Location, distribution, and quantification of myenteric plexus neurons of the jejunum of quails fed with different levels of commercial Macleaya cordata extract

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ABSTRACT: Coturniculture has been promising, progressing from a subsistence to a technical activity due to its quick production, low breeding investment, and rapid economic return. After the restriction of antimicrobials as growth promoters, some studies aimed to evaluate alternative products that would make the farming of healthy birds viable without impacting their performance, with commercial Macleaya cordata extract being one of these substitutes. The functions of the gastrointestinal tract are coordinated mainly by the enteric nervous system, and the myenteric plexus is responsible for the reflex control of contractile activities of the external muscles. Thus, this study located and demonstrated the distribution of the myenteric plexus, quantifying the total population of myenteric neurons (Giemsa+), and the subpopulation of myenteric nitrergic neurons (NADPH-d+), and evaluated the effects of commercial Macleaya cordata extract on these populations of jejunal neurons. A total of 240 one-day-old female laying quails were distributed into four treatments, with four repetitions of 15 birds each. The test groups (T1, T2, and T3) were treated with commercial Macleaya cordata extract throughout the experimental period using the following doses: T1 - test group, basal diet added with 150 ppm of the extract in the feed; T2 - test group, basal diet added with 100 ppm of the extract in the feed; and T4 - control group, basal diet with no added extract. The study included histological analysis, Giemsa+, and NADPH-d+ myenteric neuron staining. The results showed that the myenteric plexus is located between longitudinal layer fibers and in the transition region between the longitudinal and circular layers of the muscular tunic, with the myenteric population organized into ganglia and isolated in the region of neuronal fiber bundles. The commercial Macleaya cordata extract showed no quantitative changes in the myenteric Giemsa+ population and myenteric NADPH-d+ subpopulation, however, the groups that consumed the extract showed greater NADPH-d+ neuron activity compared to the control group, implying that the food remained longer in the intestinal lumen, therefore, enabling greater nutrient use and resulting in increased productive performance.

Key words: benzophenanthridine and protopine alkaloids, Coturnix coturnix japonica, enteric nervous system, Sangrovit.

Localização, distribuição e quantificação dos neurônios do plêxus mioentérico do jejuno de codornas de postura alimentadas com diferentes inclusões do extrato comercial de Macleaya cordata

RESUMO: Coturnicultura tem apresentado características promissoras, devido a ser uma atividade de subsistência e ocupando patamares tecnofificados devido a sua precocidade produtiva, baixo investimento de criação e rápido retorno econômico. A partir da restrição da utilização de antimicrobianos como promotores de crescimento, estudos foram direcionados com o objetivo de se avaliar produtos alternativos que viabilizassem a criação de aves saudáveis, sem comprometer seu desempenho, sendo o extrato comercial da Macleaya cordata um destes substitutos. As funções do trato gastrintestinal são coordenadas principalmente pelo sistema nervoso entérico, sendo o plêxus mioentérico responsável pelo controle reflexo das atividades contraóritas da musculatura externa. Desta forma, o presente trabalho teve como objetivo localizar e demonstrar a distribuição do plêxus mioentérico, quantificar a população total de neurônios mioentéricos (Giemsa+), e a subpopulação de neurônios mioentéricos nitrérgicos (NADPH-d+), além de avaliar os efeitos do extrato comercial da Macleaya cordata sobre estas populações de neurônios do jejuno de codornas. Foram alojadas 240 codornas de postura, fêmeas, com um dia de idade, distribuídas aleatoriamente em quatro tratamentos, com quatro repetições de 15 aves cada. Os grupos testes (T1, T2, e T3) foram tratados com extrato comercial de Macleaya cordata durante todo o período experimental conforme as doses indicadas, sendo: T1 - grupo teste, com dieta basal adicionado de 150 ppm do extrato na ração; T2 - grupo teste, com dieta basal adicionado de 100 ppm do extrato na ração; e T4 - grupo controle, com dieta basal isenta do extrato. Foram realizadas análises histológicas e a marcação dos neurônios mioentéricos Giemsa+ e NADPH-d+. Os resultados demonstraram que o plêxus mioentérico está localizado entre as fibras do extrato longitudinal, e na região de transição entre os estratos longitudinal e circular da ínﬁca muscular, estando a população mioentérica organizada em ganglios, e também isoladamente na região dos feixes das fibras nervosas. O extrato comercial de Macleaya cordata não alterou quantitativamente os neurônios da população mioentérica Giemsa+ e da subpopulação mioentérica NADPH-d+, mas os grupos que consumiram o extrato apresentaram maior atividade dos neurônios NADPH-d+ em relação ao grupo controle, permitindo inferir que o alimento permaneceu maior tempo no lúmen intestinal e, portanto, possibilitou um maior aproveitamento dos nutrientes, podendo reﬂetir em melhor desempenho produtivo.

Palavras-chave: alcaloides da benzofenantridina e protopina, Coturnix coturnix japonica, sistema nervoso entérico.
INTRODUCTION

Coturniculture is the segment of the poultry industry responsible for raising, promoting, and managing quails (SILVA et al., 2018), standing out from its modernization, followed by new production technologies, progressing from a subsistence to a technical activity, with positive results to investors (PASTORE et al., 2012).

Besides favorable results, coturniculture has encouraging characteristics such as low initial investment and fast economic return, with quails being fast-growing birds with good feed conversion, early sexual maturity (35 to 42 days), and productive longevity (14 and 18 months), which requires small farming spaces (MURAKAMI & ARIKI, 1998; PASTORE et al., 2012). In this context, quail farming plays an important social role, providing a source of income for family agriculture and small rural producers (SILVA et al., 2018).

In the development and modernization of coturniculture, associated with the current restriction on the use of antimicrobials as growth promoters in animal nutrition, some studies aimed to evaluate alternative products to replace growth promoters in animal feed without decreasing productivity and enabling the rearing of healthy birds (BONATO et al., 2008; OTUTUMI et al., 2009) and safety food for the consumer, eliminating the possibility of residues and bacterial resistance.

According to KANTAS et al. (2015), Macleaya cordata extract meets the requirement of animal production free of antibiotics and growth promoters, increasing performance and profitability. SANGROVIT® ED is a natural product made from plants of the family Papaveraceae, which contains 1% Macleaya cordata extract in its composition (SANGROVIT® ED, 2019).

High bird productivity depends on adequate nutrients for the body. For nutrients to be digested and absorbed, the intestinal mucosa needs satisfactory morphophysiological structures, since digestion depends on mechanisms occurring in the intestinal wall, with its preservation and integrity being of vital importance (PATRÍCIIO, 2016).

Gastrointestinal functions are coordinated mainly by intrinsic neurons of the enteric nervous system (ENS), besides the participation of autonomic extrinsic neurons of the sympathetic and parasympathetic pathways and sensory neurons (BAYLISS & STARLING, 1899; FRAUCHES et al., 2016; HANSEN, 2003; LANGLEY, 1921).

The ENS is organized in the form of plexuses, composed of numerous ganglia of various sizes along its extension, with a greater concentration of intestinal nervous cells arranged into two sets of ganglia. The ganglia of the myenteric plexus are responsible for controlling the reflex of contractile activities of the external musculature, while the ganglia of the submucous plexus are responsible for coordinating secretomotor and vasomotor activities of the mucous tunic (FRAUCHES et al, al, 2016; FURNESS, 2006; SCHEMANN, 2005).

The myenteric plexus is described as a nerve network with small ganglia (FURNESS, 2006) along the gastrointestinal tract from the esophagus to the rectum, located between the circular and longitudinal layers of the muscular tunic (FRAUCHES et al, al, 2016; FURNESS & COSTA, 1980; FURNESS, 2006; HANSEN, 2003), with neuron organization and density varying with animal species and the segment of the digestive tract being studied (GABELLA, 1981; IRWIN, 1931) and presenting an easily identifiable mesh format (GABELLA, 1981).

FRAUCHES et al. (2016) and HANSEN (2003) reported that enteric neurons express different neurotransmitters that define their neuronal function. FURNESS (2000), FURNESS (2006), and TAKEUCHI et al. (2005) reported that acetylcholine is the main excitatory neurotransmitter in smooth muscles of the digestive tract that acts through M₄ muscle receptors (TAKEUCHI et al., 2005), presenting the substance P as co-transmitter (FURNESS, 2006). In addition, FURNESS (2000) and FURNESS (2006) described that nitric oxide (NO) is an inhibitory neurotransmitter synthesized when necessary from the activation of the enzyme nitric oxide synthase (NOS), present in inhibitory enteric neurons, which, according to BROOKES (1993), promotes the relaxation of gastrointestinal smooth muscles.

This study was conceived considering the properties of Macleaya cordata extract as an alternative feed additive for production animals, due to the growing interest in coturniculture, the scarcity of morphophysiological studies on the myenteric plexus of quails, and the lack of studies on the effects of Macleaya cordata extract on the myenteric plexus of quail, this study aimed to locate the myenteric plexus, demonstrating its distribution in this plexus, quantifying the total population of myenteric neurons (Giemsa+) and the subpopulation of myenteric nitrergic neurons (NADPH-d+), and evaluating the effects of Macleaya cordata extract on these populations of laying quail jejunal neurons.
MATERIALS AND METHODS

This study was evaluated and approved by the Committee on Animal Research and Ethics (CARE) of the Paranaense University (UNIPAR) under protocol number 31660/2017.

Animals, treatment, and environment

The experiment was conducted in the Experimental Aviary, UNIPAR Campus II, with 240 one-day-old female laying quails (Coturnix coturnix japonica) with a mean weight of 6.31 g, from the Vicami Codornas, Assis, SP, being randomly distributed into four treatments, with four repetitions of 15 birds each. The test groups (T1, T2, and T3) were treated with commercial Macleaya cordata extract through out the experimental period using the following doses: T1 - test group, basal diet added with 150 ppm of the extract in the feed; T2 - test group, basal diet added with 100 ppm of the extract in the feed; T3 - test group, basal diet added with 50 ppm of the extract in the feed; and T4 - control group, basal diet with no added extract. The quails of 1 to 35 days of age were kept in boxes with bedding at an ideal comfort temperature for their age, with ad libitum water and balanced feed (ROSTAGNO et al., 2017) without the addition of anticoccidials or growth promoters produced in the Laboratory of Animal Nutrition, UNIPAR Campus II.

Commercial Macleaya cordata Extract - Sangrovit® ED

Sangrovit® ED is a feed additive consisting of the extract of the Macleaya cordata plant at 1% concentration having benzophenanthridine (sanguinarine and chelerythrine) and protopine (protopine and allocryptopine) alkaloids as bioactive compounds, which improve intestinal integrity at a 1% concentration having benzophenanthridine (sanguinarine and chelerythrine) and protopine (protopine and allocryptopine) alkaloids as bioactive compounds, which improve intestinal integrity.

Euthanasia of the animals and jejunum fragment collection

After the 35-day trial period at the Experimental Morphology Laboratory of the Graduate Program in Animal Science with Emphasis on Bioactive Products, UNIPAR central campus, one quail per repetition was randomly selected to be euthanized, totaling four birds per group, using an anesthetic protocol with xylazine hydrochloride as pre-anesthetic medication at an intramuscular dosage of 4 mg/kg, and sodium thiopental as anesthetic at an intramuscular dosage of 25 mg/kg.

Subsequently, the birds were necropsied and three jejunum fragments were collected 2 cm anteriorly to the yolk diverticulum (Meckel’s diverticulum). The first segment was used for Giemsa positive myenteric neuron staining (Giemsa+). The second fragment was used to detect NADPH positive myenteric neurons (NADPH-diaphorase), which respond to the neurotransmitter NO, also called nitrergic neurons. Lastly, the third fragment was used to locate the myenteric plexus using the hematoxylin-eosin (HE) technique.

Giemsa+ myenteric neuron staining: evidence of total neuronal population, according to BARBOSA (1978)

To stain the Giemsa+ myenteric neurons, fragments of the jejunum of each bird were washed with 0.9% saline solution. The oral and aboral ends were sutured with cotton thread, and the interior was filled with Giemsa fixative solution, being immersed in the same solution until membrane preparations were obtained. Each membrane preparation was then placed in a Giemsa staining solution containing methylene blue in Sorensen phosphate buffer (pH 6.9) for 24 hours at room temperature. Moreover, the membrane preparations were dehydrated in an alcohol sequence (95%, Absolute I, Absolute II) and were diaphanized with two consecutive Xylol immersions (Xylol I and II, for five minutes each). Each membrane preparation was placed on a slide and coverslipped with synthetic resin.

Obtaining membrane preparations

After the fixation period, the jejunum sutures were removed, and the segment was sectioned transversely to obtain a fragment of approximately 8 mm in width. The fragment was then sectioned along the longitudinal axis of the mesenteric border and micro-dissected in a glass plate using a transilluminated stereomicroscope forceps, removing the mucous and submucous coats and preserving the muscular and serosal.

Myenteric nitrergic neurons detection (NADPH-d+): evidence of nitrergic neuronal subpopulation, according to SCHERER-SINGLER et al. (1983)

To detect myenteric nitrergic neurons (NADPH-d+), fragments of the jejunum of each bird were washed with sodium phosphate buffer solution (PBS) (pH 7.4), sutured at the oral and aboral ends with cotton thread, and had their interior filled with the same solution, being subsequently immersed in 4% paraformaldehyde for 30 minutes and then washed in PBS containing Triton X-100 at 0.3% for ten minutes. The jejunum fragments were then washed again in PBS three times (ten minutes each) and incubated for
90 minutes in a reaction medium containing 50 mg Nitro Blue Tetrazolium (NBT), 100 mg β-NADPH, and 0.3% Triton X-100 in Tris-HCl buffer (0.1M, pH 7.6). After this, the jejunum fragments were again washed in PBS for three more times (five minutes each), and at the end of this period, the fragments were removed and were immersed in 4% paraformaldehyde solution to interrupt the reaction, and for fixation and storage, obtaining membrane preparations as previously described. Subsequently, the membrane preparations were dehydrated in an increasing alcohol sequence (80%, 90%, Absolute I, and Absolute II), followed by diaphanization in Xylol (I and II), and placed on aslide and coverslipped with synthetic resin.

Quantification of neurons evidenced by the Giemsa and NADPH-d histochemistry techniques

The model proposed by SANT’ANA et al. (1997) was used to guide the uniform capture of mesenteric, intermediate, and antimesenteric images. Under light microscopy (Nikon Eclipse E200), the material obtained was visualized with a 40x objective and the images were transferred from the microscope to a computer using an image analysis system coupled with a high-resolution photographic camera (Moticam 5.0 megapixels), with 120 random microscopic fields being captured per membrane preparation. In addition, the images of mesenteric, intermediate, and antimesenteric areas of the jejunum were quantitatively analyzed for myenteric neurons stained by the Giemsa and NADPH-d histochemistry techniques, with the neuron mediums being considered in alternate fields.

Statistical analysis

The data was subjected to descriptive analysis using the BioEstat 5.0 software (AYRES et al., 2007). The normality was evaluated (Lilliefors). The data regarding the NADPH-d+ neuron count showed normal distribution, which was compared using analysis of variance (ANOVA). The data on Giemsa+ neuron counts were not normally distributed, which were compared using the Kruskal-Wallis test. A 5% significance level was considered for all data.

RESULTS AND DISCUSSION

The myenteric population (Giemsa+) and the nitrergic subpopulation (NADPH-d+) of the myenteric plexus of laying quails (Coturnix coturnix japonica) fed with different levels of commercial Macleaya cordata extract were evaluated.

The histological sections showed that the myenteric plexus is located between the fibers of the longitudinal layer and in the transition between the fibers of the longitudinal and circular layers of the muscular tunic of the jejunum (Figure 1). FRAUCHES et al. (2016), FURNES (2006), and HANSENS (2003), reported that the myenteric plexus is in this position from the esophagus to the rectum, and LI et al. (1994) reported that it is in the small intestine of broiler quails (Coturnix coturnix coturnix).

The myenteric Giemsa+ population and the NADPH-d+ neuron subpopulation were arranged in ganglia, which were isolated between the neuronal fibers that establish communication between the ganglia (Figure 2). LI et al. (1994) verified in the European quail gizzards, however, data regarding their distribution in the jejunum was not found. PREVIATO DO AMARAL et al. (2017) studied NADPH-d+ neurons in the duodenum of broiler chickens (Gallus gallus domesticus) aged 21 days; they reported neurons in separated ganglionic arrangements between the fibers that interconnect the ganglia. YANG et al. (2013) analyzed the distribution arrangements between the fibers that interconnect the ganglia. YANG et al. (2013) analyzed the distribution
lower number of nitrergic neurons does not necessarily indicate a lower number of existing nitrergic neurons in the animal, because only the active neurons are demonstrated with the technique used.

Studies on other bird species, such as the one by PREVIATO DO AMARAL et al. (2017), found a density of 24.38±4.97 NADPH-d+/mm² neurons in the duodenum of broiler chickens aged 21 days. YANG et al. (2013) reported 45.92±17.51 NADPH-d+/mm² neurons in the jejunum of broiler chickens aged 40 days. These authors suggested that smaller animals may present higher neuronal density of the myenteric plexus. However, the present study showed lower myenteric nitrergic density than YANG et al. (2013) in broiler chickens; however, a higher density was reported by PREVIATO DO AMARAL et al. (2017).

SERENINI (2020) emphasized that the quantitative variations in neurons of the myenteric plexus should consider intra-species differences, such as animals of different ages, different intestinal regions studied, as well as the region of the sampled organ. In addition, interspecific differences must also be evaluated, such as the feeding habit of the species, its body mass, and the regions of intestinal fragment collection.

To stain the total myenteric neuron population, the present study used the Giemsa technique, with no significant differences between treatments, as well as with the control group (Table 1). According to BARBOSA (1978), the Giemsa staining technique, which uses the methylene blue dye in digestive tract membrane preparations, can be used as one of the available methods to evaluate the total myenteric neuronal population due to the affinity of the dye for acid structures in nerve cells (SANT’ANA et al., 2012). GÓIS et al. (2016) reported that the polyribosomes of all neurons can be stained by the Giemsa technique.

It should be noted that this staining aimed to evaluate the nitrergic subpopulation in relation to the total number of myenteric neurons in the jejunum of quails, estimating the number of active neurons in this subpopulation under the research conditions.

The mean NADPH-d+ neurons showed in groups T₁, T₂, T₃, and T₄ corresponded to 26.72%, 23.66%, 26.36%, and 17.79% of the population of Giemsa+ myenteric neurons, respectively. According to LI et al. (1994), nitrergic neurons comprised one third of the myenteric neurons in each sampled region of the gastrointestinal tract of European quails. Several studies with different animal species reported similar results, such as SERENINI (2020), with 20.4% active NADPH-d+ neurons in the jejunum of bats, FEREZIN et al. (2017), with approximately 26.0% NADPH-d+ neurons in the colon of Wistar quails.

Ciência Rural, v.51, n.11, 2021.
rats, FURNESS (2006), with 23.0% NADPH-d+ neurons in the small intestine of guinea pigs, and QU et al. (2008), with 29.0% NADPH-d+ neurons in the ileum of mice.

As NO is an extremely specific neurotransmitter produced when the NOS enzyme is activated (FURNESS, 2006), it is possible to infer that a lower nitrergic activity, as observed in the control group, increases the speed of food transit, therefore reducing its time in the intestinal lumen, consequently decreasing nutrient use and increasing food conversion (Table 2). SANDERS & WARD (1992) studied the colon of guinea pig neck and verified that NO promoted relaxation of the intestinal smooth muscle, allowing greater absorption capacity, resulting in greater fluid and electrolyte absorption (MIZUTA et al., 1999).

The bioactive compounds of Macleaya cordata extract include quaternary benzophenanthridine (sanguinarine and chelerythrine) (KOSINA et al., 2004; OLIVEIRA, 2012; SANGROVIT® ED, 2019; SIMANEK et

Table 1 - Density of myenteric neurons revealed by the Giemsa method (A to D) and by NADPH-diaphorase histochemistry, corresponding to 1.00 mm² of the jejunum of laying 35-day-old quails fed with different levels of commercial Macleaya cordata extract.

| Treatments | NADPH+ | GIEMSA+ |
|------------|--------|---------|
| T1 - 150 ppm | 42.3   | 158.3   |
| CV%  | 27.76  | 11.03  |
| T2 - 100 ppm | 34.5   | 145.8   |
| CV%  | 13.90  | 32.28  |
| T3 - 50 ppm | 42.3   | 160.5   |
| CV%  | 31.54  | 23.16  |
| T4 - Control | 40.3   | 226.5   |
| SEM | 39.62  | 14.87  |
| P-value | -5.75  | -16.90  |
| CV%  | -0.7831| -0.0664 |

CV%: Coefficient of variation; SEM: Standard error of mean.
The myenteric plexus is located between longitudinal layer fibers and longitudinal and circular layers of the muscular tunic, similar to the other species studied. Furthermore, the myenteric population is mostly organized in ganglia, however, it is isolated between the neuronal fibers that establish the communication between the ganglia.

Under this experiment conditions, the commercial _Macleaya cordata_ extract showed no quantitative neuron changes in the myenteric Giemsa+ population and myenteric NADPH-d+ subpopulation, however, the groups that consumed the extract showed greater NADPH-d+ neuron activity compared to the control group, implying that the food remained longer in the intestinal lumen, which can indicate greater nutrient use, resulting in increased performance.

The study of the myenteric Giemsa+ population and NADPH-d+ subpopulation and the effects of the commercial _Macleaya cordata_ extract on these neurons improve knowledge on the functionality of the ENS in digestive processes, serving as a reference for future research in the poultry industry.
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BIOETHICS AND BIOSECURITY COMMITTEE APPROVAL

This research was conducted in accordance with the norms edited by the National Council for the Control of Animal Experimentation (CONCEA) and was approved by the CARE of UNIPAR, under protocol No. 31660/2017.

DECLARATION OF CONFLICT OF INTEREST

The authors declare no conflicts of interest. This article is part of the PhD dissertation of the first author in the Graduate Program in Animal Science with Emphasis on Bioactive Products at UNIPAR, Umuarama, Paraná, Brazil.

AUTHORS’ CONTRIBUTIONS

All authors contributed equally for the conception and writing of the manuscript. All authors critically revised the manuscript and approved of the final.

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