Ethyl Acetate Fraction from *Trichilia catigua* Confers Partial Neuroprotection in Components of the Enteric Innervation of the Jejunum in Diabetic Rats

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**Key Words**
*Trichilia catigua* • Enteric Neuron • Enteric Glia • Diabetes mellitus • Antioxidant action • Neuroprotection

**Abstract**

**Background/Aims:** Diabetes causes damage to the enteric nervous system. The enteric nervous system consists of neurons and enteric glial cells (EGCs). The present study evaluated the effects of an ethyl-acetate fraction (EAF) from *Trichilia catigua* (*T. catigua*; 200 mg/kg) on the total population of enteric neurons (HuC/D-immunoreactive [IR]) and EGCs (S100-IR and glial fibrillary acidic protein [GFAP]-IR) in the total preparation and jejunal mucosa in diabetic rats. **Methods:** The animals were distributed into four groups: normoglycemic rats (N), diabetic rats (D), normoglycemic rats that received the EAF (NC), and diabetic rats that received the EAF (DC). The jejunum was processed for immunohistochemistry to evaluate HuC/D, S100, and GFAP immunoreactivity. The expression of S100 and GFAP proteins was also quantified by Western blot. **Results:** The D group exhibited a decrease in the number of neurons and EGCs, an increase in the area of cell bodies, an increase in S100 protein expression, a decrease in GFAP protein expression, and a decrease in S100-IR and GFAP-IR EGCs in the jejunal mucosa. The DC group exhibited a decrease in the number of neurons and EGCs, a decrease in the area of cell bodies, a decrease in S100 and GFAP protein expression, and a decrease in S100-IR and GFAP-IR EGCs in the jejunal mucosa. The NC group exhibited maintenance of the number of neurons and EGCs, an increase in the area of cell bodies, and a decrease in S100 and GFAP protein expression. **Conclusion:** The EAF from *T. catigua* partially conferred protection against diabetic neuropathy in the enteric nervous system.
Introduction

Diabetic neuropathy is the most common complication of diabetes. Approximately half of diabetic patients present symptoms of this disorder [1]. Many mechanisms have been proposed to explain neuronal dysfunction and degeneration in diabetic nerves, including impairments in fatty acid metabolism [2], the production of advanced glycation end products [3], a reduction of nerve blood flow [4], activation of the polyol pathway [5], and deficiencies in neurotrophic substances [6]. The result of all of these factors is the overproduction of reactive oxygen species (ROS). When ROS production exceeds antioxidant capacity, oxidative stress occurs. If cellular antioxidants do not scavenge free radicals, then such radicals can attack and damage proteins, lipids, and nucleic acids in the cell. When this occurs, biological activity decreases, leading to losses of energy metabolism, cell signaling, transport, and other principal functions. These altered products are also targeted for proteasome degradation, further decreasing cellular function. The accumulation of such injury can ultimately cause a cell to die through necrotic or apoptotic mechanisms [7].

Patients with diabetes mellitus (DM) present disorders of the gastrointestinal tract, such as esophageal enteropathy, gastroparesis, constipation, diarrhea, and fecal incontinence [1]. These symptoms are related to damage to the enteric nervous system (ENS). The ENS consists of enteric neurons and enteric glial cells (EGCs) [8]. Enteric glial cells were previously viewed as only a mechanical support structure for neurons. More recently, however, they have been shown to have other important functions to maintain intestinal homeostasis. Enteric glial cells have been shown to play an essential role in regulating the intestinal barrier and influencing motility and inflammatory processes [9-12]. Some studies have shown that both EGCs and enteric neurons are affected by diabetes [11, 13, 14]. Many studies have sought to verify the action of antioxidants (e.g., quercetin, vitamin E, vitamin C, L-glutamine, and L-glutathione) in diabetic neuropathies [14-18]. The present study evaluated the effects of an ethyl-acetate fraction (EAF) from *Trichilia catigua*, which has been reported to exert better antioxidant effects than the crude extract [19]. Previous studies have investigated the antioxidant activity of natural products for the treatment of DM [20-22]. A few studies have evaluated the antioxidant effects of natural products on diabetic neuropathy in the ENS. Thus, the objective of the present study was to evaluate the effects of the EAF from *T. catigua* on the total population of enteric neurons (HuC/D-immunoreactive [IR]) and EGCs (S100-IR and glial fibrillary acidic protein [GFAP]-IR) in the jejunum in diabetic rats.

Materials and Methods

All of the experimental procedures were conducted according to the ethical principles of the Brazilian Society of Animal Laboratory Science (SBCAL/COBEA). These procedures were approved by the Committee of Ethics in Animal Experimentation of the Universidade Estadual de Maringá (protocol no. 082/2012).

Plant material and extract preparation

The crude extract was prepared from the powdered bark of *T. catigua* (HUEM exsiccate no. 1943) as described by Dos Santos et al. [23] and Gomes et al. [24]. The crude extract (50 g) was dissolved in water (500 ml) and extracted with ethyl acetate (500 ml). Afterward, both fractions were concentrated in vacuo and lyophilized, yielding the EAF (8.2 g) and aqueous fraction according to Chassot et al. [19] and Longhini et al. [25]. The EAF was prepared and supplied by PALAFITO (Laboratory of Pharmaceutical Biology) under the supervision of Prof. Dr. João Carlos Palazzo de Mello.

The EAF was evaluated by measurements of procyanidin B2 and epicatechin by high-performance liquid chromatography (HPLC) according to Longhini et al. [25]. The EAF (5 mg) was weighed and dissolved in 1 ml of 20% aqueous methanol solution. This solution was eluted in a solid-phase extraction (SPE; Strata C18-E, Phenomenex, 200 mg/3 ml), preconditioned with methanol and water. A 20% aqueous methanolic solution was added to a volumetric flask to achieve a final volume of 10 ml.
The sample was analyzed by HPLC under the following conditions: Thermo device equipped with a PDA spectrophotometer detector (Finnigan Surveyor PDA Plus Detector), integral pumps and degasser (Finnigan Surveyor LC Pump Plus), automatic sampler (Finnigan Surveyor Autosampler Plus) equipped with a 10 μl loop and software controller (Chromquest), Phenomenex 100 Å column (250 x 4.6 mm, 5 μm) and precolumn (6 x 3 mm, 5 μm, Phenomenex). The EAF that was used in the present study was from the same lot as the EAF that was used by Gomes et al. [24]. The procyanidin B2 and epicatechin concentrations were 74.4 and 40.6 mg/g EAF, respectively.

**Material collection and processing**

Twenty-four male Wistar rats (Central Vivarium of Universidade Estadual de Maringá) were randomly distributed into four groups: normoglycemic rats (N group), diabetic rats (D group), normoglycemic rats that received the EAF (NC group), and diabetic rats that received the EAF (DC group). The rats were housed in individual cages for 120 days under a 12 h/12 h light/dark cycle and controlled temperature (24°C ± 2°C). Food and water were available ad libitum.

The animals in the D and DC groups underwent type 1 DM induction after a 14 h fast. Streptozotocin was administered intravenously (35 mg/kg body weight; Sigma, St. Louis, MO, USA) under anesthesia with thiopental (40 mg/kg body weight, i.p., Abbott Laboratories, Chicago, IL, USA). To verify the establishment of the experimental model of DM, 2 days after induction, one drop of blood was obtained from the tail of the animals to measure glucose photometrically based on glycoso-dye-oxidoreductase (Accu-Chek Active glycosimeter; Roche Diagnostics GmbH, Mannheim, Germany). Only rats with glycemia > 200 mg/dl were used in the subsequent experiments [16-18].

The EAF was administered daily in the NC and DC groups by gavage at a dose of 200 mg/kg [19] for 58 days. The animals in the N and D groups underwent the same administration schedule but received only the diluent.

At 150 days of age, the animals were weighed and received an intravenous injection of vincristine sulfate (0.5 mg/kg; Eurofarma Laboratórios, São Paulo, SP, Brazil) 2 h before euthanasia for microtubule stabilization. The rats were anesthetized with thiopental (40 mg/kg body weight, i.p., Abbott Laboratories, Chicago, IL, USA). Blood was collected by cardiac puncture for blood glucose measurements using the glucose-oxidase method to confirm the diabetes. Laparotomy was performed, and the jejunum was collected.

For immunohistochemistry, whole mounts of the jejunum were washed in 0.1 M phosphate-buffered saline (0.1 M PBS; pH 7.4), fixed for 18 h in Zamboni fixative solution [26], and stored at 4°C. After fixation, the segments were opened along the mesenteric border and successively washed in 80% alcohol until complete removal of the fixative. Subsequent dehydration was performed with an ascending series of alcohol (95%, 80%, and 50%). The samples were then stored in 0.1 M PBS (pH 7.4) with 0.08% sodium azide at 4°C.

For the analysis of the mucosal layer, others samples were used to perform cryostat cuts. After fixation for 18 h in Zamboni fixative [26], the samples underwent successive washes in 0.1 M PBS (pH 7.4) for 12 h, followed by cryoprotection in 18% sucrose solution in 0.1 M PBS (pH 7.4) for 24 h. After cryoprotection, the samples were soaking in O.C.T 4583 compound (Cryomatrix Shandon, Pittsburgh, PA, USA), rapidly frozen in liquid nitrogen, and stored at -80°C.

For Western blot, separate jejunum samples were collected and washed several times with Krebs-Ringer buffer solution (pH 7.4) to completely remove the contents. Tissue lysis was performed in Turrax by immersion in homogenization buffer (50 mM Tris HCl [pH 7.2], 600 mM NaCl, and 1 mM EDTA) that contained protease inhibitor (Sigma-Aldrich, St. Louis, MO, USA). The lysed samples were then centrifuged for 10 min at 10,000 x g to remove the insoluble portion. The supernatant was collected and stored at -80°C until use. The concentration of total proteins in the supernatant was determined by the Bradford method (Bio-Rad, Hercules, CA, USA).

**Immunohistochemical double-staining for HuC/D and S100 proteins**

The jejunum was cut into pieces (~1 cm) that were then micro dissected to obtain whole-mount preparations to evaluate neurons in the myenteric plexus. To immunolabel HuC/D and S100 proteins, the tissue samples were first washed with 0.1 M PBS (pH 7.4) with 0.5% Triton (Sigma-Aldrich, St. Louis, MO, USA) twice for 10 min. Following the washes, blockade was performed with 0.1 M PBS (pH 7.4) and 0.5% Triton X-100 with 2% bovine serum albumin (BSA; Sigma-Aldrich, St. Louis, MO, USA) and 10% goat serum
for 1 h. To double-label HuC/D and S100, the whole mounts were incubated in a solution that contained both primary antibodies: mouse anti-HuC/D antibody (1:400 dilution; Molecular Probes, Eugene, OR, USA) and rabbit anti-S100 antibody (1:200 dilution; Sigma, St. Louis, MO, USA). After 72 h of incubation at room temperature, the jejunum was washed three times in 0.1 M PBS (pH 7.4) that contained 0.5% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) for 5 min each and incubated for 4 h at room temperature with the following secondary antibodies: anti-mouse Alexa Fluor 488 (1:200 dilution; Molecular Probes, Eugene, OR, USA) and anti-rabbit Alexa Fluor 568 (1:200 dilution; Molecular Probes, Eugene, OR, USA). The whole mounts were washed three times in 0.1 M PBS (pH 7.4) for 5 min each, mounted on slides with Prolong Gold Anti-fade mounting medium conjugated to DAPI (Molecular Probes, Eugene, OR, USA), and stored at 4°C. For the negative control, the primary antibody was omitted. To avoid potential procedural variation, tissues from all groups of animals were stained at the same time using the same solutions.

Quantitative analysis

Quantification was performed by sampling in the intermediate jejunum region in images that were captured with microscopy with filters for immunofluorescence (Zeiss, Jena, Germany) and a high-resolution camera (Zeiss, Jena, Germany) coupled to a computer and later recorded on a flash drive. For each technique, 30 images/animal were captured with a 20x objective. All neurons and EGCs were quantified in each image. Quantification was performed using ImagePro Plus 4.5.0.29 image analysis software (Media Cybernetics, Silver Spring, MD, USA). The results are expressed as neurons and EGCs per cm².

Morphometric analysis

Morphometric analyses of the total population of neurons (HuC/D-IR) and EGCs (S100-IR) were performed using samples of the myenteric plexus of the jejunum. The areas (in μm²) of 100 neuronal cell bodies and 100 EGCs were calculated per animal using ImagePro Plus 4.5.0.29 image analysis software (Media Cybernetics, Silver Spring, MD, USA).

Immunohistochemical analysis of GFAP and S100 proteins in the mucosal layer

The samples were semi-serially sectioned at 10 μm using a cryostat, stretched on slides that were prepared with 2% organoacane adhesive in acetone, and stored at -18°C. The slides were washed three times with a solution that contained 0.5% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) in 0.1 M PBS (pH 7.4) for 5 min and blocked in a solution that contained 1% BSA (Sigma-Aldrich, St. Louis, MO, USA) and 0.5% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) in 0.1 M PBS (pH 7.4) for 2 h at room temperature. The slides were then sequentially incubated in a humid chamber in a solution that contained 1% BSA in 0.1 M PBS (pH 7.4) and 0.5% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) with the following primary antibodies for 72 h at 4°C: mouse anti-GFAP (1:200 dilution; Sigma-Aldrich, St. Louis, MO, USA) and rabbit anti-S100 (1:300 dilution; Sigma, St Louis, MO, USA). After incubation, the sections were washed three times with a solution that contained 0.5% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) in 0.1 M PBS (pH 7.4) for 5 min each. The sections were then incubated in a dark wet chamber in 2% BSA (Sigma-Aldrich, St. Louis, MO, USA) in 0.1 M PBS (pH 7.4) for 2 h at 4°C. For slides that contained anti-GFAP primary antibody, Alexa Fluor 488 goat anti-mouse secondary antibody (1:200 dilution; Molecular Probes, Eugene, OR, USA) was added to the solution. For slides that contained anti-S100 primary antibody, Alexa Fluor 568 goat anti-rabbit secondary antibody (1:300 dilution; Molecular Probes, Eugene, OR, USA) was added to the solution. The sections were then washed in a solution that contained 0.5% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) in 0.1 M PBS (pH 7.4). The sections were mounted on slides with Prolong Gold Anti-fade mounting medium conjugated to DAPI (Molecular Probes, Eugene, OR, USA) and stored at 4°C. For the negative control, the primary antibody was omitted. To avoid potential procedural variation, tissues from all groups of animals were stained at the same time using the same solutions.

Quantitative analysis

Quantification was performed using images that were captured with a microscope with filters for immunofluorescence and a high-resolution camera (Zeiss, Jena, Germany) coupled to a computer and later recorded using a flash drive. For each technique, 30 villi/animal were captured with a 20x objective and quantified. Quantification was performed using ImagePro Plus 4.5.0.29 image analysis software (Media Cybernetics, Silver Spring, MD, USA).
Western blot

Protein (30 µg/well) was separated using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (17% for S100 protein and 10% for GFAP protein) and transferred to nitrocellulose membranes (Bio-Rad, Hercules, CA, USA). Transfer efficiency was verified by Ponceau-S staining (Sigma-Aldrich). Proteins were blocked for 1 h at room temperature with a solution that contained 5% nonfat dried milk and 0.1% Tween-20 in Tris-buffered saline (TBS) buffer (2.24 g/L Tris base and 8 g/L NaCl, pH 7.6). The membranes were then incubated overnight at 4°C under stirring with rabbit anti-S100 primary antibody (1:500 dilution; Sigma, St Louis, MO, USA) and mouse anti-GFAP primary antibody (1:500 dilution; Sigma-Aldrich, St. Louis, MO, USA). The membranes were then incubated for 2 h at room temperature with horseradish peroxidase-conjugated anti-rabbit secondary antibody (1:1000 dilution; Novex, Invitrogen Rockford, IL, USA) and horseradish peroxidase-conjugated anti-mouse secondary antibody (1:2000 dilution; Novex, Invitrogen Rockford, IL, USA). Immunodetection was performed using the chemiluminescence method with a ChemiDoc MP imager according to the manufacturer’s instructions (V3 Western Workflow, Bio-Rad, Hercules, CA, USA). Immunostaining was analyzed using Image Lab software (Bio-Rad, Hercules, CA, USA). Protein levels were normalized to β-actin as the control. All of the electrophoresis and Western blot protocols were performed according to the standard Bio-Rad Mini Protean System procedures.

Statistical analysis

The statistical analysis was performed using Statistics 7.0 software (Statsoft South America, Sao Caetano do Sul, SP, Brazil) and Prism 3.1 software (GraphPad, La Jolla, CA, USA). The results are expressed as mean ± standard error of the mean. The morphometric data were subjected to delineation blocks followed by Fisher’s test. The other data were analyzed using one-way analysis of variance (ANOVA) followed by Fisher’s post hoc test. Values of p < 0.05 were considered statistically significant.

Results

Sixty days after diabetes induction, the animals in the D and DC groups were determined to be hyperglycemic. Animals in the diabetic groups exhibited the loss of body weight compared with nondiabetic rats in the N and NC groups (Table 1).

The quantitative analysis revealed a reduction of the number of HuC/D-IR myenteric neurons (p < 0.0005) and S100-IR EGCs (p < 0.0001) in the D group compared with the N group. The administration of EAF in the DC group did not alter the reduction of the density of myenteric neurons or EGCs compared with the D group (p > 0.05). The NC group exhibited preservation of the number of HuC/D-IR neurons (p < 0.05) and S100-IR EGCs (p < 0.03) compared with the N group (Fig. 1).

The cell body area of HuC/D-IR neurons and S100-IR EGCs in diabetic animals increased by 24% and 39%, respectively, in the D group compared with the N group (p < 0.00001; Fig. 2). The EAF prevented the increase in the area of cell bodies of HuC/D-IR neurons and the area of cell bodies of S100-IR EGCs in diabetic animals (16% and 12% respectively) in the DC group compared with the D group (p < 0.00001; Fig. 2). Animals in the NC group exhibited a 13% increase in the area of HuC/D-IR neurons and 33% increase in the area of S100-IR EGCs compared with the N group (p < 0.00001; Fig. 2).

S100 protein expression increased in diabetic animals compared with the N group (p < 0.03; Fig. 3A). Treatment with the EAF in the DC group reduced S100 protein expression compared with the D group (p < 0.01; Fig. 3) but not compared with the N group (p > 0.05; Fig. 3). The expression of the Table 1. Physiological parameters, including initial weight (90 days old), final weight (150 days old), and final glycemia in the following groups: normoglycemic (N), diabetic (D), normoglycemic that received EAF (NC), and diabetic that received EAF (DC). The results are expressed as mean ± SEM, n = 6 rats/group. *p < 0.05, compared with N group.

<table>
<thead>
<tr>
<th>Physiological parameter</th>
<th>N</th>
<th>NC</th>
<th>D</th>
<th>DC</th>
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<tbody>
<tr>
<td>Initial weight (g)</td>
<td>353.7 ± 8.62</td>
<td>358.3 ± 13.04</td>
<td>341.8 ± 8.18</td>
<td>362.0 ± 9.90</td>
</tr>
<tr>
<td>Final weight (g)</td>
<td>455.3 ± 18.38</td>
<td>435.3 ± 16.80</td>
<td>326.8 ± 1.74*</td>
<td>323.4 ± 19.98*</td>
</tr>
<tr>
<td>Final glycemia (mg/dL)</td>
<td>121.2 ± 12.23</td>
<td>104.5 ± 6.26</td>
<td>565.8 ± 23.12*</td>
<td>466.4 ± 37.38*</td>
</tr>
</tbody>
</table>
GFAP protein in the NC, D, and DC groups decreased compared with the N group (p > 0.0003; Fig. 3B).

The density of GFAP-IR EGCs in the jejunal mucosa decreased in the NC, D, and DC groups compared with the N group (p < 0.05; Fig. 4A, C). The number of GFAP-IR EGCs was similar in the D and DC groups (p > 0.05; Fig. 4A). The density of S100-IR EGCs in the jejunal mucosa decreased in the D group compared with the N group (p < 0.05; Fig. 4B, C). The density of S100-IR EGCs decreased in the D and DC groups compared with the NC group (p < 0.05; Fig. 4B, C), with no difference between the D and DC groups (p > 0.05; Fig. 4B).

Discussion

The present study established a model of DM. The animals in the D and DC groups had a glycemic index > 400 mg/dl and also presented significant weight loss at the end of the experiment. The results of this study were similar to previous studies that were performed by our research group [14, 16, 18].

Fig. 1. (Top) Density per unit area (cm²) of HuC/D-immunoreactive myenteric neurons and S-100-immunoreactive enteric glial cells. The results are expressed as mean ± SEM. *p<0.05, compared with N group. (Bottom) Photomicrographs of HuC/D-immunoreactive myenteric neurons and S-100-immunoreactive enteric glial cells in the jejunum in the N, NC, D, and DC groups. Scale bar = 50 μm. n = 6 rats/group.

Fig. 2. (A, B) Cell body area (μm²) of HuC/D-immunoreactive myenteric neurons (A) and S-100-immunoreactive enteric glial cells (B) in the jejunum in the N, NC, D, and DC groups. The results are expressed as the mean ± SEM. n = 6 rats/group. *p<0.0001, compared with the N group; **p<0.0001, compared with the D group.
The analysis of neuronal density showed an 18% reduction of HuC/D-IR neurons in the D group compared with the N group. This decrease may be attributable to a reduction of antioxidant defense concomitant with the intensification of cellular oxidative stress that resulted from diabetes. According to Kuyvenhoven [27], free radicals can react with DNA, proteins, and lipids, causing neuronal death. Neuronal death that is caused by diabetes-related oxidative stress is considered a main causal factor for ENS damage [28].
In addition to the reduction of HuC/D-IR neurons, we observed a reduction of S100-IR EGCs in the D group compared with the N group. A 26% reduction of S100 immunoreactivity was observed in the total preparation, with a 16% reduction of S100 immunoreactivity in the intestinal mucosa. The literature reports various reductions of EGCs that are caused by aging [29] and diet [30]. The reduction of glial cell density in diabetes may also be related to an elevation of oxidative stress. Lopes et al. [13], Qi et al. [11], and Souza et al. [14] reported decreases in glial cell density in the duodenum, stomach, and jejunum, respectively, in diabetic rats. However, other studies by our research group reported different results with regard to glial cell density. This discrepancy may be attributable to the segment that was evaluated, such as the ileum [18, 31] and cecum [32]. The impact of diabetes on EGCs varies according to the gastrointestinal segment. Reductions of EGCs and neuronal density are reflected by gastrointestinal tract physiology. Enteric glial cells represent a communication link between the ENS and local immune system, thus playing a role in the normal function of the digestive system [10]. Previous studies suggested that EGCs are the main regulators of intestinal epithelial barrier (IEB) function [33]. The loss of IEB integrity allows the translocation of normally excluded luminal contents (e.g., microorganisms, antigens, and nutrients, among others) to the mucosa, thus initiating and perpetuating inflammatory disorders and tissue injury [34]. The IEB serves as the first defensive boundary between the organism and luminal environment [35]. The reduction of EGCs could be one cause of diarrhea and intestinal constipation, in addition to the low immunity that diabetic patients present.

A previous study evaluated the importance of EGCs in intestinal physiology, showing that all adult rats died within 2 weeks after GFAP-IR EGC depletion because of fulminating jejunileitis [36]. These findings support the role of EGCs in maintaining intestinal integrity. In the present study, we observed a 20% reduction of GFAP-IR glial cell density in the intestinal mucosa in the D group compared with the N group. This may have also been caused by an increase in oxidative stress. The reduction of the density of GFAP-IR EGCs was accompanied by lower GFAP protein expression, evaluated by Western blot. GFAP expression is modulated by glial cells under conditions of differentiation, inflammation, and injury, indicating that the level of GFAP production coincides with the functional status of these cells [10].

The reductions of the density of HuC/D-IR neurons and S100-IR EGCs in the D group were accompanied by increases in the cell body area and S100-IR EGCs, with higher S100 protein expression, evaluated by Western blot. This increase in neuronal area may be attributable to an increase in neurotransmitter synthesis in an attempt to compensate for neuronal death [13]. This increase may also suggest metabolic neural alterations that are caused by diabetes through an increase in sorbitol production, which increases the osmolarity of neurons and causes neuronal edema [7]. The increase in EGCs may reflect cellular edema that is caused by the same metabolic alterations of neurons or a mechanism that compensates for the loss of neurons and seeks to maintain homeostasis. Enteric glial cells provide mechanical support for enteric neurons. They release a wide range of factors that are responsible for the development, survival, and differentiation of peripheral neurons. Enteric glial cells have also been shown to have a more complex nature, playing a role in the maintenance of intestinal homeostasis [12].

Diabetic neuropathy is related to an elevation of ROS. Antioxidants have been used to prevent the development of diabetic neuropathy. The present study used an EAF from *T. catigua* as an antioxidant agent [37]. Longhini et al. [25] found that an EAF from *T. catigua* contained two flavonoids, procyanidin B2 and epicatechin, which have good antioxidant activity [38]. Administration of the EAF did not alter S100-IR or GFAP-IR EGCs or neuronal density in the DC group, but it prevented the increase in neuronal area of S100-IR EGCs in the DC group. No increase in the area of S100-IR EGCs was found, but a reduction of the expression of the S100 protein was observed, quantified by Western blot. This lack of an increase in cellular area may be attributable to the antioxidant action of the EAF. The antioxidant activity of flavonoids occurs through different mechanisms, such as the elimination of free radicals,
metal ion chelating agents (e.g., iron and copper), and the inhibition of enzymes that are responsible for the generation of free radicals [39].

Preservation of the densities of S100-IR EGCs and HuC/D-IR neurons were observed in the NC group. The aging process causes ROS generation but to a lesser extent than diabetic individuals that present the activation of different metabolic cascades that converge to produce higher ROS production. In the NC group, the antioxidant action of the EAF protected neurons and EGCs from the deleterious actions of ROS that were generated by the aging process, preserving cell density. Notwithstanding the beneficial effect of the EAF, it altered the cell areas of both neurons and S100-IR EGCs, the density of GFAP-IRs EGCs, and GFAP protein expression. These alterations may indicate that the EAF altered the osmolarity of the cell, causing cellular edema and reducing the density of GFAP-IR cells.

Conclusion

The effects of additional doses of the EAF on diabetic neuropathy should be evaluated because many treatment responses are dose-dependent. In conclusion, the effects of the EAF from T. catigua were partially beneficial for the prevention of diabetic neuropathy in the ENS.

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Cynthia Priscilla do Nascimento Bonato Panizzon performed the experiments, did the statistical analysis and wrote the manuscript. Marcílio Hubner de Miranda Neto, revised the manuscript, Francielle Veiga Ramalho, performed the experiments, Renata Longhini, was in charge of producing the EAF, João Carlos Palazzo de Mello was in charge of producing the EAF and Jacqueline Nelisis Zanoni designed the study, wrote the manuscript and revised the manuscript.

Disclosure Statement

The authors have no competing interests.

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