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A imunomarcação positiva para c-kit está associada com a presença de células análogas às intersticiais de Cajal no músculo ciliar?

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ABSTRACT

Purpose: Interstitial cells of Cajal were identified in the gastrointestinal tract of several species, with close relation to the enteric nervous system. Since it was recognized that interstitial cells of Cajal express the gene product of c-kit, we performed immunohistochemistry for c-kit protein in ciliary muscle specimens of monkeys' eyes. Methods: Eight eyes from four adult male new world monkeys (Cebus apella) were studied. After blocking endogenous peroxidase activity and nonspecific protein binding, 1:100 dilution of mouse monoclonal antibody against c-kit human oncoprotein was applied to tissues. Antigen-antibody reaction was visualized using the avidin-biotinylated horseradish peroxidase complex in each slide. Results: We observed some groups of fusiform c-kit expressing cells located amongst muscle bundles of the ciliary muscle. Other pigment cells and mast cells were also observed. Conclusion: C-kit expressing cells observed in the ciliary muscle of Cebus apella, showed no similarity to melanocytes or mast cells and they could be associated with their gastrointestinal interstitial cells of Cajal counterpart.

Keywords: Coiled bodies; Ciliary body; Enteric nervous system/physiology; Gastrointestinal motility/physiology; Proto-oncogene proteins c-kit; Cebus

INTRODUCTION

Interstitial cells of Cajal (ICC) were identified in the gastrointestinal (GI) tract over a century ago, and several possible functions were ascribed to these cells on the basis of their morphology and close anatomic relationships with smooth muscle cells and neurons. The original studies describing the existence of a specialized cell type in the tunica muscularis of the GI were those of Cajal. These cells are closely related anatomically and functionally to the enteric nervous system (ENS). Several studies have suggested that the differentiation of ICC is dependent on ENS.

After the description of ICC in the gut wall, the presence of ICC-like cells has been demonstrated in the smooth muscle layers of various organs, such as the human ureter, rabbit portal vein and pancreas. For many years, no conclusive data were available about ICC being smooth muscle cells or fibroblasts. Recently, it was recognized that ICC express the gene product of c-kit, a proto-oncogene that encodes a tyrosine kinase receptor, c-kit. Labeling of kit receptors or c-kit mRNA has
provided an efficient way of identifying ICC throughout the GI tracts of several species, including human, using light microscopy\(^9^{10}\).

Since ICC have been described in several tissues related with the autonomic nervous system, we performed immunohistochemistry for kit protein in monkey eye specimens to study the presence of these cells in the ciliary muscle.

**METHODS**

**Tissues**

Eight eyes from four adult male new world monkeys (*Cebus apella*), of 2.0 to 2.8 kg weight, were studied in agreement with statements of the “Principles of laboratory animal care” (NIH), the OPRR Public Health Service Policy on the Humane Care and Use of Laboratory Animals, the U.S. Animal Welfare Act and the Ethics Committee of Animal Experimentation - Ribeirão Preto Medical School, University of São Paulo. The animals belonged to the Capucchin Monkeys Procreation Nucleus of the School of Odontology of Aracatuba, University of São Paulo. They were sacrificed by a lethal dose of pentobarbital sodium and the eyes fixed with 10% formaldehyde in 0.1 M sodium borate buffer (pH 9.5, 4ºC). Both buffer (pH 6.0, 4ºC) and finally with 2L of 4% paraformaldehyde in 0.1 M sodium acetate by perfusion via ascending aorta with 0.9% saline pH 7.0, by a lethal dose of pentobarbital sodium and the eyes fixed in at least three slides of each fragment. The positive c-kit cells with this specific format, characterized by multiple processes, were located among muscle bundles of the ciliary muscle and did not present toluidine blue metachromatic stain (Figure 1). The location and other observed features of these c-kit expressing cells may be associated with the counterpart of ICC in the GI tract.

Moreover, stained mast cells from the ciliary body stroma were found on all studied slides. They were confirmed specially by their histological features (Figures 2a and 2b). Other pigment c-kit staining cells of the ciliary body included pigment epithelium (Figure 2a). S-100 immunostaining cells were observed specially in the iris’ stroma and displayed different format and tissue location (Figure 3).

**Immunohistochemistry**

**C-kit**

Four-µm thick sections were prepared from two paraffin blocks taken from the dissected fragments. One block was used for hematoxylin and eosin (HE) staining and for detailed histological examination and the other was used for immunohistochemistry\(^9^{10}\). After blocking endogenous peroxidase activity with 3% hydrogen peroxidase in methanol, sections were incubated for 40 min for antigen retrieval. After treatment with 10% normal goat serum for 10 min to block nonspecific protein binding, a 1:100 dilution of mouse monoclonal antibody against c-kit human oncoprotein (CD117, Novocastra Laboratories Ltd, Newcastle-upon-Tyne, UK) was applied to the tissues and the slides were kept overnight at room temperature.

**S-100**

After blocking endogenous peroxidase and nonspecific protein binding, a 1:800 dilution of rabbit polyclonal antibody against S100 (S100 - Ab-2, Lab Vision, USA) was applied to the tissues and the slides were incubated for 60 minutes, at room temperature.

**Antigen-antibody reaction**

The antigen-antibody reaction was visualized in both assays using the avidin-biotinylated horseradish peroxidase complex (Novostain Super Kit ABC\(^TM\), Universal, Novocastra Laboratories Ltd.) and diaminobenzidine as the chromogen (DAB, Zymed Laboratories, San Francisco, CA, USA). Positive and negative controls were performed as recommended by both laboratories.

Slides were counterstained with toluidine blue, in order to identify mast cells’ metachromatic granules.

**RESULTS**

Some groups of elongated c-kit expressing cells with small perikarya and long branching processes were observed in at least three slides of each fragment. The positive c-kit cells with this specific format, characterized by multiple processes, were located among muscle bundles of the ciliary muscle and did not present toluidine blue metachromatic stain (Figure 1). The location and other observed features of these c-kit expressing cells may be associated with the counterpart of ICC in the GI tract.

Moreover, stained mast cells from the ciliary body stroma were found on all studied slides. They were confirmed specially by their histological features (Figures 2a and 2b). Other pigment c-kit staining cells of the ciliary body included pigment epithelium (Figure 2a). S-100 immunostaining cells were observed specially in the iris’ stroma and displayed different format and tissue location (Figure 3).

**DISCUSSION**

In the deep muscular plexus, Cajal\(^2\) described cells with small perikarya and long, branching processes. On the basis of the staining characteristics of ICC with methylene blue and silver chromate, he believed ICC were a type of primitive neuron. In addition, these cells were located close to the nerve strands of the deep muscular plexus and those of the myenteric plexus.

Many of the ultrastructural features of ICC, including a basal lamina, smooth and rough endoplasmic reticulum, dense filaments, cell-to-cell contacts with other ICC and smooth muscle cells, and close contacts with nerve endings, seemed to exclude a nerve or connective tissue nature. Some investigators suggested that ICC are specialized or primitive smooth muscle cells, whereas others thought that ICC possessed the ultrastructural characteristics of fibroblasts\(^1\). To avoid ambiguity, the whole shape of a cell type as well as its relation to nerve and muscle should correspond closely to that originally described by Cajal.

The c-kit gene encodes a transmembrane receptor that has tyrosine kinase activity. C-kit plays a role in hematopoiesis, gametogenesis, and melanogenesis\(^1\). In fact, different cell types express c-kit, such as melanocytes and mast cells\(^1\).
Is the positive c-kit immunostaining associated with the presence of cells analogous to the interstitial cells of Cajal in the ciliary muscle?

Immunohistochemical staining for c-kit became to be accepted as the best marker of ICC on light microscopy. This technique let to an undoubted differentiation between ICC and other local similar cells\(^7,^8\).

We observed, in this study, the presence of some grouped fusiform c-kit expressing cells with small perikarya and long branching processes. We called these cells in the ciliary muscle as “ICC analogous”. They are located amongst muscular bundles of the ciliary body. The presence of mast cells and melanocytes in the ciliary body was described in a previous study, with no description of local ICC\(^9\). That study described “large dendritic mast cells” with numerous long processes, in close association with nerve fibers. Since those c-kit-expressing cells showed no morphological and staining similarities to melanocytes or mast cells (observed with S-100 and toluidine blue, respectively), and were near nerve fibers, they could be associated with their ICC counterparts.

Some specific issues were not well elucidated and should be studied in order to confirm the rationale of the present data. These issues include in vitro physiological tests involving muscular activity, and double or triple identification of all cell types using conventional immunofluorescence assays and serial confocal microscopy.

Until recently, the function of ICC has been speculative and not based on physiological tests. C-kit labeling has improved the understanding of the anatomic relationships between ICC and enteric neurons, smooth muscle cells, and other resident cells in the tunica muscularis. Some physiological studies showed that ICC are involved in the development of electrical rhythmicity and in the regulation of a GI pacemaker system\(^15,^16\).

Moreover, there is evidence suggesting that ICC are essential for rhythmic slow waves of the smooth muscle layers of the small intestine\(^11,^16\).

Because of the close relationship between ICC and ENS of the GI tract\(^16\), we hypothesize that ciliary body ICC analogues could be associated with a sort of physiological mechanism involving the autonomic nervous system and ciliary muscle function, such as maintenance of basal muscle tonus in visual accommodation.

The present study discusses for the first time the possible presence of c-kit-positive ICC in the normal monkey ciliary muscle. It may be hypothesized that a lack of ICC in the ciliary muscle could contribute to dysfunction of this muscle in the aging process in some species. Further studies are needed to confirm and refine the understanding of the relationship of the mechanisms of ICC and ciliary muscle function.

Figure 1 - Immunohistochemical localization of c-kit cells in the monkey ciliary muscle, showing possible interstitial cells of Cajal (ICC) analogous, with a fusiform shape (black arrows) and branches between muscular bundles; microphotography, X1000

Figure 2 - Microphotography of other c-kit staining cells in the ciliary body of Cebus apella; a) note dark-staining cells in the pigmented epithelium (black arrow) and scattered intermediate-staining mast cells in the stroma (white arrows), X400; b) immunostained mast cells around blood vessels in the ciliary muscle (black arrows), X400

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Figure 3 - S-100 immunohistochemistry displaying a typical melanocyte (black arrow) with shorter branches and abundant cytoplasm in the connective tissue of iris; microphotograph, X400

CONCLUSIONS

As immunohistochemical staining of characteristic c-kit cells is accepted as a marker of ICC on light microscopy, we accept that those elongated c-kit-expressing cells visualized in the ciliary muscle could be ICC analogous of the eye.

Functional experiments involving the accommodation process and double or triple immunostaining for c-kit and other local cells are the next step to study these hypotheses.

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