ABSTRACT - This study had as its purpose to assess the effects of acute diabetes induced by streptozotocin (35 mg/kg body weight) on the number and size of the myenteric neurons of the duodenum of adult rats considering equally the antimesenteric and intermediate regions of the intestinal circumference. Experimental period extended for a week. Neuronal counts were carried out on the same number of fields of both regions of the duodenal circumference and measurements of neuronal and nuclear areas on equal numbers of cells. Number and size of the myenteric neurons stained with Giemsa were not significantly different between groups. On the other hand, the proportion of NADH-positive neurons increased from 18.54% on the controls to 39.33% on the diabetics. The authors discuss that this increased reactivity probably results from a greater NADH/NAD+ ratio, described in many tissues of diabetic animals, which has consequences on the modulation of the enzymes that use these cofactors and whose activity is detected by the NADH-diaphorase technique.

KEY WORDS: duodenum, myenteric neurons, acute diabetes, NADH-diaphorase.

Número e tamanho dos neurônios mientéricos do duodeno de ratos adultos com diabetes agudo

RESUMO - Este estudo teve como objetivo avaliar os efeitos do diabetes agudo induzido por estreptozootocina (35 mg/kg de peso corporal) sobre o número e tamanho dos neurônios mientéricos do duodeno de ratos adultos considerando de forma equivalente as regiões antimesentérica e intermediária da circunferência intestinal. O período experimental se estendeu por uma semana. As contagens neuronais foram feitas em igual número de campos nas duas regiões da circunferência duodenal e as mensurações das áreas neuronais e nucleares em igual número de células. O número e o tamanho dos neurônios corados por Giemsa não foram significativamente diferentes entre os grupos. Por outro lado, a proporção de neurônios NADH-positivos aumentou de 18,54% nos animais controles para 39,33% nos diabéticos. Os autores discutem que essa maior reatividade possivelmente resultou do aumento da proporção NADH/NAD+, descrita em diversos tecidos de animais diabéticos, que repercute na modulação das enzimas que utilizam esses cofatores e cuja atividade é detectada pela técnica da NADH-diaforase.

PALAVRAS-CHAVE: duodeno, neurônios mientéricos, diabetes agudo, NADH-diaforase.

The knowledge about the mammalian enteric nervous system (ENS) can be considered, currently, as wide and in some instances, quite deep. Descriptions found on the literature assess several aspects of this subdivision of the autonomic nervous system and reveal the richness of structure and organization of this neuronal population of magnitude similar to that of the spinal
Systematization of the studies on the enteric neurons, especially those of the myenteric plexus of species such as the rat, mouse and guinea-pig, have permitted to determine, with relative precision, the functions of specific groups of neurons. Due to the accessibility of the ENS, investigations about the alterations caused by pathophysiological states, such as diabetes, have been equally productive; there are reports on the literature about the differential effects of diabetes on specific neurotransmitter systems and on neurons of diverse intestinal segments.

Among the various approaches used on the study of the neurons of the ENS one of the most common and of good reproducibility is the morphoquantitative approach, i.e., the evaluation of the enteric neurons through neuron counts, measurements and morphological parameters. As these studies proceed, differences are observed between the gastrointestinal segments concerning the number and size of the myenteric neurons. More recently, it was also found that a single portion of the gastrointestinal tract can show significant neuronal differences among its circumferential regions. This fact must be considered of fundamental importance on the morphological and quantitative analyses, especially when experimental states are being assessed, once inherent differences of the enteric neurons can be mistaken for pathophysiological differences.

A manner of avoiding this kind of difficulty is to consider, on the morphoquantitative analyses, a single circumferential region of the intestinal segment of interest in all experimental groups. Alternatively, it is possible to take into account two or more regions of the intestinal circumference, as long as they are equally evaluated.

Considering that there are many reports about morphological and quantitative alterations of the myenteric plexus which are caused by diabetes, but that do not take into account the circumferential regionalization of the studied segment(s), this study analyzed the effects of acute diabetes on the number and size of the myenteric neurons of the duodenum of adult rats considering equally the antimesenteric and intermediate regions of the duodenal circumference.

**METHOD**

Twenty male Rattus norvegicus, aging seven months, from the Central Biotery of the State University of Maringá, were divided into two groups: group C, control, composed of ten animals weighting 443.6±3.06 gr and group D, diabetic, composed of ten animals of 437.4±3.04 gr body weight. All the animals were initially subjected to overnight fast, but only the animals of group D received i.v. injection of streptozotocin (35 mg/kg body weight) in citrate buffer pH 4.5 in the morning. Groups were kept in separate compartments with water and ration ad libitum for one week.

Overnight fast preceeded the sacrifice, so as to facilitate cleaning of the segments. Animals were killed under ethylc ether anesthesia, blood samples were collected for the glucose oxidase test, and laparotomy was carried out.

The duodenum was entirely removed and the excess of mesenteric fat was dissected out. Five segments of each experimental group were washed in 0.9% saline solution, and filled and immersed in fixative for further staining according to the Giemsa technique. Fixation was, at least, of 48 hours, and staining was carried out for 16 to 20 hours. The other five segments of each group were washed and filled with Krebs solution, pH 7.3, and prepared according to the NADH-diaphorase technique. Segments were kept on the incubation medium for 45 minutes, and fixation in 10% buffered formol solution lasted for a minimum of 48 hours. Distension of the segments was kept by ligature of the extremities.

For microscope examination, rings of duodenum were opened by the mesenteric margin and the mucosa and submucosa, as well as part of the circular muscle layer, were removed under stereomicroscope. Slide preparation of the resultant whole-mounts followed the routine histological procedure of dehydration and diaphanization.

Neuron counts were carried out on Olympus BX40 microscope under 40X objective. In each whole-mount preparation, either those stained by Giemsa or those subjected to the NADH-diaphorase technique, it was counted all neurons seen in 80 microscopic fields, 40 fields on the antimesenteric region and 40 on the intermediate region (Fig 1). Half-seen neurons were counted on alternate fields. Total area of the 80 fields was 17.68 mm².
The proportion of neurons stained by the NADH-diaphorase technique (NADH-positive neurons) relative to the Giemsa-stained neurons was calculated.

Measurements of the areas of cell body and nucleus profiles were made using image analyser (Image-Pro Plus 3.0.1) coupled to the microscope. In each Giemsa-stained whole-mount 100 neurons were measured, 50 in each circumferential region.

Data are presented as mean ± SE. The two regions were considered altogether, both on the quantitative and the morphological analyses. Comparisons of the results used the single-tailed t test for unpaired data (Prism 2.0) with significance level of 5%.

RESULTS

Plasma glucose of the control group (102.0±4.51 mg/dl) was significantly lower (p<0.0001) than that of the diabetic animals (214.4±9.15 mg/dl). During removal and dissection of the duodenum, alterations of tonus, hypertrophy or hypotrophy of tunicas or lesions were observed in none of the animals.

In 17.68 mm² of the Giemsa-stained whole-mounts it was found 3615±62.01 neurons on the control animals; on the diabetic animals 3460±123.9 neurons were counted. The difference between groups did not attain significance (p=0.2957), in spite of the number of neurons found on the diabetic animals being smaller than that found on the controls.

As for the NADH-positive neurons, it was found in 17.68 mm² 670.2±129.4 neurons on the controls (Fig 2) and 1361±181.8 neurons on the diabetics (Fig 3), this being a significantly higher
value (p=0.0148). In proportional terms, the NADH-positive neurons represented 18.54% of the Giemsa-stained neurons on the control animals and 39.33% on the diabetic animals.

The mean area of cell body profile of the myenteric neurons on the duodenum of the control animals was 229.7±3.75 \(\mu\text{m}^2\) and that of the nuclear profile was 88.02±1.23 \(\mu\text{m}^2\). These areas were not significantly different from those of the diabetic animals, where the corresponding values were 224.0±3.88 \(\mu\text{m}^2\) and 86.52±1.23 \(\mu\text{m}^2\) (p=0.2911 for cell body area and p=0.3887 for nuclear area).

**DISCUSSION**

Streptozotocin is being increasingly used in experimental procedures as diabetogenic agent. Its mode of action and its potential are still subjects of research and assays. In this study, it was observed that streptozotocin caused a consistent diabetic state on the animals, although plasma
glucose level (214.4 mg/dl) has not been markedly elevated, contrary to the observations made with
the same dose of the drug applied to rats aging 75 days and kept diabetic for periods of two and eight
months21. Studies about the dose-response relationship of streptozotocin discuss that blood glucose
levels vary during the first weeks after diabetes induction22, which can explain the relatively low
glycemic levels observed in this study.

The number of Giemsa-stained myenteric neurons found on the duodenum of the control
animals is about 38% smaller than that reported by Buttow et al.23, with values corrected for differences
on the counting areas. The explanation is based on the difference of age between the rats: while the
animals of this study had seven months of age, those had three and five months on the sacrifice. This
decrease on the neuronal population is also described by other authors on the intestine of rats15 and
guinea-pigs24, and on the small and large intestines of humans25,26, and is due to aging by itself,
independently of pathophysiological conditions.

In spite of the diabetic animals exhibiting fewer Giemsa-stained neurons than the controls,
the difference between counts did not attain significance. On the other hand, there are reports of
approximately 18% decrease on the number of myenteric neurons in the duodenum of diabetic rats23.
It must be stressed, however, that those authors used the same dose of streptozotocin but in
considerably younger animals (mean weight of 250 gr and 75 days of age). Considering that peripheral
nerves of rats are more susceptible to morphological and physiological changes until the 26th week of
age27, it is possible that the myenteric plexus of younger rats is more sensitive to experimental
diabetes than that of mature ones.

Opposite to what was observed on the Giemsa-stained neurons, the number of NADH-positive
neurons increased substantially on the diabetic animals, rising the proportion of this neuronal sub-
population, relative to the Giemsa-stained population, from 18.54% on the controls to 39.33% on
the diabetics. On the literature, there are reports stating that high glycemic levels28,29 and their metabolic
and vascular consequences30 are the cause of an increased NADH/NAD\(^+\) ratio, which influences the
activity of the enzymes using these cofactors28. In this way, it is possible that the greater neuronal
reactivity observed in this study was due to a greater activity of the enzyme(s) involved on the
evidenciation of these cells by the NADH-diaphorase technique, elicited during the experimental
period by the hyperglycemic state, and not by streptozotocin, once the glucose uptake by neurons is
insulin-independent, and thus is a function of the plasma concentration of the substance. On the
hepatic cells, which are also insulin-independent for glucose uptake, it was observed that in periods
of diabetes from five days to eight months, the relationship NADH/NAD\(^+\) is always changed and
probably irreversibly29. The disponibility of substrates and the reaction time during incubation of
the segments can both be discarded as cause of the greater neuronal reactivity on the diabetic animals,
because incubation medium contained, in equal amounts for both groups, the substrates nitro blue
tetrazolium (NBT) and \(\beta\)-NADH, and all segments were identically processed.

In humans whose plasma glucose levels reach between 300 and 500 mg/dl, the glucose loss on
the urine is considered as causing extra- and intracellular dehydration31. In this study, the morphometric
analyses of the areas of cell body and nucleus profiles of the myenteric neurons did not demonstrate a
statistically significant difference between the control and diabetic groups, although on the latter the
neurons and their nuclei had been marginally smaller. The lack of difference on the morphometry of
these cells can be related to the glycemic level which, despite being above normal, was below that
observed in chronically-diabetic animals21, and thus was not high enough to elicit an intracellular
dehydration capable of affecting the dimensions of the myenteric neurons and their nuclei.

CONCLUSIONS

1) In mature adult rats, streptozotocin-induced diabetes does not provoke significant decrease
on the duodenal neuronal population during the one-week period.
2) On the streptozotocin-induced diabetes, there is an increase on the respiratory metabolism of the myenteric neurons, rising the proportion of neurons evidenced by the NADH-diaphorase.

3) Moderate plasma glucose levels in diabetic rats are not sufficient to induce intracellular dehydration capable of significantly reducing the size of the myenteric neurons and their nuclei.

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