere with the cysticercoid structures and metabolic pathways, which are essential in cysticercoid functioning and excystation. However, the exact mode of action of CT and the active compounds responsible for the anthelmintic activity are still unknown (Novobilský et al., 2013) and could differ depending on the species of parasite, its developmental stage, and possibly the biochemical characters and structures of the forage species (Min and Hart, 2003).

The different potency of CT from these plant extracts may be associated with the percentage of procyanidin monomer units or the mDP of the CT polymer, as the CT content in all three applied samples was standardized in the assays. However, HN also contains a small percentage of galloylated CTs (Irene Mueller-Harvey and Christos Fryganas, unpublished results), which may also influence anti-parasitic activity (Brunet and Hoste, 2006). The preliminary conclusion from this CT series is that the procyanidin tannins were more effective than prodelpphinidin tannins in the in vitro and in vivo experiments. This is an unexpected finding as most other studies ascribed higher anthelmintic activities to the prodelpphinidins (Brunet and Hoste, 2006). There may be two possible reasons: 1) the presence of galloylated CT in HN or 2) the fact that the pH values of insect guts tend to be alkaline (Gullan and Cranston, 2010) and are therefore quite different from the rumen or abomasum of ruminants. Relatively little is known about the reactivity of PC and PD tannins under alkaline conditions with constituents of
the insect gut. Anthelminthic activities of CT from different plants or plant extracts are known to have markedly different effects on parasites. For example, grazing of sheep on *Lotus pendunculatus* reduced nematode fecal egg counts more effectively than grazing on *L. corniculatus* (Niezen et al., 1998), as *L. pedunculatus* has a higher PD/PC ratio than *L. corniculatus* (Foo et al., 1997). Thus, the multitude of different CT structures (Mueller-Harvey and McAllan, 1992) may influence their biological activities (Athanasiadou et al., 2001). Further experiments are needed to determine the relative contributions of mDP and PD/PC ratio on cysticercoid excystation.

Availability of free CT in the intestine may be important factor for CT to be effective. Formation and dissociation of the protein-CT complex is highly pH dependent. Stable protein-CT complexes are formed at pH 5 - 7, but the complexes easily dissociate and release proteins at higher and lower pH (Mueller-Harvey and McAllan, 1992). Optimum complex formation occurs at the isoelectric point of the protein, but little is known about the isoelectric point of proteins in the beetle gut. There is thus a possibility of lack of formation of protein-CT complex in the intestine of beetles or CT being subjected to oxidative changes.

In summary, our *in vitro* results indicate concentration dependent inhibitory effect of all tested CT in plant extracts on cysticercoid excystation, the HN extract being most potent, followed by PB and WC. Anti-excystation activity appeared to be positively linked to the presence of procyanidin tannins. The *in vivo* treatment with HN reduced cysticercoid establishment, and is the first observation on anti-cestodal properties of CT from plant extracts. Although the invertebrate-parasite model is not fully representative of the biological action of CT in the mammalian system, this model could be useful for
a first screening of potentially interesting compounds. This invertebrate model has several advantages over vertebrate models as far as, ethical clearance, legislation, time and cost are concerned. Future studies will need to address the mechanism of CT action and include comparative studies with vertebrate animals in order to explore their effects against the different lifecycle stages and species of tapeworms.

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Fig. 1. *In vitro* mean percentage (n=5, ±SEM) of cysticercoids excystation treated with praziquantel (A) and condensed tannin (CT) extract from pine bark (B) at different concentrations. Ten cysticercoids were used for each concentration. Control refers to 2% DMSO for (A) and to Milli-Q™ water for (B). Different letters within each figure represent statistical significance (α = 0.05).

Fig. 2. *In vitro* mean percentage (n=5, ±SEM) of cysticercoids excystation treated with three different condensed tannin (CT) extracts [white clover flower (WC), pine bark (PB), and hazelnut skin (HN)] at different concentrations. Ten cysticercoids were used for each concentration. Control refers to Milli-Q™ water. Different letters within each figure represent statistical significance (α = 0.05).

Fig. 3. *In vitro* mean percentage (n=3, ±SEM) of cysticercoid excystation treated with three different condensed tannin (CT) depleted solutions (depleted) and condensed tannin extract solutions (control) at 2.5mg CT/ml concentration. Ten cysticercoids were used for each solution of white clover flower (WC), pine bark (PB) and hazelnut pericarp (HN). *refers to a significant difference relative to the respective controls.

Fig. 4. *In vivo* mean number of cysticercoids (n=5 for A and n=3 for B, ±SEM) treated with praziquantel (A), and a condensed tannin (CT) extract from pine tree bark (B) at different concentrations. Average numbers of cysticercoids from four beetles were used for each treatment. Control refers to 2% DMSO in (A) and to Milli-Q™ water in (B).
Figure 1A

Mean % of excystation (+SEM)

Concentration (mg praziquantel/ml)

Control  0.0001  0.0010  0.0100
Figure 4A

Mean number of cysticercoids (+SEM)

Control  25   50   100
Concentration (mg praziquantel/kg of beetle body weight)
Figure 4B

B.

Mean number of cysticercoids (+SEM)

Concentration (mg CT/kg of beetle body weight)

Control 125 250 500