Echinococcus multilocularis infection affects risk-taking behaviours in Microtus arvalis: adaptive manipulation?

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Abstract

Manipulation of host behaviour by parasites to enhance transmission to the next host is a fascinating phenomenon that has interested scientists since the 1970s. It has been proposed that infection with the cestode *Echinococcus multilocularis* produces an impairment of the antipredatory behaviour in the rodent intermediate host common vole, *Microtus arvalis*, which may facilitate transmission of the tapeworm to the canid final host. In this study, we observed the behaviour of infected common voles at 12 weeks post-infection, when protoscoleces production and maturation commonly occurs, in order to assess behavioural changes compared to uninfected controls, that might ease predation in the wild. Infected and uninfected voles were monitored for 24 hours to observe their spontaneous activity. In addition, the next day, both infected and uninfected voles were subjected to four different behavioural tests: open field test, barrier test, platform test and air-puff test in a running wheel. No significant difference between uninfected and infected voles emerged during the behavioural tests. However, observation of spontaneous activity revealed that infected voles increased their feeding frequency and spent significantly more time above bedding even when not eating, compared to the uninfected controls. In the wild, these behavioural changes increase the animals exposure to predators, raising their chance of becoming prey. These findings are the first direct evidence consistent with behavioural manipulation by *E. multilocularis* on common voles.

**Keywords:** behavioural alteration, *Echinococcus multilocularis*, *Microtus arvalis*, predator-prey relationship, trophically-transmitted parasites
Introduction

Understanding the transmission dynamics of zoonotic parasites is essential to design efficient public health control strategies. The relationship between a parasite and its host(s) has been the subject of many investigations. For some parasites completion of the life cycle occurs within a single host (simple life-cycle parasites; SLCs), whereas others need different host species (complex life-cycle parasites; CLCs) (Brown et al., 2001). For the latter, one or more hosts acts as Intermediate Host (IH), in which the parasite reproduces asexually, and one host becomes the Definitive Host (DH), in which the parasite reaches sexual maturity (Brown et al., 2001). For a CLC parasite, the most common form of host-to-host transmission is trophic, in which the prey plays the role of IH and the predator DH (Bethel and Holmes, 1973). In the 20th century, the so-called ‘manipulation hypothesis’ was proposed to explain the evolution of the parasite-host relationship in some CLC parasites. This hypothesis predicts that the ability of a parasite to induce phenotypic alteration in the IH produces increased parasite transmission rates to the DH final host (Dobson, 1988; Freedman, 1990; Moore, 2002). The ‘manipulation hypothesis’ is supported by several studies showing an advantage for the parasite presenting an infection-induced phenotypic alteration (e.g. behaviour, morphology and/or physiology) in the IH (Bethel and Holmes, 1973; Lafferty and Morris, 1996; Lefèvre et al., 2009).

Echinococcus multilocularis (Leuckart, 1863), the third most relevant human food-borne parasite in the world (FAO/WHO, 2014), is a cestode presenting a CLC with IH-DH transmission through predator-prey relationship and could represent a model to study parasite transmission strategies. E. multilocularis is a gastrointestinal parasite whose adult develops in the small intestine of wild and domestic canids (e.g. coyote Canis latrans, red fox Vulpes vulpes, raccoon dog Nyctereutes procyonoides, golden jackal (Canis aureus), wolf Canis lupus, domestic dog Canis lupus familiaris) and, to a lesser extent, in cats (Felis catus) (Romig et al., 2017). In the intestinal lumen of the DH, the adult worm produces fertilized eggs which are
shed in the environment with the host faeces (Romig et al., 2017). If accidentally ingested by a competent IH (mostly rodent), the eggs hatch in the host's gastrointestinal tract, releasing an oncosphere which migrates to the target organs, most frequently the liver, through the bloodstream of the portal vein (Eckert et al., 2011). Once in the liver, the oncosphere develops into the metacestode (Heath and Lawrence, 1976) which starts asexual multiplication, causing multilocular cyst-like lesions (Kamiya, 2008) and produces the infectious larval stage: the protoscolex. The cycle is completed only when a competent predator consumes an infectious prey (i.e., with fertile lesions containing mature protoscoleces).

Although it has been hypothesized that E. multilocularis infected rodents might have a diminished ability to escape predation, so far there are no data supporting such a hypothesis (Vervaeke et al., 2006). Therefore, clarifying the effects of E. multilocularis infection on IH anti-predatory behaviour may be a further key feature to consider in the epidemiology and transmission studies of this parasite. The overall aim of this study was to preliminarily investigate possible behavioural alterations caused by E. multilocularis in the common vole Microtus arvalis, a key host species for the maintenance of the parasitic cycle in Europe (see Oksanen et al., 2016 and references therein). With this pilot study, we aimed to obtain the proof of concept to support the hypothesis of parasitic manipulation of IH behaviour in E. multilocularis so as to facilitate its transmission. Specifically, that if any behavioural alteration is detected, this should be in the direction of an increased risk to predation of infected animals compared to controls; this in the natural cycle would translate into increased transmission rate. Our working hypotheses were: i) the parasite induces alteration of specific behavioural patterns of the IHs, facilitating its predation (behavioural alteration); ii) this alteration occurs when the IH is infective (i.e. with viable protostrongylid larvae).
Materials and methods

Experimental animals

The experimental groups included 16 *M. arvalis* aged between 9 and 14 weeks, bred and kept in captivity at the Unit of Parasitology of the University of Hohenheim, Stuttgart, Germany. The animals were randomly assigned to one of two experimental groups: one (treated, T) consisted of 10 animals (5 males and 5 females) infected with 500 viable embryonated eggs of *E. multilocularis*, whereas the other (control, C) was made of 6 sham-treated animals (3 males and 3 females). One infected and one control animal died or had to be euthanized by CO2, according to legal requirements, before the experiment at 12 weeks p.i. Therefore, the experiment was done with 14 animals: 9 infected (4 male, 5 female) and 5 control animals (2 male, 3 female). Animals were individually housed, under standard conditions of 12 hours of daylight and 12 hours of darkness (12-12 DL); Light (L): 7 a.m.-7 p.m.; 22°C; humidity 40-60%, in standard polycarbonate cages (Type III: 42.0 x 26.5 x 15.5 cm). Cages were provided with bedding (wood shavings and hay) that was changed once a week. Water and food pellets (pellet food for rats and mice; Altromin®, Germany, Lage) were available *ad libitum*.

Experimental infection

*E. multilocularis* eggs were obtained from adults collected from the intestines of naturally infected foxes hunted in Switzerland. Before inoculation, the viability of the eggs was tested using the ‘Sodium Hypochlorite Resistance’ test (*Joekel and Deplazes, 2017*), resulting in a viability of 12.5%. Each treated vole was infected with about 500 eggs by oral gavage, using a metal feeding needle, with the viable eggs suspended in 0.5 ml of PBS. The control animals were sham-inoculated with 0.5 ml of distilled water. After the infection, the voles were housed under BIO III security level for 3 days due to the risk of regurgitation of parasite eggs. After this period, the animals were brought back to the Animal Facility and monitored daily for post-
inocula trauma or illness.

**Behavioural observation and tests**

All behavioural observation and tests were performed at 12 weeks post infection (w.p.i.), to allow for protoscolex (infectious larva stage) production and maturation (Woolsey *et al.*, 2015). Initially, we planned to carry out behavioural observations also at 18 w.p.i., but due to progressing disease several animals had to be euthanized and the remaining group size did not allow for adequate replication.

Prior to the experimental phase, the voles were observed for 24 hours to assess spontaneous behaviour in the cage. Afterwards, the animals were subjected to four behavioural tests: open field, barrier test, platform test and air puff test in a running wheel (Fig. 1). All tests targeted behaviours potentially associated with anti-predatory strategies. Both open field and barrier tests were conducted to observe exploratory and risk-taking behaviours. The platform test aimed at assessing an anti-predatory response. The observation in a running wheel system aimed at assessing the physical performance, whereas the air puff test, conducted when the animal was stationary in the running wheel, aimed to assess the vole reaction time to a physical stimulus. Many behavioural variables were chosen as recommended by a previous study of Herde and Eccard (2013). All behavioural tests were carried out in succession, in a different room from the one in which the animals had been kept till then, but under the same conditions of (12-12 DL, L: 7 a.m.-7 p.m.; 22°C and 40-60% humidity). The animals were allowed to rest in a cage for two hours between each trial. All tests were video-recorded, and operators left the test room during the trials.

**Behavioural observation (24-hr monitoring)**

The spontaneous behaviour of treated and control voles was recorded for 24 hours at 12 w.p.i.
starting from 6 a.m. To increase the view of the animal in the home cage during the observations, the hay, but not the bedding, was removed from each cage three days before the start of monitoring, to accustom the animal to its absence. The 24-hour observation was carried out using one night-vision surveillance video camera (Lorex, Flir 8-Channel 1080p DVR) per cage (Fig. 1A). The setup allowed monitoring a maximum of 8 home cages simultaneously, so two sessions were needed, one day apart. The monitoring was carried out under controlled conditions (12-12 DL, L: 7 a.m.-7 p.m.; 22 °C, and 40-60% humidity). No operators entered the room during the 24 hours of monitoring. The time spent above the bedding and the frequency and duration of eating and drinking events were recorded (see Table 1).

Open Field Test
According to Herde and Eccard (2013), the open field test apparatus was built as an arena consisting of a round aluminium box (100 cm diameter) with an open top. The grey wall of the arena was 35 cm high, preventing any animals from escaping. A circle of 80 cm in diameter was drawn around the centre of the arena on the white floor of the arena so as to allow the identification of two areas: a safe peripheral zone near the outer wall (a 20 cm wide ring) and an unsafe zone (the centre of the arena; Fig. 1B). Areas far from walls are known to be perceived as “unsafe” for small mammals (Prior and Sachser, 1995). Each animal was transported in the test room in a white polycarbonate plastic tube (5.08 x 10.16 cm) and placed in the center of the arena. The test lasted 10 minutes and started when the animal reached the arena wall for the first time. The time spent by the vole in each of the two areas was recorded (see Table 1).

Barrier Test
The barrier test consisted of a semi-transparent plastic box (36 x 26 x 21 cm) with sterilized

https://doi.org/10.1017/S0031182024000507 Published online by Cambridge University Press
sand as bedding. The plastic box was divided into two equal compartments by a white, 4.5-cm high wood barrier (Fig. 1C). Each animal was carried to the test room in a white polycarbonate plastic tube (5.08 x 10.16 cm). The animal was randomly placed in one of the two compartments and the latency to cross the barrier the first time was measured. The duration of the test was 5 min. The other measured variables are reported in Table 1.

**Platform Test**

The platform test was conducted in a Type IV cage Type (59 x 380 x 200 cm) with a small white platform (an overturned box of 30 x 17.5 x 6 cm) put in the centre of the home-cage filled with 3 cm of bedding layer (Fig. 1D). Each animal was transported in the test room in a white polycarbonate plastic tube (5.08 x 10.16 cm). The test started when the animal was placed in the centre of the platform and the latency to hide in the bedding was measured. The test ended after 3 minutes.

**Running wheel and Air Puff Test**

The apparatus used in this test consisted of a voluntary running wheel (11.5 cm diameter). The vole was forced to stay in the running wheel by three wooden walls (one large wall measuring 17 x 14.5 cm, and two small walls measuring 11 x 14.5 cm) covered with black cardboard and one transparent plastic wall allowing video-monitoring during the test. The air puff was delivered through two holes (3 mm diameter) in the largest wooden wall located at 4.5 cm from the bottom of the wheel, so that the air puff stimulus (delivered from compressed air bottle, Air Duster 125 ml, Vivanco®, Germany, Ahrensburg) reached the right side of the vole (Fig. 1E). The number of rotations was recorded by a Liquid Crystal Display (LCD) activity counter (STARR®, life science group; United States, Oakmont) connected to the running wheel. In the described apparatus, the animal was forced to stay inside the wheel, but unlike the standard
forced running system, it was free to stay still or move. Each animal was transported in the test room in a white polycarbonate plastic tube (5.08 x 10.16 cm). The test started when the animal was placed in the running wheel. The air puff was delivered when the vole was stationary. The animal received an air-puff stimulus of 1 second after at least 5 minutes on the running wheel. The vole was video-monitored from the beginning of the test till 1 minute after the stimulus application. The spontaneous activity in the running wheel before the stimulus application and the freezing time after the air puff were recorded. The variables measured are listed in Table 1.

Post-mortem analyses

All the experimental animals were sacrificed after 18 weeks by CO2, according to legal requirements, and subjected to a post-mortem visual inspection of the lesions. For estimating the number of mature protoscoleces in each subject the portion of the liver presenting lesions was cut off and homogenised using a pestle in 1 ml of distilled water. The homogenate was filtered through a nylon mesh of 1 mm pore size (the filter was washed with 1 ml of distilled water) onto a squared petri dish (12 x 12 cm) with a 5 x 5 mm squared grid on the bottom for a total of 576 cells. The protoscoleces inside the cells of the grid laying on both diagonals of the petri dish (48 cells) were manually counted under a microscope. An estimate of the number of protoscoleces was then computed by multiplying the average number of protoscoleces calculated for the inspected cells by the total number of cells.

Behaviour quantitative analysis

The videos were exported in .asf format and imported using Boris version 6.3.5 software (Friard and Gamba, 2016). Each video was checked at normal speed, and the total duration in seconds (state behaviour) and the occurrence of an event(s) of each variable (event behaviour) was considered. Software R 4.2.2 (R Core Team, 2022) was used for all data analyses. Each
variable reported in Table 1 was compared between infected and control groups using a two sample bootstrap t-test (Efron and Tibshirani, 1994) by means of the boot.t.test function in the package MKinfer (Kohl, 2019), setting the number of bootstrap samples = 10,000. This resampling approach was chosen as it is more robust and powerful with small and unbalanced samples, as well as for hypothesis testing for groups with different patterns of non-normality but equal variance (Dwivedi et al., 2017). Sex was excluded from the analysis because we did not expect a differential effect of infection on different sexes (Woolsey et al., 2015). Furthermore, a Fisher exact test was performed to compare the frequency of infected and uninfected animals that showed spontaneous activity in the wheel, before the application of the air puff stimulus. The same test was also applied to compare the frequency of infected and uninfected vole behavioural reaction within 3 seconds after the air puff stimulus application. We used an alpha level of 0.05 for all statistical tests.

Results

Behavioural observation (24-hr monitoring)

During the 24-hour in-cage monitoring period, infected voles spent, on average, 5.6 hours more above bedding compared to intact voles (32.4% increase, T-voles = 22.93 ± 0.87 vs. C-voles = 17.29 ± 2.61, p=0.03; Table 2, Fig. 2B). Furthermore, infected voles showed a mean increase of 62.5 eating events per 24 hours compared to control animals (41.37% increase, T-voles = 151.11 ± 15.52 vs. C-voles = 88.60 ± 5.05; p=0.002; Table 2, Fig. 2A).

Behavioural tests

No significant differences between control and treated voles were found for any of the behavioural tests (open field, platform, barrier, running wheel) (Table 2). Moreover, in spontaneous running activity in the wheel, infected and uninfected voles did not show any
significant difference (Fisher exact test, p>0.05). In fact, all control voles displayed spontaneous running activity in the running wheel before the air puff test, as did 8 out of 9 infected voles. As regards the voles that engaged in some spontaneous running, we did not detect any significant difference between control and treated voles in the time they spent doing this activity and the running events (Tab. 2). The most common reaction to the air puff observed within 3 seconds from the stimulus was a freezing reaction (3 control and 7 infected voles froze after the stimulus; Fisher exact test, p>0.05). In addition, the time spent freezing in the 60 seconds after stimulus delivery did not differ in the two experimental groups (Tab. 2).

Post-mortem analyses

Seven of the infected voles were euthanized and necropsied at 18 weeks p.i., two had to be euthanized and necropsied earlier due to disease progress in line with animals welfare requirements (13 weeks p.i.). All infected voles had *E. multilocularis* lesions in the liver and mature protoscoleces. One animal necropsied at 13 weeks p.i. presented 5 discrete lesions with 44,904 protoscoleces, one necropsied at 13 weeks p.i. had three discrete lesions with 10,836 protoscoleces, while animals necropsied at 18 weeks p.i. presented an average ± SE of 2.7 ± 0.9 lesions each with a mean ± SE of 113,773 ± 59,431 (min = 1,342, max = 452,687) mature protoscoleces, N = 7. Lesions were present in all the liver lobes, with 9 lesions found in the Left Lateral Lobe, 4 in the Left Medial Lobe, 6 in the Right Medial, 7 in the Right Lateral Lobes and 1 in the Caudate Lobe. All control voles were euthanized 15-18 weeks p.i., no lesions and protoscoleces were present in any of these animals.

Discussion

Our study aimed to assess whether and which behavioural alterations might occur following *E. multilocularis* infections in one of its IHs, *M. arvalis*. Specifically, we compared the behaviour
of voles experimentally infected with *E. multilocularis* eggs to the behaviour of uninfected ones at 12 weeks post-infection.

Both infected and uninfected voles displayed comparable behaviours when tested in arenas, experimental cages, or running wheels. However, despite the small sample size, a clear difference emerged between the two groups in spontaneous behaviours in their home cage. In fact, we observed a significant increase in the time spent above bedding by infected animals compared to intact voles (Fig. 2B). This noteworthy result highlights that the condition of being infected may have affected the animals’ hiding behaviour, which is their spontaneous anti-predator response (*Sundell and Ylönen, 2004*), resulting in a significant increase in time spent in the open space of the cage. Moreover, the infected voles, when above bedding, fed more frequently compared to the controls (Fig. 2A). In the wild this behavioural alteration would translate to longer time spent in the open, outside the nest, and more frequent feeding with an increased likelihood of predation by a DH. This is consistent with the hypothesis that the parasite can induce behavioural modification, making the IH more susceptible to predation. This conclusion is in line with previous studies reporting that cestodes alter IH’s activity by decreasing or increasing the host activity level depending on the species involved (*Lafferty and Shaw, 2013*). For instance, *Echinococcus granulosus* (sensu lato) infection may cause a moose physical impairment due to lung lesions induced by the parasite larval stage (*Messier et al., 1989*). This case can be explained by the side effect of being infected (*Adamo, 2013; Dantzer et al., 2008*) rather than a manipulation of the parasite on the IH. Differently, fishes infected by *Schistocephalus solidus* larval stages display increased surfacing activity, consequently increasing the probability of being detected by a predator (*Øverli et al., 2001*). In this case the effect of the parasite can be more plausibly explained by a specific manipulation aimed at increasing the transmission probability. This seems to be the same effect we observed in our voles infected by *E. multilocularis*. As a matter of fact, spontaneous activity in the running
wheel (before the air puff test) did not differ between infected and uninfected voles, suggesting that the parasite did not seem to cause (at least up to 12 weeks from infection onset) a general debilitation of the voles. Furthermore, the freezing reaction, an anti-predatory strategy displayed by rodents, is triggered by traumatic or stressful experiences (Roelofs, 2017). Both control and treated animals comparably showed freezing reaction to the air puff stimulus, suggesting that the parasite infection did not significantly alter fear elicited behavioural responses.

Although we did not measure the amount of food eaten by intact and infected animals when above bedding, it would not be surprising if the more frequent feeding of the infected voles resulted in greater food consumption. However, the observed increase in frequency of feeding might be consistent with one or both of two different mechanisms linked to an increase in energy demand in infected animals: the parasite’s direct draw of energy from the host or the host’s immune response to the infection (Schmid-Hempel, 2005). Referring to the former mechanism, an alteration of the insulin pathway and, possibly, of the glucose levels might induce a nutritional needs impairment and an alteration of food intake requirements. It is known that *E. multilocularis* has receptors for insulin to target the liver as an optimal environment for the settlement and establishment of metacestodes (Hemer et al., 2014). However, it is unknown whether the parasite might have nutritional dependence on host glucose, as is the case of the trematode *Schistosoma mansoni* (Ahier et al., 2008) or what the role of insulin receptors in modulating the glucose parasite level could be (Brehm et al., 2006). In addition, the time spent above bedding might be altered via physiological systems other than the digestive system. For instance, the parasite might increase the host oxygenation demand, as studied in swines affected by pulmonary metastrongylosis, a parasitosis caused by nematode *Metastrongylus* species (Pavlović et al., 2021). Another example of pulmonary debilitation which involves the helminth-amphibian host system, regards infected toads (*Bufo bufo*) which might have

https://doi.org/10.1017/S0031182024000507 Published online by Cambridge University Press
decreased lung function and, consequently, an altered oxygenation demand. (Goater and Ward, 1992). In our case, the altered oxygen demand can directly induce the vole to stay above bedding almost all the time. In this case, the increased feeding frequency might be caused simply by the continuous availability of food close to the animal. Another possibility could be that liver tissue damage (in this case induced by larvae proliferation) might cause a host blood intoxication (Lockwood et al., 1979). Toxicants in the blood, such as ammonia, might have a neurotoxic effect, impairing the activity of the IH, its reactivity to stimuli and mobility, thus making it more susceptible to predation (Lockwood, 2004; Ong et al., 2003). Our findings are not consistent with this hypothesis as the infected voles do not seem to be less mobile nor less reactive than control voles.

Despite the small sample size and the unclear physiological mechanisms underlying the increased above-bedding activity in *E. multilocularis*-infected IHs, the pilot study provided the first direct evidence of IH behavioural alteration following *E. multilocularis* infection. In particular, the parasite infection increased both the time spent above bedding as well as the time the voles spent feeding compared to the intact controls, suggesting a specific effect of the parasite on the activity of the IH. The study is intended as a starting point for investigating the effect of *E. multilocularis* on the physiology of rodent IHs.

**Data availability.** All data and code used for the analyses are available at GitHub repository link: https://github.com/MatildeUNIBO/ParManPilot.git

**Acknowledgements.** We thank Dr. Jens Jacob of the Julius Kühn-Institut for valuable suggestions for the study protocol and technical support. We thank Prof. Dr. Jana Anja Eccard of the University of Potsdam for help in the lab for behavioural apparatus design. We thank Giulia Ferrucci for help in the lab. We thank the reviewers for helping with useful comments.
that improved the manuscript.

**Author contributions.** MM conceived and designed the study, conducted data gathering, performed statistical analyses and wrote the article, TC conducted data gathering and wrote the article, TR conceived and designed the study, reviewed and edited the article and supervised the project, AG designed the study, reviewed and edited the article, DG designed the study, performed statistical analyses, reviewed and edited the article, MZ reviewed and edited the article, AM conceived and designed the study, reviewed and edited the article and supervised and coordinated the project, acquired funding.

**Financial support.** This work was partially funded by PhD grant from the Natural Sciences & Engineering Research Council of Canada (NSERC300), project number: 10013904, project title: ‘‘The ecology of parasites with complex lifecycles in multi-host systems: *Echinococcus multilocularis* transmission dynamics at multiple spatial scales’’.

**Competing interests.** The authors declare there are no conflicts of interest.

**Ethical standards.** Our study was done with animals that had been infected with *E. multilocularis* for other purposes (in vivo propagation of *Echinococcus*, permit no. A391/14 PA, Regierungspräsidium Stuttgart, Germany). The behavioural tests were approved by the animal welfare commission of the Regierungspräsidium Stuttgart, Germany on 10.12.2018, permit no. V350/18 PA.
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Figure 1. Schematic drawing of the behavioral observation and tests conducted on 9-14 weeks old males and females *Microtus arvalis* to test behavioral manipulation by *Echinococcus multilocularis* infection. Structure of A) Behavioural observation (24-hr monitoring) apparatus; B) Open Field test; C) Barrier test; D) Platform test; D) Running wheel and Air Puff test (APH, air puff holes). Treated group was subjected to oral injection of 500 *E. multilocularis* eggs (estimated viability of 12.5%), whereas control group was sham-inoculated with distilled water. Figure was created with BioRender.com.
Figure 2. Results of 24-hr behavioral observation conducted on 9-14 weeks old males and females *Microtus arvalis* to test behavioural manipulation by *Echinococcus multilocularis* infection. Boxplots of: A) Eating events; B) Time spent above bedding within 24 hours (duration). Treated group (white colour) was injected with 500 *E. multilocularis* eggs (estimated viability of 12.5%), whereas control group (grey colour) was treated only with distilled water.
**Table 1.** List of tested behavioural variables for the in-cage monitoring and behavioural trials performed in the study conducted on 9-14 weeks old males and females *M. arvalis* to test behavioural alteration by *E. multilocularis* infection. Treated group was injected with 500 *E. multilocularis* eggs (estimated viability of 12.5%), whereas control group was sham-inoculated with distilled water. The variables reflect mainly anti-predatory behaviours and activity of the tested animals.

<table>
<thead>
<tr>
<th>Test</th>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Behavioural observation (24-hr monitoring)</td>
<td>Drinking</td>
<td>Events and duration in minutes that the animal spends sucking the bottle of water</td>
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<tr>
<td></td>
<td>Eating</td>
<td>Events and duration in minutes that the animal spends chewing the pellet</td>
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<td></td>
<td>Above bedding</td>
<td>Total duration in hours that the animal spends above bedding</td>
</tr>
<tr>
<td>Open Field test</td>
<td>Peripheral time</td>
<td>Total duration in seconds that the animal spends in the peripheral (safe zone) of the arena</td>
</tr>
<tr>
<td></td>
<td>Centre time</td>
<td>Total duration in seconds that the animal spends in the centre (unsafe zone) of the arena</td>
</tr>
<tr>
<td>Barrier test</td>
<td>Freezing</td>
<td>Events and duration in seconds that the animal spends completely stationary in the two sides of the barrier for at least 1 second</td>
</tr>
<tr>
<td></td>
<td>Latency</td>
<td>Latency (in seconds) to jump over the barrier</td>
</tr>
<tr>
<td>Test</td>
<td>Outcome</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>----------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Platform test</td>
<td>Latency to hide</td>
<td>Latency (in seconds) to hide into the bedding the first time</td>
</tr>
<tr>
<td>Running wheel and Air Puff test</td>
<td>Spontaneous activity</td>
<td>Presence, events and duration in seconds that the animal spends running into the wheel before the stimulus application</td>
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<tr>
<td></td>
<td>Freezing after stimulus</td>
<td>Duration in seconds of freezing reaction within 1 minute after the stimulus application</td>
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<td></td>
<td>Reaction</td>
<td>Presence of reaction (running) within 3 seconds after the stimulus application</td>
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</table>
Table 2. Results of the bootstrap t-test calculated on the variables measured in the behavioural tests on 9-14 weeks old males and females *M. arvalis* aimed at testing behavioural manipulation by *E. multilocularis* infection. Treated group was subjected to oral injection of 500 *E. multilocularis* eggs (estimated viability of 12.5%), whereas control group was sham-inoculated with distilled water. The bootstrap p-value is based on 10,000 bootstrap replications. Significant p-values ($P < 0.05$) are in bold. Difference of means = control group mean – treated group mean.

<table>
<thead>
<tr>
<th>Test</th>
<th>Variables (measure unit)</th>
<th>Difference of means</th>
<th>95% CIs</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>24-hr monitoring</strong></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td></td>
<td>Drinking Events (frequency)</td>
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<td>Drinking Duration (minute)</td>
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<td>Above Bedding (hour)</td>
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<td>-10.8</td>
<td>-0.9</td>
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<tr>
<td><strong>Open Field test</strong></td>
<td>Peripheral time (second)</td>
<td>2.1</td>
<td>-16.3</td>
<td>19.4</td>
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<tr>
<td></td>
<td>Centre time (second)</td>
<td>-2.3</td>
<td>-19.3</td>
<td>16.4</td>
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<tr>
<td><strong>Barrier test</strong></td>
<td>Freezing Events (frequency)</td>
<td>3.5</td>
<td>-1.0</td>
<td>8.7</td>
</tr>
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<td>Freezing Duration (second)</td>
<td>6.5</td>
<td>-48.5</td>
<td>61.6</td>
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<td>Latency (second)</td>
<td>-41.5</td>
<td>-127.0</td>
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<td>Digging Events (frequency)</td>
<td>-0.7</td>
<td>-6.5</td>
<td>5.9</td>
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<td>Digging Duration (second)</td>
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<td>34.3</td>
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<tr>
<td><strong>Platform test</strong></td>
<td>Latency to hide (second)</td>
<td>-10.9</td>
<td>-44.5</td>
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<td><strong>Running wheel and Air Puff test</strong></td>
<td>Spontaneous activity Events (frequency)</td>
<td>0.04</td>
<td>-5.5</td>
<td>5.7</td>
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<td>Spontaneous activity Duration (second)</td>
<td>-25.1</td>
<td>-88.1</td>
<td>39.9</td>
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<td>Freezing after stimulus (second)</td>
<td>-8.2</td>
<td>-37.5</td>
<td>21.0</td>
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