Biological control of horse cyathostomin (Nematoda: Cyathostominae) using the nematophagous fungus *Duddingtonia flagrans* in tropical southeastern Brazil

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1 CNPq scholarship.

**A R T I C L E   I N F O**

Article history:
Received 18 April 2008  
Received in revised form 4 May 2009  
Accepted 5 May 2009

**Keywords:**  
Nematophagous fungus  
*Duddingtonia flagrans*  
Cyathostomin  
Horse  
Biological control

**A B S T R A C T**

The viability of a fungal formulation using the nematode-trapping fungus *Duddingtonia flagrans* was assessed for the biological control of horse cyathostomin. Two groups (fungus-treated and control without fungus treatment), consisting of eight crossbred mares (3–18 years of age) were fed on *Cynodon* sp. pasture naturally infected with equine cyathostome larvae. Each animal of the treated group received oral doses of sodium alginate mycelial pellets (1 g/(10 kg live weight week)), during 6 months. Significant reduction ($p < 0.01$) in the number of eggs per gram of feces and coprocultures was found for animals of the fungus-treated group compared with the control group. There was difference ($p < 0.01$) of 78.5% reduction in herbage samples collected up to (0–20 cm) between the fungus-treated group and the control group, during the experimental period (May–October). Difference of 82.5% ($p < 0.01$) was found between the fungus-treated group and the control group in the sampling distance (20–40 cm) from fecal pats. During the last 3 months of the experimental period (August, September and October), fungus-treated mares had significant weight gain ($p < 0.01$) compared with the control group, an increment of 38 kg. The treatment with sodium alginate pellets containing the nematode-trapping fungus *D. flagrans* reduced cyathostomin in tropical southeastern Brazil and could be an effective tool for biological control of this parasitic nematode in horses.

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1. Introduction

A large variety of helminths are known to parasite horses. Nematodes, mainly cyathostomin species, are the most common and important among them. Also known as small strongyles, cyathostomin infections are responsible for causing anemia, weight loss, intestinal colic, and death in horses (Assis and Araújo, 2003). They are the most prevalent parasites in horses, present throughout the year in the pasture, with a wide distribution in different age groups (Barbosa et al., 2001; Quinelato et al., 2008).

Klei and Chapman (1999) reported field data suggesting that horses can acquire resistance to helminths with age, which is confirmed by the reduced parasite load and egg count in feces. This response is slow and inconsistent in most animals and unrelated to the intensity of previous contact with parasite. Kaplan (2002) and Matthews et al. (2004) discussed that worm control in horses is usually carried out with
anthelmintic drugs, which have not been totally effective for the control of these nematodes since their action is restricted to adult parasites and there is occurrence of resistance.

The continued use of the same anthelmintic class, as well as the rapid rotation of compound groups, introduction of resistant worms and the use of doses lower than the recommendation should be avoided (Mota et al., 2003). Biological control using natural nematode antagonistic fungi is among the most viable alternatives. These organisms comprise different types of fungi classified into predators, endoparasites and opportunists, whose action is concentrated in the fecal environment and directed against free-living parasitic larvae. Within the predator group, the species Duddingtonia flagrans stands out as the most promising for the control of gastrointestinal nematodiasis in domestic animals (Terrill et al., 2004; Dias et al., 2007a). However, to be used as a biological control agent, nematophagous fungi must have ability for nematode capture and survive passage through gastrointestinal tract (Waller et al., 1994).

Sodium alginate-based formulations containing D. flagrans mycelial mass have been experimentally evaluated against parasitic nematodes of animals in laboratory and field conditions (Ararjo and Sampaio, 2000; Ararjo et al., 2000; Dias et al., 2007b), but none these formulations have been developed for the control of parasitic nematodes of horses in the field.

The objective of the present study was to test an alginate pellet formulation containing D. flagrans for the biological control of cyathostomin in horses raised in fields.

2. Materials and methods

2.1. Fungal cultures

Isolate (AC001) of D. flagrans, a nematode-trapping fungus belonging to the genus Duddingtonia, was kept in test tubes at 4 °C containing 2% corn meal-agar (2% CMA) in the dark. The isolated was obtained from a Brazilian soil using the soil sprinkling method (Duddington, 1955), modified by Santos et al. (1991).

Fungal mycelia were obtained by transferring culture disks (approximately 5 mm in diameter) of fungal isolates in 2% CMA to 250 mL Erlenmeyer flasks with 150 mL liquid potato-dextrose medium (Difco), pH 6.5, and incubated under agitation (120 rpm), in the dark at 26 °C, for 10 days. Mycelia were then removed for pelletizing using sodium alginate as described by Walker and Connick (1983) and modified by Lackey et al. (1993).

2.2. In vivo experimental assay

The experiment was conducted at the horse experimental sector of the Federal University of Viçosa, Viçosa, MG, Brazil, latitude 20° 45’20”S, longitude 42° 52’40”W, from May to October 2007.

In the beginning of the experiment, the 3–18 year old crossbred mares were previously dewormed with 200 μg/kg live weight Ivermectin 1% and 6.6 mg/kg live weight Pyrantel Pamoate (Centurion Valné®), Montes Claros-Minas Gerais, Brazil.

Fourteen days after the anthelmintic treatment, the mares were separated into two groups (fungus-treated and control) of eight animals each on the basis of age and weight. Mean age and mean weight of the fungus-treated group were 6.3 (±6.1) and 386.2 (±54.07) respectively, and 7.1 (±4.7) and 381.1 (±53.91) of the control group respectively. Mares were allocated to two 2.5 ha paddocks of Cynodon sp., that had been previously grazed by young and adult horses and were naturally infested with equine cyathostomin larvae. Then, each animal of the treated group received twice a week 1 g pellets/10 kg live weight, containing D. flagrans mycelial mass combined with 100 g of horse commercial ration, as described by Assis and Araújo (2003). The treatment was offered during 6 months starting from May 2007. Animals of the control group received 1 g pellets/10 kg live weight without fungus. From the beginning (May) to the end (October) of the experiment, animals from both groups were monthly weighed. During the experiment, mares were fed daily with 2 kg of commercial ration with 14% soybean meal, 83.1% corn meal, 14.5% salt, 1.5% limestone and 14% protein.

After the mares had been moved to the paddocks, samples of fresh feces were collected once a week directly from the rectum, 72 h after the treatment, to determine egg per gram of feces (EPG), according to Gordon and Whitlock (1939) and modified by Lima (1989).

Coprocultures were established together with EPG counts; 20 g of feces were mixed with ground, moistened and autoclaved industrial vermiculite (NS Barbosa Ind. Com.,®) and taken to an oven at 26 °C, for 8 days, to obtain cyathostome larvae. Larvae were identified to the genus level as described by Bevilaqua et al. (1993). EPG and larvae recovered from coprocultures of animals of both treated and control groups were recorded and percentage of larval reduction was determined according to Mendoza-De-Guives et al. (1999):

\[
\text{reduction (\%)} = \frac{\text{mean } L_3 \text{ recovered from control group} - \text{mean } L_3 \text{ recovered from treated group}}{\text{mean } L_3 \text{ recovered from control group}} \times 100
\]

Every 15 days, two herbage samples were collected from both the treated and control groups, from each paddock, in a zigzag pattern from several and alternated points, 0–20 and 20–40 cm away from fecal pats, in each paddock of the different groups, according to Amarante et al. (1996). Herbage samples were always collected in the morning at 8 a.m. Then, a 500 g herbage sample was weighed, and parasitic nematode larvae were recovered following the procedure of Lima (1989). The samples were incubated in a drying oven at 100 °C, for 3 days, to determine dry matter. Data were transformed into larvae per kg of dry matter.

Climate data referring to averages of maximum, average and minimum monthly temperatures, air relative humidity and monthly rainfall were daily recorded in a meteorological station in the area.

The egg count curves (EPG) originated from the coprocultures, number of infective larvae recovered from paddocks (L3), correlation between EPG and recovered L3
and animal weight were compared over the experimental period. Data were transformed into log \((x + 1)\) and then examined by analyses of variance (ANOVA) and Tukey's multiple comparison test with 1% probability. The analyses were performed using the BioEstat 3.0 Software (Ayres et al., 2003).

3. Results

Fig. 1 shows the monthly mean EPG counts. EPG of animals treated with *D. flagrans* was significantly lower \((p < 0.01)\) than the control group, especially in the last 4 months of the experiment, in which the EPG monthly mean of the treated group was 46.2% lower than the control group. July, August, September and October showed smaller percentages of EPG reduction for fungus-treated animals than the control group: 35.4%, 73.2%, 64.3% and 30.5%, respectively. Additionally, fungus-treated animals had EPG values lower than the control group throughout the experiment. Fig. 2 shows the coproculture data. There was significant difference \((p < 0.01)\) between the results of fungus-treated animals and the control group in the last 4 months of the experiment (July, August, September and October) with larval reduction of 57.2%, 59.4%, 68.5% and 51% respectively.

Fig. 3 shows the number of larvae recovered from paddocks in the distances (0–20 cm) and (20–40 cm) away from the fecal pats. There was a significant difference \((p < 0.01)\) of 78.5% for the 0–20 cm samples between the treated group and the control, from May to October. Significant difference \((p < 0.01)\) of 82.5% was also found for the distance 20–40 cm from the fecal pats between the treated group and the control in the same period.

Fig. 4 shows the weights of animals from both groups. There was no significant difference \((p > 0.01)\) for animal weight during the first 3 months of the year (May, June and July) between the two groups. However, in the last 3 months of the experiment (August, September and October), significant differences \((p < 0.01)\) of 9.74%, 10.26% and 12.21%, respectively, were found for the weight between treated and non-treated animals.

4. Discussion

Amarante et al. (1996) states that the parameter EPG count allows evaluation of infection levels in animals and levels of pasture infestation by gastrointestinal nematode parasites. A number of studies on *D. flagrans* using horses and ruminants recorded average monthly EPG counts lower for treated animals than for non-treated groups (Baudena et al., 2000b; Knox and Faedo, 2001; Fontenot et al., 2003; Araújo et al., 2006; Paraud et al., 2007). The efficacy of *D. flagrans* on gastrointestinal parasites of...
ruminants was also demonstrated in the work of Dimander et al. (2003). These findings are in agreement with results obtained in the present work, confirming that the fungus acts on the infective forms in the fecal environment, with consequently decrease in EPG. There is nevertheless a lack of studies involving nematophagous fungi and equine cyathostomin (Bird and Herd, 1985; Baudena et al., 2000b).

Results seen in Fig. 2 suggest that there was a direct action of *D. flagrans* on infective cyathostomin larvae present in the pasture and a consequent lower parasitic infection of fungus-treated animals (Baudena et al., 2000a; Waghorn et al., 2003; Araújo et al., 2006). Only the occurrence of small strongyles (Cyathostominae) was observed after the coprocultures, according to the parameters described by Bevilaqua et al. (1993). Silva et al. (1993) reported that the subfamily Cyathostominae is highly prevalent in a large part of the Brazilian territory, and Carvalho et al. (1998) identified 19 species of small strongyles in necropsied horses in the state of Minas Gerais. The importance of these parasites for horses is directly related with larval cyathostomosis, a potentially fatal syndrome in most cases, and the high resistance of most gastrointestinal nematode parasites to routine antihelmintics (Reinemeyer, 1986; Reinemeyer and Herd, 1986).

The number of larvae recovered in the distances 0–20 and 20–40 cm from fecal pats (Fig. 3) is likely to be directly related with the use of nematophagous fungi that act directly on the *L*$_3$ present in pastures, confirming that *D. flagrans* was responsible for the satisfactory reduction of environmental contamination (Araújo et al., 2006).

In a work carried out to evaluate the survival and migration of cyathostomin in Tifton 85 grass (*Cynodon* spp.) at three collection times (8:00 a.m., 1:00 p.m. and 5:00 p.m.), Bezerra et al. (2007) recorded the largest number of recovered cyathostomin at 8:00 a.m., however, no statistical difference was found (*p* < 0.01) among the three times. Langrová et al. (2003), in a similar study in the Czech Republic, reported difference among collection times, with a higher cyathostome recovery at 8:00, 7:00 and 6:00 a.m. respectively. Hasslinger and Bittner (1984) discussed that temperature and moisture in the mornings favor the large number of *L*$_3$ recovered from pastures. In the present work, the largest number of infective larvae was recovered within the distance 0–20 cm away from the fecal pats. This result agrees with findings reported by Quinelato et al. (2008) and Dias et al. (2007b) who recorded larger numbers of larvae recovered within 0–20 cm from fecal pats, confirming that the few larvae that leave the feces migrate to the herbage beyond 0–20 cm. Stromberg (1997) points out that temperature and moisture are essential for the development of infective larvae. Only cyathostomin larvae were found in the herbage over the experimental period (May–October). Climatic conditions, such as temperature, relative humidity and rainfall favored the development of free-living stages and migration to the herbage (Figs. 5 and 6). The lowest rainfall rates occurred in July and August (12.64 and 16.96 mm$^3$ respectively), however, the larval count was high in this period due to accumulated larval loads. June and September had the highest rainfall rates (25.25 and 35.31 mm$^3$ respectively), with the smallest larval number recorded, possibly because the *L*$_3$ were washed off by rain (Figs. 3 and 6). Quinelato et al. (2008), working in the tropical southeastern Brazil, reported higher recovery of cyathostomin larvae from herbage and later from feces in the dry period, observing that the environmental conditions were favorable for recovering these larvae. The authors also argued that horses might be infected throughout the year in tropical climates, since *L*$_3$ are always present in the pastures and that the grass type can affect larval recovery. Langrová et al. (2003), in central Europe, suggests that *L*$_3$ respond to rain through dispersion within the vegetation, occurring a moderate correlation between moisture and *L*$_3$ number in the pasture.

Courtney (1999) observed that during the dry period, the *L*$_3$ development is slower, but they survive longer. Still, Fernández et al. (1997) and Baudena et al. (2000a) suggest that the survival of these parasites in the environment is strongly related with temperature and that few larvae would be found in feces in the summer. Baudena et al. (2000a) recorded field data in southern Louisiana, a region with subtropical climate in The United States, appearing that there is a larger number of infective larvae in the pasture in months with mild temperatures. This agrees with the results found in this work, in which the largest number of larvae recovered in pastures was found during months of mild temperatures (Fig. 3). Peña et al. (2002) and Chandrawathani et al. (2004) reported reduction of more than 90% of infective larvae present in fecal pats of ruminants using *D. flagrans*.

Fontenot et al. (2003) also discussed that besides *D. flagrans* decreasing infectious forms of gastrointestinal nematode parasites in pastures, it would avoid contamination of new animals entering these sites.
The correlation coefficient between EPG and infective larvae recovered from paddocks of group 1 within 0–20 cm from fecal pats was 0.0662; and for the distance 20–40 cm was 0.0416. For group 2, the correlation coefficient between EPG and infective larvae recovered within 0–20 cm from the fecal pats was −0.0394 and within 20–40 cm was 0.0401. These results showed weak, non-significant correlations, close to zero, nevertheless, as Dias et al. (2007b) pointed out, there might be dependence between EPG and infective larvae recovered from pastures even if the correlations are null. Besides, the availability of larvae on pasture may be determined by contamination from animals, as well as environmental factors, parasite and host (Lima et al., 1997).

*D. flagrans* is considered the most promising species in for biological control of gastrointestinal nematode parasites of livestock (Faedo et al., 2002). It has been used successfully in several laboratory and field studies (Araújo et al., 2006). Baudena et al. (2000b) proved the effectiveness of *D. flagrans* to reduce recovery of cyathostomin larvae from pastures. The authors reported reduction in the percentage of recovered larvae in animals that received doses of 2 × 10^6 spores/kg of live weight during 4 days compared with the control.

In a work testing two fungal isolates of the genus *Monacrosporium, Assis and Araújo* (2003) found fungal mycelia in horse feces up to 96 h after passing through the gastrointestinal tract of horses. In this work, *D. flagrans* was offered twice a week, for an efficient weekly coverage.

In Malaysia, Chandrawathani et al. (2003) confirmed the effectiveness of daily administration of *D. flagrans* to sheep. Terrill et al. (2004) also reported reduction of larvae in feces of goats infected with predominantly *Haemonchus contortus*. They also found that the daily administration of fungi (*D. flagrans*) was more effective than every 2 or 3 days. The frequency of treatments in this work promoted reduction of pasture contamination, mainly the weekly treatment.

The difference (*p < 0.01*) found in weight gain of treated animals compared to the control group may have been caused by a lower parasite load in animals that received pellets containing *D. flagrans* mycelia, which may have contributed to a better food conversion of treated animals. These results are similar to those found by Dias et al. (2007a) on weight gain of cattle treated with pellets containing *D. flagrans* mycelia.

The findings reported in this study suggest that the nematophagous fungus *D. flagrans* could be used in an integrated program to control horse cyathostomin in southeastern Brazil. It demonstrated the usefulness of a previous anthelmintic treatment to reduce the parasite load in animals and consequently the EPG, and starting from that to supply animal feed combined with the fungus to control the larval forms present in the environment and thus prevent reinfection.

5. Conclusion

Treatment of horses with pellets containing mycelial mass of the nematophagous fungus *D. flagrans* can be effective to control cyathostomin in tropical southeastern Brazil.

Acknowledgements

The authors would like to thank Fapemig and CNPq for the financial support and grant concession.

References


