Immunohistochemical study of the digestive tract of Oligosarcus hepsetus

Danielle A Vieira-Lopes, Nadja L Pinheiro, Armando Sales, Adriana Ventura, Francisco G Araújo, Iracema D Gomes, Aparecida A Nascimento

AIM: To describe the histology of the digestive tract and to investigate the occurrence of endocrine cells in Oligosarcus hepsetus (O. hepsetus).

METHODS: The digestive tract (DT) of O. hepsetus was divided into esophagus, two stomach regions (glandular and non-glandular) and two intestinal regions (anterior and posterior). These specimens were processed by routine histological techniques and stained with hematoxylin-eosin, Gomori’s trichrome, periodic acid Schiff (PAS) and Alcian blue (AB). An immunohistochemical method using avidin-biotin-peroxidase was employed.

RESULTS: The esophagus is lined with a non-keratinized stratified squamous epithelium that is reactive to PAS and AB. The stomach has a mucosa lined with a simple columnar epithelium with mucus-secreting cells that are reactive only to PAS. The intestine has a simple columnar epithelium with a brush border and goblet cells that are reactive to PAS and AB. Somatostatin, serotonin and cholecystokinin immunoreactive cells were identified throughout the DT.

CONCLUSION: This study revealed adaptations for the species’ diet and showed that the distribution and relative frequency of immunoreactive cells are similar to those of other fish.

INTRODUCTION

The literature stresses the importance of knowledge of the anatomy of the digestive tract (DT) of fishes, because this structure is highly variable, related to the diversity of feeding habits, type of food and lifestyles. The histological architecture of the DT includes a layer of mucus-secreting cells, observed by histochemical techniques in various studies of teleosts. The secretions vary among different fish species and also according to the location in the DT within the same species. These secretions play an important role in lubricating the or-
Digestive tract of *Oligosarcus hepsetus* 

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The specimens were collected from two reservoirs: Funil (22°30’S, 44°45’W), and Ribeirão das Lajes (22°43’S, 44°23’W), all located in Rio de Janeiro state, Brazil. 

The histochemical analysis involved periodic acid Schiff (PAS) and Alcian blue (AB) pH 2.5 staining to reveal the neutral and acid glycoconjugates (GCs), respectively. Five slides from each specimen were prepared for each protocol, one from each of the five sectioned regions. 

**Histological and histochemical analysis** 

The sections obtained from the DT of *O. hepsetus* were stained with hematoxylin-eosin for analysis of the tissue architecture and with Gomori’s trichrome for differential visualization of the connective tissue and collagen fibers. 

The histochemical analysis involved periodic acid Schiff (PAS) and Alcian blue (AB) pH 2.5 staining to reveal the neutral and acid glycoconjugates (GCs), respectively. Five slides from each specimen were prepared for each protocol, one from each of the five sectioned regions. 

**Immunohistochemistry** 

For the immunohistochemical procedure, 5 μm-thick sections were cut by microtome and mounted on glass slides precoated with 0.1% poly-L-lysine, after being de-waxed and dehydrated by the routine protocol. They were incubated in citrate buffer (pH 6.0-0.01 M) and placed in a microwave oven for 15 min to recover the antigen, then they were incubated with a solution of 3% H2O2 in methanol for 15 min to block any endogenous peroxidase. Subsequently, the sections were incubated at room temperature in a humid chamber with a 1:100 μL dilution of bovine serum albumin in phosphate buffered saline (PBS) solution for 30 min. The sections were then incubated overnight at 4°C with the primary antisera against the individual gastrointestinal hormones (Table 1). The sections were then incubated with biotinylated “Universal” secondary antibody diluted to 1:200 μL for 30 min at room temperature, then with avidin-biotin-peroxidase complex, and to investigate immunohistochemically the Ilha dos Pombos and Santa Cecília sites of the Paraíba do Sul River, and three each from the Ilha dos Pombos and Santa Cecília sites of the Paraíba do Sul River. The fish were dissected in the field, after being anesthetized with benzocaine hydrochloride (50 mg/L) and killed rapidly by hypothermia for immediate removal of the DT. Fragments of the DT were fixed for 8 h in Bouin's fluid and then placed in 70% alcohol. The esophagus, two stomach regions (glandular and non-glandular) and two intestinal regions (anterior and posterior) were obtained from each specimen (Figure 1). These materials were taken to the Histology and Embryology Laboratory of the Federal Rural University of Rio de Janeiro, and the DT segments, endocrine cells compose a complex system disseminated among the epithelial components, with the ability to secrete physiologically active polypeptide hormones and amines. According to Deveney *et al.*, hormones have important functions in the overall regulation of the digestive process, such as nutrient absorption, the secretion of intestinal and associated glands, gut motility and intestinal blood flow. 

Table 1. These studies are essential for efforts to restock native species to improve the condition of ecosystems. 

**MATERIALS AND METHODS**

**Collection area** 

The specimens were collected from two reservoirs: Funil (22°30’S, 44°45’W), and Ribeirão das Lajes (22°43’S, 44°23’W), as well as two points of the Paraíba do Sul River: Ilha dos Pombos (21°84’S, 42°58’W) and Santa Cecília (22°48’S, 44°45’W), all located in Rio de Janeiro state, Brazil. 

**Tissue processing** 

The specimens were collected from two reservoirs: Funil (22°30’S, 44°45’W), and Ribeirão das Lajes (22°43’S, 44°23’W), all located in Rio de Janeiro state, Brazil. 

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Table 1  Details of primary antisera used in this study

<table>
<thead>
<tr>
<th>Primary antiserum</th>
<th>Donor</th>
<th>Code No.</th>
<th>Working dilution</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Somatostatin</td>
<td>Rabbit</td>
<td>A366</td>
<td>1:300</td>
<td>Dako Corp., CA, United States</td>
</tr>
<tr>
<td>Serotonin</td>
<td>Rabbit</td>
<td>S5545</td>
<td>1.6.000</td>
<td>Sigma-Aldrich, Inc., United States</td>
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<tr>
<td>Cholecystokinin</td>
<td>Rabbit</td>
<td>C2581</td>
<td>1.8.000</td>
<td>Sigma-Aldrich, Inc., United States</td>
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<tr>
<td>Gastrin</td>
<td>Rabbit</td>
<td>G0785</td>
<td>1.1.000</td>
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<tr>
<td>Glucagon</td>
<td>Mouse</td>
<td>G2654</td>
<td>1.2.000</td>
<td>Sigma-Aldrich, Inc., United States</td>
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<tr>
<td>Insulin</td>
<td>Mouse</td>
<td>I2018</td>
<td>1.1.000</td>
<td>Sigma-Aldrich, Inc., United States</td>
</tr>
</tbody>
</table>

Intensity of immunoreactions: (-), absent; (+), low; (++), medium; (+++), strong. GI: Glandular; NG: Non-glandular; ANT: Anterior; POST: Posterior.

Table 2  Regional distribution and intensity of immunoreaction in endocrine cells in the digestive tract of Oligosarcus hepsetus

<table>
<thead>
<tr>
<th>Antisera</th>
<th>Segment of the esophagus</th>
<th>Segments of the stomach</th>
<th>Segments of the gut</th>
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<tr>
<td></td>
<td>GI</td>
<td>NG</td>
<td>ANT</td>
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<tr>
<td>Somatostatin</td>
<td>+++</td>
<td>++</td>
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<td>Serotonin</td>
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<td>Cholecystokinin</td>
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<tr>
<td>Glucagon</td>
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<td>Gastrin</td>
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<td>Insulin</td>
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</tr>
</tbody>
</table>

diluted at 1:200 μL for 30 min at room temperature. Subsequently, the peroxidase label was revealed by reaction with Stable DAB/Plus, prepared according to the kit’s instructions. All dilutions and thorough washes between stages were performed using PBS (pH 7.4). The sections were counterstained with Harris hematoxylin, rinsed with deionized water, dehydrated through a series of ethanol and methylcyclohexane solutions and mounted using Entellan. To investigate the specificity of the reactions, negative and positive controls were used. The negative control was prepared by replacement of the primary antibody with non-immune serum and PBS (pH 7.4). Positive controls were produced using tissue sections for each respective antisera, as indicated in the product data sheet.

Observation and photomicrography
Photomicrographs of all samples from each of the fourteen specimens were obtained with a digital camera Nikon Coolpix 4300 attached to a microscope Olympus BX41. The number of immunoreactive endocrine cells to each antisem per analyzed segment was recorded and the intensity of immunoreaction was classified: absent (-) or low (+), medium (+++) and strong (++++) immunoreactivity (Table 2).

RESULTS
Histological and histochemical study of digestive tract
The following layers were observed in the DT of O. hepsetus: mucosa, submucosa, muscular and adventitia or serosa. The muscularis mucosae is absent in this species.

Esophagus
Histological examination revealed that the mucosa of the esophagus of O. hepsetus has many longitudinal folds and is lined with a stratified epithelium with non-keratinized squamous surface cells. The majority of the mucous-secreting cells are interspersed with a smaller number of non-secretory cells (Figure 2A).

The secretory cells reacted positively to PAS and AB staining, indicating the presence of neutral and acid GCs, respectively. The lamina propria is composed of connective tissue and does not have glands. The muscular layer is formed by two sub-layers of striated skeletal muscle, with an internal longitudinal and an external circular layer. Externally, the esophagus is enveloped by an adventitia, composed of connective tissue with some nerve fibers and blood vessels (Figure 2).

Stomach
In the stomach of O. hepsetus, the mucosa is lined with a simple epithelium layer composed of columnar mucous-secreting cells with basal nuclei. These were reactive to PAS but not to AB, revealing the presence of only neutral GCs. The stomach epithelium forms crypts along the gastric mucosa. The mucosa layer projects toward the organ’s lumen, forming various gastric folds, arranged longitudinally. In the non-glandular region, the submucosa and muscular layers accompany the mucosa in forming these folds, making the lumen very small (Figure 3).

The division of the stomach observed in the present work is in accordance with the structural characteristics of the two regions. The glandular region is characterized by having well-developed tubular gastric glands, composed of oxynticopetic cells, occupying the entire lamina propria (Figure 3B). These are smaller and less numerous in the initial portion and increase in number and size in the direction of the non-glandular region. The non-glandular region has a well-developed muscular layer (Figure 3D and E). The submucosa layer is composed of connective tissue and blood vessels.

In the stomach, the muscular layer is composed of smooth muscle fibers arranged in two directions: internal circular and external longitudinal (Figure 3E). The glandular region contains myenteric plexuses arranged in sparse groups, composing the enteric nervous system and located between the muscular sub-layers, and there is a serosa layer which surrounds these structures.

Intestine
Just as for the stomach, the adopted division of the intestine follows specific structural patterns in relation to the mucosa layer. The histological analysis of the intestine of O. hepsetus revealed that the pattern of folds varies, characterizing two distinct regions: anterior and posterior. The anterior region has numerous thin and elongated folds with villi (Figure 4A). In contrast, in the posterior
The intestinal mucosa is lined by a simple columnar epithelium with a striated border and goblet cells, which were positive to PAS and AB stainings, with pink (PAS) and blue (AB) coloration (Figure 4), indicating the presence of neutral and acid GCs, respectively.

The limits between the lamina propria and the submucosa layer are not evident, with connective tissue and blood vessels present in both these regions. In both parts of the intestine, the muscular layer has the same organization as in the stomach, with an internal circular layer and an external longitudinal one, observed in the cross-sections, both composed of smooth muscle cells (Figure 4A). The posterior part of the intestine contains a continuous layer of nervous tissue and a myenteric plexus between the muscular sub-layers (Figure 4E). Externally there is a serosa layer.

**Immunohistochemical study of digestive tract**

SOM-, 5-HT- and CCK-immunoreactive (IR) cells were identified in the DT of *O. hepsetus* but GAS-, GLUC- and INS-IR cells were not present (Table 2).

**Esophagus**

Somatostatin immunoreactivity: In the esophagus, SOM-IR cells were detected in the basal layer of the stratified squamous epithelium (Figure 5). Morphologically, these cells were totally colored by chromogen, making it impossible to visualize a nuclear halo. The nucleus of these cells is very small, occupying a tiny area inside them.

**Stomach**

Serotonin and somatostatin immunoreactivity: Serotonin (5-HT)-IR cells (Figure 6) and SOM-IR cells (Figure 7) were observed in the lining epithelium and glandular epithelium of the stomach. Regarding the morphology of immunoreactive cells, two types were found in this portion, namely closed-type cells and open-type cells.

**Intestine**

Cholecystokinin immunoreactivity: CCK-IR cells were only observed in the lining epithelium of the posterior part of the intestine of *O. hepsetus* (Figure 8). Closed-type and open-type immunoreactive cells were found.

**DISCUSSION**

The stratification of the wall of the DT of *O. hepsetus* has the same organization observed in the majority of other teleosts, with some modifications associated with the species’ feeding habits. We observed four layers: mucosa, submucosa, muscular and serosa. The muscularis mucosae is not present in the examined areas of the DT, unlike that observed in *Pimelodus maculatus* (*P. maculatus*)\(^1\) and *Semaprochilodus insignis*\(^2\). These authors assumed that the existence of muscular tissue between the lamina propria and submucosa aids in the elimination of the substances produced by the glands.

The very large longitudinal folds of the mucosa layer

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**Figure 2 Transversal sections of the esophagus.**

A: Non-keratinized stratified squamous epithelium (arrow). Hematoxylin and eosin stain; B: Presence of adipose tissue (arrowheads) in the submucosa layer. Muscular layer formed by two sub-layers, internal longitudinal (IL) and external circular (EC). Gomori’s trichrome stain; C: Presence of neutral glycoconjugates (GCs). Periodic acid-Schiff stain; D: Acid GCs. Alcian blue stain.
of the esophagus of *O. hepsetus* substantially increases the organ’s capacity for distension, an effect described by other researchers\(^\text{[16,17]}\). The digestive capacity is related to the volume of the folds in the mucosa, with a greater number of folds implying more efficient digestion.

The lining of the mucosa by a stratified squamous epithelium, according to Hunbert *et al*\(^\text{[18]}\), acts to protect the fish against mechanical aggression and invasive bacteria. The same pattern was found in *Prionotus carolinus* by Blake\(^\text{[19]}\), but was not observed in *Salmo trutta* by Burnstock\(^\text{[20]}\) or in *P. maculatus* by Santos *et al*\(^\text{[14]}\). The submucosa contains bundles of adipose tissue and blood vessels; the same was found in *Dentex dentex* (*D. dentex*)\(^\text{[21]}\).

The positive reaction of the epithelium to PAS and AB staining revealed the production of neutral and acid GCs, respectively, in the esophagus. The first type of mucus has low viscosity and is important to assure laminar flow during the lubrication and treatment of particles, to enable digestion to be conducted by the esophagus until the upper region of the stomach. In turn, acid GCs have high viscosity and are fundamental to trap particles\(^\text{[22]}\).

The presence of these GCs was also reported in *Anguilla anguilla*\(^\text{[23]}\), *D. dentex*\(^\text{[21]}\), *P. maculatus*\(^\text{[14]}\), *Cynoscion guatucupa* (*C. guatucupa*)\(^\text{[24]}\), *Pelteobagrus fulvidraco* (*P. fulvidraco*)\(^\text{[25]}\) and *Hyphessobrycon anisitsi* (*H. anisitsi*)\(^\text{[26]}\).

The transition from the esophagus to the stomach is characterized by an abrupt change in the lining epithelium, which starts to present a single layer of columnar cells secreting mucus. This type of stomach lining epithelium has been observed in the majority of other teleosts as...
well\(^{[14,13,12,24]}\), but in *Plecostomus plecostomus\(^{[27]}\) and *Epinephelus marginatus\(^{[28]}\), the epithelium described at the beginning of the stomach was of the squamous and cubic type, respectively, becoming simple columnar in the posterior regions. The mucus-secreting cells were only reactive to PAS in the two stomach regions; the same was found in *C. guatucupa\(^{[26]}\), *P. fulvidraco\(^{[25]}\) and *H. anisitsi\(^{[26]}\), but was not like that observed in *Anguilla anguilla* (*A. anguilla*)\(^{[23]}\) and *Chanos cha-

**Figure 4** Longitudinal sections of the intestine. A-C: Anterior portion; D-F: Posterior portion; A: Overview of the anterior intestine, showing the arrangement of small folds (arrowhead) presenting villi (arrow). Organization of the internal circular (IC) and external longitudinal (EL) muscular layers. Hematoxylin and eosin (HE) stain; B: Simple columnar epithelium with brush border and goblet cells, indicating the presence of acid glycoconjugates (GCs) (arrow). Alcian blue (AB) stain; C: Neutral GCs (arrows). Periodic acid Schiff (PAS) stain; D: Mucosa layer with thick folds (arrowhead). HE stain; E: Myenteric plexus (arrow). HE stain; F: Simple columnar epithelium with striated border and goblet cells with acid GCs (arrow). AB stain; G: Neutral GCs (arrow). PAS stain.
As mentioned, the probable function of this mucus is to promote a flow able to carry the food bolus to the intestine. Besides this, since cells producing hydrochloric acid (HCL), essential for digestion, were identified in the stomach region, this mucus also functions as a layer to protect the epithelial cells.

The invaginations formed by the lining epithelium are called gastric crypts, which in the glandular region communicate with well-developed and branched tubular glands, as also observed by Díaz et al. in *C. guatucupa* and by Domeneghini et al. in *A. anguilla*. These are common characteristics of carnivorous fish species. The gastric glands are composed of oxynticopeptic cells, which play a role similar to that of the principal and parietal cells in mammals, by synthesizing HCL and pepsinogen. In this case, we believe the glandular region has digestive functions while the non-glandular region only acts to carry the food to the gut with the epithelial secretions, with the help of the muscular layer, which in this region is thicker. As seen in the esophagus, the stomach regions also contain well-developed longitudinal folds, whose function is to allow expansion of the organ’s diameter to store a large volume of food, another common characteristic of carnivorous fish species.

The intestine of *O. hepsetus* has two distinct parts, the same as in *Tilapia* spp. The anterior part is characterized by a larger number of thin and elongated longitudinal folds,
which branch out to form villi, making the organ’s lumen very small. The posterior region, endowed with thicker and less sinuous longitudinal folds, contains a larger number of goblet cells, similar to the pattern observed in the large intestine of mammals. This pattern of ample folds of the mucosa is common in carnivores, acting to expand the surface absorption area, since carnivores have a relatively short intestine compared to animals with other feeding habits[39]. Sections of the lining epithelium of both regions, when submitted to the PAS and AB histochemical protocols, reacted positively, showing cells with pink (PAS) and blue (AB) coloring, indicating the presence of neutral and acid GCs, respectively, as also observed by Cinar et al[33] in Pseudophycis antarctica, by Carrasón et al[34] in D. dentex and by Leknes[28] in H. anisitsi, but unlike that observed in P. fulvidraco[30] and Chanos chanos[39].

This allows the hindgut to lubricate the tube and to trap particles to be eliminated, permitting the food bolus to reach this region in dehydrated form[29].

The organization of the muscular layer along the entire DT of O. hepsetus is the same as observed in P. maculatus[31] and P. fulvidraco[28], except in the esophagus, where the pattern resembled that found in mammals. The function of this layer is to promote motility in the DT, carrying and mixing food with the digestive secretions. This motility and also the release of these secretions are favored by the existence of a myenteric plexus between the muscular tissue sub-layers, observed in the glandular region of the stomach as well as in the posterior intestine. Unlike in mammals, the myenteric plexus does not have the form of ganglia, but rather appears in sparse form or in continuous layers, as seen in Pimelodus maculatus[31], but unlike that observed in Salmo trutta[30] and P. fulvidraco[28].

The results obtained in this study demonstrate that the DT of O. hepsetus has three types (5-HT-, SOM- and CCK) of endocrine cells, but three other types (GAS-, GLUC- and INS-IR cells) were not present. However, these cells have been observed in other fish, such as: GLU-IR cells in the gastric mucosa of cartilaginous fishes[33]; GAS-IR cells in the stomach pyloric region of Otolithes mykiss; and INS-IR cells in the stomach pyloric region of Monopterus albus (M. albus) and the stomach cardiac and pyloric regions of Peltocephalus fulvidraco[36]. The reason for the absence of these endocrine cells in the DT of O. hepsetus may be related to its digestive histophysiology, but further studies should be conducted to confirm this relationship.

The peptide SOM is a component responsible for inhibiting many substances, such as GAS, CCK, GLUC, INS, secretin, motilin and gastric acid[37]. In mammals, it also controls the absorption of amino acids and glucose[38]. It is thus essential in the DT, since it participates in basic mechanisms for efficient food processing. Ku et al[39] identified the production of this hormone along the DT of the reptile Trachemys scripta elegans, including in the esophagus, where we found SOM-IR cells in O. hepsetus. Other studies investigating the presence of this hormone in the DT have been performed by Lee et al[40] and Pan et al[41], the latter analyzing the presence of endocrine cells in eight fish species: P. fulvidraco, M. albus, Siniperca chuatsi (S. chuatsi), Colossoma brachypterus and Tilapia nilotica, all of which presented SOM-IR cells in the gastric mucosa, as observed in O. hepsetus. Although we did not visualize this hormone in the intestinal parts of the thin dogfish, it was reported in the gut of the spiny dogfish Squalus acanthias[41], Otolithes mykiss[42], P. fulvidraco, M. albus, S. chuatsi[42], Zacco platypus[43] and Coreoperca bergi (C. bergi)[44].

5-HT-IR cells have been detected in the DT of various vertebrates: fish[42], amphibians[45], reptiles[44], birds[44] and mammals[46,49]. Researchers state that all vertebrates have this type of endocrine cell in the DT, assuming that these cells’ location is based on the evolution of these higher life forms[44]. It is known that in fish, serotonin promotes gastrointestinal motility[50] and blood flow, in the latter case by triggering vasocoonstriction[51,52]. In O. hepsetus, 5-HT-IR cells were observed only in the regions of the stomach, both in the lining epithelium and the glandular epithelium, as also observed in C. bergi by Lee et al[40].

As was described for the fish species Otolithes mykiss[42], Salmo trutta[40], Odontesthes bonariensis[46] and Rhamdia quelen[53], in O. hepsetus we observed CCK-IR cells in the intestine in this case only in the posterior part, while in O. bonariensis these cells were observed throughout the gut, but with greater concentration in the hindgut. There were no CCK-IR cells in the other regions of the DT of O. hepsetus. This hormone controls intestinal motility, by stimulating the release of pancreatic juice and inhibiting gastric emptying[54,55]. The existence of these cells has been verified in fish[56,57], reptiles[58], birds[49] and mammals[52].

In conclusion, our histological and histochemical study of the DT of O. hepsetus revealed adaptation for the species’ feeding habits, to protect the tract and increase the absorptive processes. The immunohistochemical study showed that the DT of this fish species contains different types of endocrine cells similar to those found in other vertebrate species. This study will help comprehension of the digestive physiology of this species and provide a basis for diagnosing diseases that affect the digestive tract of carnivorous teleosts.
Acknowledgments

We thank Iza Lucas Coelho Meirelles for her technical assistance.

Comments

Background

Studies indicate the importance of knowledge of the morphology of the digestive tract (DT) of various species of fish. This method is fundamental for the reduction of food into simpler particles that can be absorbed. The integration of the motor, secretory, and absorptive phenotypes of the DT is essential for the production of food into simpler particles that can be absorbed. This association is achieved by actions and interactions of the nervous and endocrine systems, which in the digestive system are represented by the endocrine cells. The diffuse neuroendocrine system (DNS) acts to control the motility and transit rate of the ingesta, the various secretions of the DT, the absorption of nutrients, and the blood flow, to assure the activation and action of enzymes at the proper moment. Therefore, the products of the digestive system can be absorbed by the organism and reach the blood and lymphatic circulation systems.

Innovations and breakthroughs

The avidin-biotin-peroxidase complex method was applied to study the endocrine cells in the DT of Oligosarcus hepetus (O. hepetus). This method involves the use of three reagents: the primary antibody, which binds to the receptor of the specific hormone of interest; the secondary antibody, which is produced linked to a molecule of the vitamin biotin (C) and binds to receptors of the primary antibody; and the glycoprotein-avidin complex, produced from biotin and peroxidase, with joins with the previous reagent, the secondary antibody.

Applications

The immunohistochemical study showed that the DT of O. hepetus contains different types of endocrine cells similar to those found in vertebrate species. This study will help comprehension of the digestive physiology of this species and provide a basis for diagnosing diseases that affect the digestive tube of carnivorous teleosts.

Terminology

The gastrointestinal epithelium is permeated by a cell, originating from the DNS, called endocrine cells. The secretions of these cells control the digestion of food to ensure it is efficient, by regulating the digestive processes. Besides controlling the absorption of nutrients, they play an important role in determining secretions from the gut and associated glands and in regulating the intestinal blood flow.

Peer review

In this study, the authors describe the microscopic anatomy of the DT of a carnivorous fish species and analyze the functional components that aid the digestion of food. The results are relevant by enabling comparison with other fish species, contributing to phylogenetic studies.

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