Folliculostellate cells in pituitary pars distalis of male viscacha: immunohistochemical, morphometric and ultrastructural study

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Introduction

Pituitary folliculostellate cells (FSC) have been reported in several mammalian species.1 At present, however, there is no agreement in the literature in relation with the embryological origin and exact function of these cells. In fact, FSC have been reported to play a supporting role, to have phagocytic activity, and to act as regulators of hormone secretion and as stem cells.2-5 FSC have been found lining the lumen of follicles scattered throughout the pituitary pars distalis.5-9 The FSC have been studied using the antibody against S-100 protein, in both mammalian10-12 and non-mammalian species.1,3,13 In addition, some authors have reported in human,11 cat, rat and rabbit14 and mink15 that FSC were immunostained with anti-glial fibrillary acidic protein (GFAP) and anti-vimentin. However, in pituitary of dog and horse these cells did not express GFAP and vimentin, but they expressed the S-100 protein.10 Soji et al.16 have shown an increase in the number of these cells in relation with age, suggesting that the expression of the S-100 protein indicates the cell morphologic maturation stage in rats. Moreover, several authors have demonstrated the existence of different physiological stages of FSC based on their morphological characteristics and immunophenotype.2,7,11,16

On the other hand, there are few reports in the literature on the distribution of FSC in the pituitary parenchyma. Sato et al.12 have reported that FSC are mainly localized in the basal region and transition zone of rat pituitary and also showed an association between FSC distribution and vascularization.

Furthermore, FSC have been reported to synthesize part of the follicular colloid and a close relationship has been described between these cells, the follicular colloid and hormone secreting cells.3,16,18

Ultrastructurally, FSC share distinct characteristics in all studied species. They exhibit scanty cytoplasm with poorly developed endoplasmic reticulum, inconspicuous Golgi complex, moderate amount of mitochondria and absence of the secretory granules.1 Some studies in pituitary have also been performed with lanthanum as electrodense tracer.19,20 Our experimental model is the viscacha (Lagostomus maximus maximus), the largest member of the Chinchillidae family. This rodent inhabits the southern hemisphere from Paraguay through central Argentina.2,21 In its natural habitat, the viscacha is a seasonal breeder2,22. The pituitary pars intermedia (PI), pineal gland, and blood-testis barrier of this rodent have been studied using lanthanum.20,22 Piezzi et al.24 have demonstrated the permeability to lanthanum and the communication of the PI parenchyma with Rathke’s pouch. In this pituitary zone, the FSC originated follicles with PAS-positive colloid inside, expressed S-100, GFAP and vimentin, and were closely associated with melanotrophs.27 Moreover, an ultrastructural study of the pituitary PD has demonstrated the existence of two types of colloid-containing structures: 1) follicles, originated by the FSC, and 2) colloid extracellular accumulations, limited by different types of granulated cells.22 The follicular lumen was filled with abundant electrodense colloid and several cellular debris comprising vacuoles, altered membranes, mitochondria and secretory granules, which are probably a product of FSC phagocytic activity.22,23 In addition, the hormone secreting cells have exhibited regionalization within the PD parenchyma.22,23

Based on the above data and our previous knowledge about the viscacha pituitary, the aim of this work was to perform an immunohistochemical, morphometric and ultrastructural study of the pituitary pars distalis FSC in adult male viscacha to analyze their relationship with hormone secreting cells.
Materials and Methods

Fourteen adult male viscayas weighing 6.0-6.5 kg were captured in their habitat near San Luis, Argentina (33° 20’ south latitude, 760 m altitude) during February-March 2008 (summer), using traps placed in their burrows. According to the results published by Llanos and Crespo and Branch et al., the body weight of this rodent is in relation with its age. Then, the viscayas used in this study were in the same growth stage. After being captured, animals were immediately taken to the laboratory, anaesthetized with Nembutal (pentobarbital; 25 mg/Kg ip.) and killed by decapitation. The brains were rapidly exposed and the pituitaries (weighing 28-30 mg) were excised. The experimental design was approved by the Local Ethics Committee and was in agreement with the Guidelines of the National Institutes of Health (Bethesda, Md., USA) for the use of experimental animals.

For light microscopy, the pituitaries (n=6) were sectioned in the middle plane (Figure 1, line M). Both halves of each pituitary were fixed in Bouin’s fluid, dehydrated in increasing graded ethanol, cleared in xylene, embedded in paraffin and five-µm thick sections were obtained with a microtome (Microm HM 325). Periodic acid-Schiff (PAS) technique was obtained with a microtome (Microm HM 325) fixed in Bouin’s fluid, dehydrated in increasing graded ethanol, cleared in xylene, embedded in paraffin and five-µm thick sections were obtained with a microtome (Microm HM 325). The details, suppliers, dilution, time and temperature of the incubation of antibodies used are reported in Table 1. The tissue sections were first deparaffinized with xylene and graded ethanol, cleared in xylene and mounted with Entellan (Merck, Germany). Labeling was assessed using an Olympus BX-40 light microscope. To confirm the specificity of the immunoreactions the following control procedures were carried out: 1) replacement of primary antibody. No positive structures or cells were found in these sections.

Immunohistochemistry

The streptavidin-biotin immunoperoxidase method was used as described previously. The details, suppliers, dilution, time and temperature of the incubation of antibodies used are reported in Table 1. The tissue sections were first deparaffinized with xylene and hydrated by decreasing concentrations of ethanol. The microwave pre-treatment (antigen retrieval) was performed by incubating the sections in 0.01M citrate buffer (pH 6.0). After incubation for 20 min in a solution of 3% H2O2 in water in order to inhibit endogenous peroxidase activity, they were washed (3x10 min) in phosphate-buffered saline (PBS, 0.01 M, pH 7.4). Non-specific binding sites for immunostaining were blocked by 5% normal rabbit serum or 1% bovine serum albumin in PBS. The immunohistochemical visualization was carried out using the Super Sensitive Ready-to-Use Immunostaining Kit (Catalog n°: QD000-5L, BioGenex, San Ramon, Calif., USA) at 20°C. The sections were incubated for 30 min with biotinylated anti-IgG, and after being washed (3x5 min) in PBS, they were incubated for 30 min with horseradish peroxidase-conjugated streptavidin, and finally washed in PBS.

The reaction site was revealed by 100 µL 3,3'-diaminobenzidine tetrahydrochloride (DAB) chromogen solution in 2.5 ml PBS and 50 µL H2O2 substrate solution, resulting in a brown precipitate. The sections were counterstained with Harris’ hematoxylin for 30 s, washed 10 min in running water, dehydrated in increasing graded ethanol, cleared in xylene and mounted with Entellan (Merck, Germany). Labeling was assessed using an Olympus BX-40 light microscope. To confirm the specificity of the immunoreactions the following control procedures were carried out: 1) replacement of primary antibody with normal goat serum, and 2) omission of primary antibody. No positive structures or cells were found in these sections.

Morphometric analysis

Morphometric parameters were measured with a computer-assisted image analysis system consisting of an Olympus BX-40 binocular microscope, interfaced with a host computer, image processing and recording system. The images were captured by a Sony SSC-DC50A camera and processed with Image Pro Plus 5.0 software under control of a Pentium IV computer. The software allowed the following processes: images acquisition, automatic analogous adjust, thresholding, background subtraction, distance calibration, area measuring and diskette data logging. Before counting, a reference area of 3,000 µm² (×100 objective) was defined on the colour monitor, and distance calibration was performed using a slide with a micrometric scale for microscopy (Reichert, Austria).

For the morphometric analysis, all hemipituitaries were cut from the line M (Figure 1) to the glandular periphery (lines P and P’), and 400 sections (5 µm thick) were obtained in each one. All the serial sections were grouped into 5 sectors (80 sections per sector) and 8 regularly spaced serial sections (50 µm each) were analyzed in every sector. All the microscopic fields captured with ×100 objective were analyzed in each section (340 microscopic fields per section). Therefore, 13,600 microscopic fields were analyzed in each hemipituitary. The following morphometric parameters were determined: *Percentage of S-100-positive total area (%IA S-100): calculated using the formula %IA = ΣRA/Σ100 × 100, where ΣRA was the sum of the S-100-positive area and Σ100 was the sum of the PD area of every microscop ic field. *Percentage of S-100-positive cellular area (%IA S-100): calculated using the formula %IA = ΣAcel/ΣIRA × 100, where ΣAcel was the sum of the area of S-100-positive cells and ΣIRA was the sum of the PD area of every microscopic field. *Percentage of S-100-positive loboidal area (%IA S-100 col): calculated using the formula %IA = ΣAcol/ΣIRA × 100, where ΣAcol was the sum of the area of S-100-positive follicular coloids and ΣIRA was the sum of the PD area of every microscopic field.

Table 1. Features of the antibodies used for immunohistochemical study.

| Antibody | Immunostained cellular type | Clone, catalog and source | Time, Temperature Incubation |
|----------|-----------------------------|--------------------------|-----------------------------|
| Anti-S-100 protein | Folliculo-stellate cell (FSC) | Polyclonal, AR058-5R, Ready-to-Use. BioGenex | 12 h, 4°C in a moist chamber |
| Anti-GFAP | FSC (glial fibrillary acidic protein) | Ready-to-Use. BioGenex | 12 h, 4°C in a moist chamber |
| Anti-Vimentin | FSC | V9, AM074-5M, Ready-to-Use. BioGenex | 12 h, 4°C in a moist chamber |
| Anti-Prl (prolactin) | Lactotroph | Ready-to-Use. DakoCytomation | 60 min, 4°C in a moist chamber |
| Anti-FSHβ (follicle stimulating hormone) | FSH-gonadotroph | 83/72/2A8/2C7. | 12 h, 4°C in a moist chamber |
| Anti-LHβ (luteinizing hormone) | LH-gonadotroph | 3LH 5B6/YH. AM030-5M. | 12 h, 4°C in a moist chamber |
| Anti-ACH (1-24) (adrenocorticotropic hormone) | Corticotroph | Polyclonal, AR035-5R, Ready-to-Use. BioGenex | 60 min, 4°C in a moist chamber |
| Anti-GH (growth hormone) | Somatotroph | Polyclonal, N1561, Ready-to-Use. DakoCytomation | 12 h, 4°C in a moist chamber |
| Anti-TSH (thyroid stimulating hormone) | Thyrotroph | 540H. AM033-5M, Ready-to-Use. BioGenex | 12 h, 4°C in a moist chamber |
Statistical analysis

The results were expressed as mean ± standard error of the mean (SEM) for all data sets. The different groups were evaluated using one-way analysis of variance (ANOVA) followed by Tukey-Kramer multiple comparison test. A probability of less than 0.05 was assumed to be significant.

Double-immunohistochemistry

This technique was performed to examine the association between pituitary hormone secreting cells and S-100-positive FSC. The features of the antibodies are reported in Table 1. DAB and New Fuchsin were selected as chromogens to visualize the antigens because this combination is known to give good contrast. The reaction sites of the first primary antibody (against Prl, FSH, LH, ACTH, GH and TSH) were revealed following the procedure described in the Immunohistochemistry section, using DAB as chromogen. For the second labeling, the slides were washed (3x10 min) in 0.1 M glycine-HCl buffer (pH 2-2.2) at 20ºC for 70 min, washed in PBS, and finally incubated with the second primary antibody (against S-100 protein). Then, the slides were washed (3x10 min) in PBS, and the sections were incubated for 30 min with biotinylated anti-IgG, washed (3x5 min) in PBS, and incubated 30 min with alkaline phosphatase-conjugated streptavidin. The sections were washed in PBS for 10 min, and the reaction sites were revealed by 100 μL New Fuchsin Chromogen Kit (Catalog nº HK 183-5K; BioGenex), resulting in a fuchsia precipitate. The sections were counterstained with Harris’ hematoxylin for 30 s, washed 10 min in running water and mounted with permanent aqueous mounting medium (SuperMount, BioGenex). Labeling was assessed using an Olympus BX-40 light microscope. The specificity of the immunoreaction was confirmed following the above described protocol (see Immunohistochemistry).

Figure 1. Recently removed pituitary of adult male viscacha captured in February (summer). Line M represents vertical cut to obtain two hemipituitaries, and the lines P and P’ correspond to the glandular periphery extremes. The regularly spaced serial sections were grouped in five sectors (1 to 5) in each hemipituitary. Scale bar = 0.1 mm.

Figure 2. Sections of the pituitary of adult male viscacha stained with Hematoxylin-PAS technique. (A) The image shows: pars distalis (PD), pars intermedia (PI), pars nervosa (PN) and Rathke’s pouch (r). The regions and extremes of the pituitary PD are shown: vr, ventral region; dr, dorsal region; ce, cephalic extreme; ca, caudal extreme. (B) Higher magnification of the upper insert in Figure A. Long blood vessels (v) branch out in PD are observed. Arrowheads: PAS-positive colloid. (C) The follicles with PAS-positive colloids inside (arrowheads) are surrounded by one or two layers of endocrine cells (arrows) and blood vessels (v). (D) Higher magnification of the lower insert in Figure A. Blood vessels (v) that communicate pars distalis (PD) with both pars intermedia (PI) and pars nervosa (PN) are observed. r: Rathke’s pouch. Scale bars: A = 500 μm; B and D = 100 μm; C = 25 μm.
Transmission electron microscopy

Eight adult male viscachas were anaesthetized with Nembutal (pentobarbital; 25 mg/kg ip.), killed by decapitation and the brain was rapidly exposed. The pituitaries (n=4) were fixed in a mixture of 3% glutaraldehyde and 4% formaldehyde, pH 7.6 with PBS, then post-fixed in cold 2% OsO4, processed for transmission electron microscopy, dehydrated in acetone, and embedded in Spurr’s resin.

In the remaining animals (n=4), a small volume of the fixative-lanthanum mixture prepared as described below was injected in situ into the pituitary cleft. This procedure was controlled under a stereo microscope. In order to use lanthanum as an intercellular tracer, the method described by Revel and Karnovsky was adopted. A 4% lanthanum nitrate solution was adjusted to pH 7.7-7.8 by slowly adding sodium hydroxide (0.05 N). The lanthanum hydroxide thus obtained was incorporated to an equal volume of s-collidine buffered glutaraldehyde, so that the final concentration of lanthanum was 2%. Then, the pituitary was removed and immersed for an additional 6-hour period in the same solution, post-fixed in cold 2% OsO4, for 2 hours, dehydrated in acetone, and embedded in Spurr’s resin. The experimental design was carried out according to protocols previously used in viscachas in our laboratory.

Sections of 1 μm were stained with toluidine blue and observed under a light microscope (Olympus BX-40) in order to localize the area of interest. Ultra thin sections of approximately 70 nm thick were cut in a Porter Blum ultramicrotome, mounted on copper grids and stained with uranyl acetate and lead citrate. Specimens were finally examined using a transmission electron microscope (Siemens Elmiskop I).

Immunohistochemistry

Immunostaining for the S-100 protein was positive in FSC and some follicular colloid of PD and PI, as well as in the pituicytes of PN (Figure 3 A). Most of the FSC of PD were immunostained with anti-S-100 protein in both nucleus and cytoplasm, while a few cells expressed this protein only in the nucleus or only in the cytoplasm. FSC originated follicles and were occasionally isolated or in small groups. They exhibited a stellate-like shape, irregular nucleus and short cytoplasmatic processes that contacted endocrine cells and blood vessels. Most of the follicles in both pituitary extremes showed heterogeneous S-100 protein immunostaining of the follicular colloid (Figure 3 B-D). On the other hand, a small number of FSC in the PD parenchyma were immunostained with anti-GFAP in the cytoplasm surrounding the nucleus (Figure 3 E-F), and with anti-vimentin in the cytoplasm, mainly in the cytoplasmic processes (Figure 3 G-H). In addition, in each follicle only a few FSC exhibited immunostaining for GFAP and vimentin. Neither GFAP nor vimentin were expressed in the follicular colloid (Figure 3 E-H).

Morphometric analysis

The morphometric parameters varied according to the analyzed sectors. These parameters (%IA S-100, %IA S-100 cel, %IA S-100 col) gradually increased from sector 1 (0.63±0.01, 0.22±0.01, 0.40±0.02, respectively) to sector 3 (0.97±0.02, 0.46±0.01, 0.51±0.03, respectively), and then decreased towards sector 5 (0.47±0.01, 0.28±0.01, 0.19±0.02, respectively). The comparison between the morphometric parameters of the different sectors is shown in Figure 4.

Double-immunohistochemistry

The hormone secreting cells of PD, mainly lactotrophs (Figure 5 A), gonadotrophs (Figure 5 B-C) and corticotrophs (Figure 5 D) were found to be closely associated with FSC and follicles. Frequently, FSC cytoplasmatic processes involved endocrine cells and/or were in contact with the blood vessels. Moreover, the follicular colloid was occasionally immunostained with anti-Prl and anti-LH. The somatotrophs, although widely distributed in the PD parenchyma, did not show a close association with follicular structures (Figure 5 E). Thyrotrophs, which were mainly distributed in the medial and cephalic regions, were not found in the proximity of the FSC or follicles. However, a few thyrotrophs in the caudal extreme were near the follicles (Figure 5 F).

Transmission electron microscopy

FSC exhibited an irregular euchromatinic nucleus with some invaginations into the nuclear membrane, a fine cytoplasm with moderate number of mitochondria, scarce granular endoplasmatic reticulum and absence of secretory granules. They completely surrounded the follicular colloid, exhibited scarce cytoplasm in the apical region, and presented cytoplasmatic processes that contacted secretory cells and blood vessels. FSC were found to develop junctional complexes and desmosomes between their lateral membranes. A large number of microvilli protruded from the apical pole to the follicular lumen (Figure 6 A-B). Lanthanum injected into pituitary cleft freely penetrated between endocrine cells and FSC, but stopped below the FSC junctional complexes without penetrating the follicular lumen (Figure 6 C-F). Occasionally, endocrine cells in different involution or degeneration states presenting irregular and picnotic nucleus were reached by lanthanum (Figure 6 G), but the tracer did not penetrate inside healthy cells.

Discussion

In this work we have described the FSC immunostaining pattern, distribution and association with hormone secreting cells in the pituitary PD of adult male viscacha. Besides, we have shown the existence of an extensive and permeate network of interconnected intercellular spaces between endocrine cells and FSC. In different mammalian species, FSC have been shown to exhibit both nuclear and cytoplasmic immunostaining for S-100 protein. In addition, some authors have reported that they were immunostained with anti-GFAP and anti-vimentin. However, Méndez et al. showed that FSC did not express GFAP and vimentin in pituitary of dogs and horses. Reports in dog and mink pituitary PD have described the presence of two types of S-100-positive FSC. Marin et al. have suggested the existence of different types of FSC in human pituitary on the basis of the immunostaining pattern and localization within the parenchyma. In PI of viscacha, the FSC exhibited irregular nucleus, stellate-like shape and long cytoplasmatic processes which contacted melanotrophs or communicating follicles either with each other or with Rathke’s pouch. In this pituitary zone, scarce vascularization and large follicles (major colloidal diameter: 39.87±0.49 μm) have been informed. On the other hand, Mohamed et al. have described in pituitary PD follicles of smaller size (major colloidal diameter: 12.87±0.38 μm). In the present work, FSC of PD were pleomorphic and showed short cytoplasmatic processes which contacted endocrine cells and blood vessels. The smaller follicle size and the FSC short cytoplasmatic processes...
Figure 3. Immunohistochemistry for S-100 protein (A-D), GFAP (E-F) and vimentin (G-H) in pituitary pars distalis of viscacha. (A) The immunostaining for S-100 protein is observed in the FSC and some follicular colloid (arrowheads) of pars distalis (PD) and pars intermedia (PI). The pituicytes (arrows) of pars nervosa (PN) also exhibit immunostaining for this protein. r: Rathke’s pouch. (B) FSC (arrows) immunostained with anti-S-100 protein forming follicles surrounded by blood vessels (v). The follicular colloid (F) presents heterogeneous immunostaining for this protein. (C) Short cytoplasmic processes (P) of the FSC are in contact with blood vessels (v) and surround endocrine cells (arrowheads). F: immunostained follicular colloid. (D) Immunostaining for S-100 protein is observed in the nucleus (n), cytoplasm (c) and cytoplasmic process (P) of the FSC. The follicular colloid (F) exhibits a heterogeneous immunostaining pattern. v: blood vessel. Inset: A FSC (arrow) with only immunostained cytoplasm and other with a cytoplasmic process (P) surrounding an endocrine cell (arrowhead). (E-F) Scarce FSC (arrows) are immunostained with anti-GFAP in the cytoplasm around the nucleus. F: follicles without immunostaining for GFAP. Inset and Figure F, Short cytoplasmic processes (P) of FSC in contact with blood vessel (v) are observed. n: nucleus without immunostaining for GFAP. (G-H) A few FSC (arrows) show cytoplasmic immunostaining for vimentin, mainly in the cytoplasmic processes (P). The follicles (F) and nucleus (n) of FSC are not immunostained with anti-vimentin. (I) Negative control of immunoperoxidase staining. PD: pars distalis. r: Rathke’s pouch. PI: pars intermedia. Scale bars: A = 250 µm; B-C = 12.5 µm; D-H and Insets = 5 µm; I = 100 µm.
Figure 4. Distribution of S-100 protein in pituitary pars distalis of adult male viscacha. Graph: The values are expressed as mean ± SEM. %IA S-100: percentage of S-100-positive total area; %IA S-100 cel: percentage of S-100-positive cellular area; %IA S-100 col: percentage of S-100-positive colloidal area. Significant differences were determined by analysis of variance followed by Tukey-Kramer multiple comparison test. %IA S-100: a, P<0.01; 3 versus 1 and 5. %IA S-100 cel: b, P<0.01; 3 versus 1 and 5. %IA S-100 col: c, P<0.01; 3 versus 5. The images show details of the pituitary cephalic extremes in the sectors 1 (A), 3 (B) and 5 (C) immunostained with anti-S-100 protein. In sector 3 (B) an increase of the immunopositive area is observed. Arrows: FSC. F: follicular colloid. Scale bars: A-C = 12.5 µm.

Figure 5. Micrographs of double-immunohistochemistry of anterior pituitary hormones (Prl, FSH, LH, ACTH, GH and TSH, brown) and S-100 protein (fuchsia). (A) A close association between lactotrophs (arrowheads) and FSC (arrow) originating follicles (F) are shown. Inset: Single-immunostaining for prolactin. Follicular colloid immunostained with anti-PRL (asterisk) and lactotrophs (arrowhead) near the follicle are observed. (B - Inset) The FSH-gonadotrophs (arrowheads) are contacting FSC (arrow) and follicles (F). v: blood vessel; n: S-100-positive nucleus of FSC. (C) Several LH-gonadotrophs (arrowheads) are associated to FSC (arrows) and follicles (F). Inset: The cytoplasmic process (P) of FSC is in contact with blood vessel (v). Asterisk: Follicular colloid immunostained with anti-LH. (D) The corticotrophs (arrowheads) are located near the FSC (arrows) and follicles (F). Inset: a cytoplasmic process (P) of FSC (arrow) surrounding a corticotroph (arrowhead) is observed. (E) Somatotrophs (arrowheads) exhibit a wide distribution in pituitary pars distalis, but they are only observed in the proximity of follicles (F). Arrow: isolated FSC with a short cytoplasmic process. Inset: Somatotroph (arrowhead), in the second layer limiting the follicle, and a cytoplasmic process (P) of FSC (arrow) reaching the blood vessel (v) are observed. (F - Inset) In the pars distalis caudal extreme, scarce thyrotrophs (arrowheads) in the vicinity of the follicles (F) are observed. Scale bars: A-F = 25 µm; Insets of all Figures = 5 µm.
Figure 6. Electron micrographs of pituitary pars distalis of adult male viscachas. (A-B) Conventional transmission electron microscopy. The folliculostellate cells (FSC) form follicles (F) have irregular nuclei (N), scarce cytoplasm with short cytoplasmic process (P) and absence of secretory granules. The different granulated cellular types (G) are closely associated with FSC, but without contacting the follicular lumen. Well developed junctional complex (arrows) are observed at the apical lateral surface of the FSC. These cells exhibit long microvilli (M) that protrude into the follicular lumen (F). (C-G) Transmission electron microscopy with lanthanum. (C) The tracer stops below the junctional complex (arrowheads) without penetrating in the follicular colloid. (D-F) Lanthanum freely penetrates into the intercellular spaces (arrowheads) between granulated endocrine cells (G) and FSC. The tracer does not penetrate into any healthy cellular type. On the contrary, lanthanum is observed inside the cells in different states of involution and degeneration (G). Scale bars: A = 7 μm; B-C = 4 μm; D-F = 3.33 μm; G = 2 μm.
processes are probably due to the abundant irrigation and close relation between vascularization and follicles that characterize the PD. FSC expressed the S-100 protein and occasionally GFAP and vimentin, which are considered as markers of glial, astrocyte and immature glial cells, respectively. These results in PD support our previous hypothesis for viscacha PI. Moreover, the differential expression of the tested proteins and the immunostaining patterns suggest the presence of FSC in different functional stages, as has also been proposed for viscacha PI and for other species. Vimentin-positive FSC might represent a small reserve population of these cells in an immature state. It has been reported that FSC generate follicles and synthesize part of the follicular colloid. The characterization of these colloidal substances is necessary to establish the FSC precise functions. It has been suggested that colloid is derived from the interstitial fluid of the PD. An alternative hypothesis suggested that the intra and extracellular occur-rence indicate a differential function of this process of colloid formations in pituitary PD of viscacha. In viscacha, permeability to lanthanum and communication with Rathke’s pouch has been demonstrated in PI. This link might function as an alternative route to vascular irrigation and could be involved in the regulation of secretory activity in both adenohypophysial zones. In addition, FSC exhibited junctional complexes, which stopped the penetration of lanthanum, and abundant microvilli in the apical lateral surface, suggesting that they can form and mobilize the follicular colloid. Furthermore, degenerating cells reached by lanthanum may represent an early stage in the process of colloid formations in pituitary PD of viscacha. To sum up, FSC exhibited differential expression for the S-100 protein, GFAP and vimentin, indicating the presence of cells in different physiological stages; the expression of these proteins supports the hypothesis of the neuroectodermic origin of FSC; the S-100 protein was expressed in FSC and the follicular colloid, suggesting different functions of this protein according to its localization; the FSC spatial distribution, their association with endocrine cells, and the generation of an intercellular communication network suggest that these cells are involved in pituitary paracrine regulation. Therefore, the PD of viscacha constitutes a novel and interesting model for further studies of FSC. In addition, the analysis of the S-100 protein accumulated in the follicular colloid could also be of great interest in understanding its extracellular function in the pituitary gland.

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