Evaluation of immune responses and oxidative stress in donkeys: Immunological studies provoked by Parascaris equorum infection

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Abstract

This study aimed to assess the effects of Parascaris equorum (P. equorum) in infected donkeys through evaluation the oxidative stress and different gene parameters in infected tissues. Fifty donkeys were examined in Giza Zoo abattoir from the period of January to March 2021. Blood and sera samples were collected from each examined donkey. P. equorum were subjected for identification through scanning electron microscope study and the infected tissues were subjected into gene expression analysis using two genes; interleukin 1β (IL-1β); and pro-inflammatory cytokines (TNF-α) with assessment of the antioxidant and free radicals released from the animals during the infection. Eighteen donkeys were positive for P. equorum adult or larvae by postmortem examination of the intestine and abdomen with prevalence rate of 36 %. The examined infected donkeys with P. equorum showed significantly higher of Total antioxidant capacity (TAC) levels and the serum malondialdehyde (MDA) 2.45 ± 0.53 than that in non-infected control donkeys. The levels of AST enzyme were 278.54 ± 0.45 while ALT enzyme was 14.97 ± 0.87 which was significantly higher than that of control negative donkeys. The infected donkeys exerted at least 100 eggs of parasite in feces. The fecal egg count was marked decreased after treatment with moxidectin. Moxidectin is considered a novel active ingredient that has a marvelous result with high persistency and protection for long time, in addition to, broad spectrum activity and low or no resistance. We recommend the periodical deworming with different molecules as more economic and lifesaving over a single treatment every 12 months parallel with parasitic testing.

1. Introduction

Donkeys (Equus asinus) are one of early-domesticated equines, they are easy to rear, which representing a cheap mean for human transportation; about 40 million donkeys distributed world-wide and 13 million in Africa, while around 3 million working donkeys approximately in Egypt (Hilali et al., 2015; Attia et al., 2018; Attia and Mahdy, 2021). They are accustomed to low quality forages or grass and spread widely in Africa and rural areas (Ahmed et al., 2011). Unfortunately, donkeys subjected to bad management, work for long hours under stress, malnutrition and fed with low quality grass and forages which considered as source for their infection with a variety of parasites (Parsani et al., 2013; Jajere et al., 2016; Mahdy and Attia, 2021). In developing countries, one of the most prominent factors limiting the health and performance of donkeys is gastrointestinal parasites, where it considered as the most serious disease problem for horses and donkeys (Pereira and Attia, 2021).
Donkeys infected with a wide range of internal parasites as roundworms (families: Ascaridae, Trichostrongylidae, Oxyuridae and Strongylidae) and tapeworm (family: Anoplocephalidae) which induce severe gastrointestinal damages that appears in shape of diarrhea, inappetance, weight loss, poor performance, general weakness, colic, and may end with death in case of heavy infestation subsequent to intestinal impaction, obstruction or perforation (Pereira and Vianna, 2006; Getahun and Kassa, 2017; Attia et al., 2018). *Parascaris equorum* is one of gut helmhnts that represent a serious problem for equines, it has a direct life cycle takes about three months in which the adult round worm lives in the small intestine lies millions of rebellious eggs that passed daily in equine feces to the environment and develop into infective stage within 10–14 days that ingested by a new host and hatch in its intestine, larvae liberate and migrate through liver, lungs and may attack other tissues for 2–4 weeks, then return to small intestine and mature (Urquhart et al., 1988; Rose and Hodgson, 1993; Getachew et al., 2008). Parascaris equorum infection revealed inflammatory lesions mainly in small intestine, gut ischemia, severe colic, intestinal obstruction, intestinal rupture, toxemias, and death. Also, respiratory manifestation (nasal discharge, intermitent cough, and dyspnea); liver lesions and nervous manifestations may be occasionally observed due to larval migration (Rose and Hodgson, 1993; Getachew et al., 2008). Many diagnostic tools were used for parasitic detection as fecal examination, ELISA, and PCR. Enzyme Linked Immunosorbent Assay (ELISA) is a quantitative, highly sensitive technique customized for the detection of target analyte found in biological samples as serum, and plasma, where ELISA be used for detection and accurate quantification of substances such as cytokines, hematological factors, hormones, peptides, and immunoglobulins (Shalaby et al., 2008). Anthelmintic resistance specially against cyathostomine (small strongyle) and other nematodes has been extended to many active ingredients as benzimidazoles or pyrantels (50% of populations) and occasional Piperazine (Drudge et al., 1983; Kaplan, 2004; Kaplan et al., 2004; Bzikaz et al., 2006). Similarly, *P. equorum* has been found to be resistant to fenbendazole, Piperazine and macrocyclic lactones, particularly Ivermectin, which has been observed in numerous countries (Peregrine et al., 2014) and Egypt (Ali et al., 2015). On the other hand, moxidectin alone or in combination with praziquantel found to be effective treatment against internal worms in horses (Cobb and Boeckh, 2009). As a result, there is a need to quantify found to be effective treatment against internal worms in horses (Cobb and Boeckh, 2009). As a result, there is a need to quantify found to be effective treatment against internal worms in horses (Cobb and Boeckh, 2009). As a result, there is a need to quantify found to be effective treatment against internal worms in horses (Cobb and Boeckh, 2009). As a result, there is a need to quantify targeted capacity released through *P. equorum* infection with treatment trial to evaluate the efficacy of moxidectin in the treatment of such infection.

### 2. Materials and methods

#### 2.1. Animal and sampling

Fifty donkeys were examined in Giza Zoo abattoir from the period of January to March 2021. Blood was collected from each examined donkey in sterilized tubes with and without EDTA. The samples were collected and preserved as recorded in Attia et al. (2020).

The abdomen of each donkey was opened, and the intestine was removed and opened to examine the presence of *P. equorum*. As well as; collection of fecal samples from each examined donkey. The labeled fecal samples and the *Parascaris* nematodes as well as the blood and sera were collected and examined in the Faculty of Veterinary Medicine, Cairo University for further analysis.

#### 2.2. Fecal samples

Examination of fecal samples were done to exclude other internal parasitic infection where eggs or oocyst were traced using direct fecal smear and flotation technique to exclude their presence (Soulsby, 1986). Therefore, the only used donkeys’ sera as reference was that free from all parasites except presence of *P. equorum* either eggs or adults.

#### 2.3. Blood and sera samples

Ten ml of blood samples from each animal were collected during slaughtering from jugular vein; on plain tube which used for estimation of different biochemical analysis as aspartate amino transferase (AST) and alanine amino transferase (ALT) following the specific test kits instructions (spectrum diagnostics, Egypt). Also, thin blood film from each animal whole blood with EDTA was done and stained with Giemsa stain for detecting any blood parasites (Zaki et al. 2021).

#### 2.3.1. Identification of *P. equorum* Adult and Eggs

All the collected nematodes from intestine and its eggs from feces were examined in the parasitology laboratory of the Faculty of Veterinary Medicine, Cairo University for larval identification. The identification of the collected nematodes was done following Soulsby, (1986); Attia et al. (2018).

#### 2.3.2. Ultrastructure Identification of The Collected *P. equorum* using Scanning Electron Microscopy (SEM) study (JOEL)

Adult *P. equorum* were washed several times using 0.9% saline (Attia and Salem, 2021). The collected worms were fixed in 2.5% Glutaraldehyde following to Abdelsalam et al., (2020); then the worms were removed from Glutaraldehyde and dehydrated using ethanol series (50%; 70%; 90%; 100%); then the fixed Nematoda was dried in CO2 critical point drier (Autosamdri-815, Germany). The adult was glued over stubs (as anterior end and posterior end) and then; coated with 20 nm gold (Abu-Elala et al., 2018) in a sputter coater (Spi-Module sputter Coater, UK). All the specimens were photographed with a scanning electron microscope (JSM 5200, Electron probe); Microanalyzer Jeol, Japan; at Faculty of Agriculture, Cairo University, as described by Salem and Attia, (2021).

#### 2.4. Assessment of the oxidative stress markers

Oxidative stress markers were studied in sera samples as malondialdehyde (MDA); Total antioxidant capacity (TAC) according to Aytekin and Unubol Aypak (2011); Aktas et al., (2017); Salem et al., (2018); Attia et al., (2019).

#### 2.5. Evaluation of pro-inflammatory cytokines (TNF-α) and interleukin 1β

Infected intestine with the parasites were aseptically dissected and then preserved in freezer in –20°C. Samples from 5 uninfected donkeys were free from any parasites and have no gross lesion used as negative controls (Attia et al., 2020).
2.5.1. RNA Isolation
Isolation of mRNA by total RNA kit (Ambion, Applied Biosystems), from 200 mg of infected intestine with *P. equorum*. Homogenization of the intestinal tissues were applied in Lysing Matrix D tubes (MP Biomedicals) using a FastPrep-24 homogenizer (MP Biomedicals, 2 cycles of 30 s at 6 m/s). Nanodrop (Thermo Scientific) were assessed the mRNA purity and quantity. A 500 ng of mRNA were made with DNaseI amplification grade (Invitrogen) following the manufacturer’s instructions. The reverse transcription of treated RNA was performed by High-Capacity cDNA Archive Kit (Applied Biosystems) (Attia et al., 2020; Younis et al., 2020).

2.5.2. Quantitive Real-Time PCR protocol (qRT-PCR)
PCR primer sets were designed according to presence in Gene Bank specific for donkeys (*Table 1*). The reference gene used was the β-actin and used for sample normalization. Real-time PCR run protocol were followed according to Attia and Mahdy, (2021).

2.6. Treatment trials
Ten naturally infected donkeys with *P. equorum* in a private collecting station were subdivided into two groups; 1st group was treated with moxidectin (Equest®, Zoetis), at a dose of 0.4 mg/kg oral paste with interval of 3 months and 2nd group kept untreated. The efficacy of the treatment was estimated by fecal egg count (FECR) test. Individual freshly voided fecal samples were collected weekly for one month using veterinary gloves. All fecal samples were kept in labeled plastic pages and transported on ice box rapidly to Parasitology Department, Faculty of Veterinary Medicine, Cairo University for further investigations. Fecal samples were stored under refrigeration and examined within 24 h. Fecal egg counts were estimated using modified McMaster technique (Raynaud, 1970) using saturated solution of sodium chloride (specific gravity 1.20).

2.7. Statistical analysis
Data were expressed as means ± standard errors and the data were statistically analyzed using independent t-test of ANOVA test using SPSS Inc., Chicago, IL, USA; Version 18.0 software. *P*-value was considered significant when it less than 0.05.

3. Results
Eighteen examined donkeys were positive for *P. equorum* adult or larvae by postmortem examination of the intestine and abdomen with prevalence rate of 36% (*Figs. 1 and 2*).

The adult *P. equorum* were whitish in color, and cylindrical; the anterior end had 3 large lips with deep transverse groove, lips were large; crown and prominent, 3 in number, one dorsal and two sub-

| Table 1 | The sequences of the forward and reverse primer used in the quantitative real-time PCR. |
|---|---|---|---|
| **Genes** | **sequence** | **Accession number** | **References** |
| IL-1β | F: AAAACAGTGAGGGAGAAATT  
R: AGAAACTTCTTCTTGGGTAG | XM_014852743 | Huang et al., 2015 |
| TNF-α | F: ATGTTTCAGTCACATTTCAG  
R: CCTACCGGTTCCCATCTCAA | XM_014831267 | Huang et al., 2015 |
| β-Actin | F: CAGCAAGCAGGAGTACGATGAG  
R: TGTGTGGTGTGTGGTTTTG | AF035774 | Swiderski et al. 1999 |

![Fig. 1](https://example.com/f1.png) **Fig. 1.** Life cycle of *P. equorum*; the adult worms inhabit the small intestine of donkey. The worm lay eggs that passed into the feces. Eggs have a thick shell as appeared under microscope. Eggs are then ingested as in the infective form as egg containing second stage larvae within the contaminated grass or drinking water by the donkey. After that, the ingested infective stage become a free larva that penetrate the small intestines and migrate into blood stream to the liver and further to the lung to irritate the animal to coughed up and re-swallowed again.
In the infected donkeys with 20 worms of *P. equorum* significantly showed upregulation of the TNF-α by 18 than that of control non-infected donkeys while the gene expression analysis of TNF-α increased by 25 in donkeys with 21–30 collected worms; the TNF-α raised to 28-fold when the donkeys harbored more than 30 worms (Table 3).

In the infected donkeys with 20 worms of *P. equorum* showed significantly upregulation of the IL-1β by 10 than that of control non-infected donkeys while the gene expression analysis of IL-1β increased by 17 in donkeys with 21–30 collected worms; the IL-1β raised to 26-fold when the donkeys harbored more than 30 worms (Table 3). Naturally infected donkeys with *P. equorum* shed at least 100 eggs per gram feces. The results of fecal samples examination are summarized in Table 4. The fecal egg count was marked decreased after treatment with moxidectin as seen in Fig. 4.

4. Discussion

*P. equorum* is a common ascarid nematode which inhabits the small intestine of family Equidae mainly the young animals which is highly susceptible. The female adult Nematoda lays its eggs in small intestine; these eggs develop into eggs with larva (L2) in the environment full mature eggs takes 10 days to become infective in temperature between 25 and 35 °C. Donkeys are prone to endemic infection with gastrointestinal helminths (Jajere et al., 2016). From our results, the prevalence of *P. equorum* infection was 36 % among examined donkeys. This prevalence was higher than that recorded by Attia et al., (2018) and Shrikhande et al.
and E levels were significant decrease \((\text{CAT})\) were significantly increased \((p<0.05)\) in pneumonic \(P.\ equorum\) eggs during fecal examination using saturated salt solution with floatation technique before treatment. While \(P.\ equorum\) eggs after treatment.

From our results, moxidectin is an effective treatment for \(P.\ equorum\) in donkeys. This result agreed with Cobb and Boeckh, (2009) as they found combination between moxidectin and prazi-quantel is effective broad spectrum, prolonging acting, enabling prolonged treatment interval against intestinal helminths in horses, while Reinemeyer and Marchiondo, (2007) recorded that moxidectin failed to reduce fecal egg count of \(P.\ equorum\) in horses. Handling the parasitic problem in animals should be achieved using the biosecurity measures as well as the improved health status of the animals using environment-friendly products such as probiotics (Abd El-Hack et al., 2021a; Alagawany et al., 2021a; El-Saadony et al., 2021a), prebiotics (Abd El-Hack et al., 2021b; Yaqoob et al., 2021), essential oil (El-Tarabily et al., 2021; Alagawany et al., 2021b), bioactive peptides (El-Saadony et al., 2021b; El-Saadony et al., 2021c), herbal extracts (Abou-Kassem et al., 2021; Saad et al., 2021; Abd El-Hack et al., 2021c), green synthesized nanoparticles (Attia et al., 2017; El-Saadony et al., 2021d, e,f) should be adopted in the animal facility.

### Table 2

| Parameters | Infected donkeys (n = 18) | Control donkeys (n = 5) |
|------------|---------------------------|-------------------------|
| AST (U/l)  | 278.54 ± 0.45*            | 240.5 ± 0.18            |
| ALT (U/l)  | 14.97 ± 0.87*             | 8.98 ± 0.96             |
| TAC (mmol/L) | 0.57 ± 0.08*             | 0.78 ± 0.57             |
| MDA (mmol/L) | 2.45 ± 0.53*             | 0.88 ± 0.67             |

*Mean fecal egg count per gram feces.

### Table 3

| Parameters | Infected donkeys (n = 18) | Control donkeys (n = 5) |
|------------|---------------------------|-------------------------|
| TNF-α      |                          |                         |
| IL-1β      |                          |                         |

(e,f) should be adopted in the animal facility.
5. Conclusions

The development of resistance against anthelmintic against parasites at broad and *P. equorum* as well means that traditional selection of anthelmintics must be shifted routinely. In the future, every 12 months to combat and control parasitic infection in equine in Egypt parallel with parasitic testing to identify the risky animals that considered as source of infection.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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