Ependymal cells surface of human third brain ventricle by scanning electron microscopy

Lorencova M¹, Mitro A¹, Jurikova M¹, Galfiova P¹, Mikusova R¹, Krivosikova L², Janegova A², Palkovic M², Polak S¹

Institute of Histology and Embryology, Faculty of Medicine, Comenius University, Bratislava, Slovakia.
paulina.galfiova@fmed.uniba.sk

ABSTRACT
OBJECTIVES: The ependymal lining of the human brain ventricular system displays distinct structural differences and functional heterogeneity among individual ependymal cells (ECs). To date, multi-ciliated ECs (E1 cells), bi-ciliated ECs (E2 cells), uni-ciliated ECs (E3 cells), ECs without cilia, and ECs with cytoplasmic protrusions have been described in human brain ventricles.

METHOD: Using scanning electron microscopy (SEM), we evaluated ependymal samples from 6 defined regions of the third ventricle from 9 human brains. These regions were strictly defined according to the periventricular structures they neighbour with.

RESULTS: We observed different structures on the apical surface of the ECs. Various ECs differed from each other by the presence of microvilli, secretory bodies, and a variable number of cilia, which led us to divide the ECs into several exactly specified types according to their apical morphology.

CONCLUSION: We found all types of ECs in every examined region with a predominance of particular types of apical surface of ECs in the individual areas (Tab. 4, Fig. 7, Ref. 22). Text in PDF www.elis.sk.

KEY WORDS: human central nervous system, third brain ventricle, ependymal areas, surface of ependymal cells.

Introduction

Ependymal cells (ECs) form the lining of the walls of the brain ventricles. They are special glial cells of the central nervous system. In humans as well as animals, ECs originate from the subpopulation of radial glial cells and differentiate at the exact time of development (1-4).

In general, mature ECs display a squamous, cuboidal, or columnar shape, a regular round slightly hyperchromatic nucleus (granular chromatin), and an inconspicuous nucleolus. They exhibit morphological and functional polarity. The apical portion of cells’ cytoplasm lodges numerous mitochondria. On the apical surface of ECs many microvilli, cytoplasmic protrusions, and clusters of motile cilia are present in most of the cells. The ECs are in direct contact with the environments of different physical properties. The apical side of ECs is in contact with the cerebrospinal fluid (CSF) while the basal portion borders various structures of the underlying nerve tissue (neurons, nerve fibres, glial cells, blood vessels, and others) (5-8).

When considering the apical surface of ECs, three types of cells were classified in the brain ventricles of human and in mice (9, 10). The most common type are multi-ciliated ECs (E1) with numerous motile cilia exhibiting a typical 9+2 microtubule structure. Bi-ciliated ECs (E2) with one or two motile cilia also exhibit the 9+2 microtubule structure, but they differ from E1 cells by the presence of specific basal bodies. These basal bodies exhibit raceme-like appendages and cytoskeletal filaments from their sidewall. Uni-ciliated ECs (E3) represent the third type of ECs. They contain one primary non-motile cilia with a 9+0 arrangement of microtubules and a centriole which is orthogonally positioned to the basal body. A different distribution of ECs was detected in the third ventricle of mice and humans. While the mouse’s ventral part of the third ventricle is covered exclusively with E2 and E3 cells and the dorsal part is covered with E1 cells and the border between them is sharp, human ependymal cells do not occupy such definite locations (9, 10).

Studies in rats using SEM revealed the presence of cilia, microvilli, cytoplasmic protrusions, supraependymal nerve fibres, and supraependymal cells of various shapes on the apical surface of ECs of the third ventricle (11, 12). The ciliary arrangement with 20-30 cilia per EC in the dorsal two thirds of the third ventricle was described as continuous or in tufts. In the ventral third of the third ventricle, ECs mainly possessed microvilli and were mostly devoid of cilia or they displayed scattered solitary cilia. At the
edge of the apical cell surfaces, the microvilli formed a delicate mosaic pattern while non ciliated cells resembled pebbles. The base of the third ventricle (mainly the infundibular recess) contained numerous spherical blebs (also known as cytoplasmic protrusions, apical extrusions, or spheroid formations). In the lower portions and the base of the third ventricle supraependymal nerve fibres were most commonly observed randomly running over the apical surfaces of tanycytes and often contained varicose enlargements (13, 14). Supraependymal cells (including neurons, glial cells, and brain phagocytic cells) were mostly seen in the ventral part of the third ventricle; they can also be present in the dorsal part, but they are not visible through the thick ciliary cover (12). The tran-

Tab. 1. Characteristics of the individuals whose third brain ventricles were analysed.

| Patient's number | Age (years) | Sex  | Cause of death                  |
|------------------|-------------|------|---------------------------------|
| 1                | 51          | male | generalised kidney tumour       |
| 2                | 64          | male | perforated duodenal ulcer       |
| 3                | 80          | male | multiple organ failure          |
| 4                | 78          | female | myocardial infarction          |
| 5                | 61          | male | oesophageal tumour              |
| 6                | 72          | female | hepatic failure                |
| 7                | 73          | female | cardiac failure                |
| 8                | 70          | male | cardiopulmonary failure         |
| 9                | 69          | male | cardiopulmonary failure         |

Fig. 1. Frontal sections (1 – 5) of human third ventricle with designated regions (see red rectangles) of examined ependymal lining (see Table 2).
sition between an area with high ciliary density and a non-ciliated ependyma may progress as a gradual change (15, 16) and is often designated as a transitional zone with a decreased number of cilia and an abundance of microvilli and miniblebs (17).

Since the ependyma displays many structural variations within the ventricular system (11, 17), it is extremely important to determine the exact location of ependyma for tissue sampling so that the samples from different brains can be compared in a relevant way. Because the labelling of individual ependymal areas is not uniform, rather than designating the ependymal areas according to the morphology of cells, we prefer to do so according to the periventricular structures they border (18). Human brain atlases describe 30–35 periventricular structures bordering the third ventricle (19, 20). In our previous study, we used 15 periventricular structures to designate individual ependymal areas of the third ventricle (21). With the help of SEM, we examined the apical surface of 6 defined areas, each of which was examined in 3 brains of different patients. This allowed us to compare the individual regions and to define apical surface of ECs.

Material and methods

In this study, we used tissue from 9 human brains of men and women aged 51 to 80 without any pathological evidence of the central nervous system (Tab. 1). Tissue sampling was performed in accordance with the Health Care Surveillance Authority and the Ethical Committee of the Faculty of Medicine, Comenius University and Bratislava University Hospital (number EK 62/2019).

The brains were sliced within 24 hours post mortem in the auxiliary device to obtain frontal brain slices of equal thickness (0.5 cm). The slices were numbered 1 to 19. The first slice was performed at a distance of 2 cm from the frontal pole of the brain. The distance of each following frontal slice from the previous one was 0.5 cm and the last (19th) slice was at a distance of 11 cm from the frontal pole of the brain. The third ventricle was included in 5 (frontal) sections (Figs 1, 2), beginning in the region of the chiasma opticum. The exact regions of sampling, i.e. parts of the third brain ventricle walls, were defined in relation to periventricular structures as reference nerve tissue (18). We used atlases of the human brain for the terminology of individual periventricular structures, (19, 20) (Tab. 2) (Figs 1, 2) and the previously published suggestion for the labelling of individual ependymal areas (18). The labelling consists of: 1) the designation of the third ventricle (3v), 2) the letter E (as ependyma), and 3) the abbreviation of the Latin name of the periventricular structure bordering the area of interest in the frontal slice. For example, the labelling of the ependyma over the chiasma opticum is 3v-E-CHO. The final designations of the 6 examined ependymal areas are listed in Table 2.

After excision the samples (of up to 50 mm³) were fixed at room temperature by the solution of 3 % glutaraldehyde in an 0.2 M phosphate buffer for 4 hours. After a triple wash with a 0.05 M phosphate buffer, the samples were post-fixed by a 1 % solution of osmium tetroxide, dehydrated in a graded ethanol series to 100 % ethanol, dried at critical point of CO₂ using the BAL-TEC CPD 030, sputter coated with a 15 nm gold layer in LEICA EM ACE 200, and examined with a SEM ZEISS EVO LS 15.

Results

With the help of SEM, we studied the apical surface of ECs in 6 examined areas (Tab. 2). Each area with a surface of 0.25 mm² (which corresponds to apical surfaces of approximately 2500 ECs) was examined in 3 different patients. We focused on the shape and size of the apical membrane of the ECs and the presence of microvilli, cilia, short processes, and secretory bodies. We detected the presence of cilia, 0.3 μm in width and of various lengths. Those longer than 3 μm were designated as long cilia, while those equal to or shorter than 3 μm were designated as short cilia. Cytoplasmic processes with a thickness of 0.3 μm and a length less than 1 μm

Tab. 2. Designation of examined ependymal regions of the 3rd ventricle of five frontal slices defined in accordance with neighbouring periventricular structures (vw – ventral wall, lw – lateral wall).

| Frontal section of the 3rd ventricle | Wall   | Periventricular structure | Abbreviation | Labelling of ependyma | Number of examined patient |
|-------------------------------------|--------|---------------------------|--------------|----------------------|--------------------------|
| 1                                   | vw     | columna fornecis           | CF           | 3v-E-CF              | 2, 3, 5                  |
| 2                                   | vw     | chiasma opticum           | CHO          | 3v-E-CHO             | 4, 7, 8                  |
| 3                                   | lw     | corpus mammillare         | CM           | 3v-E-CM              | 3, 6, 8                  |
| 4                                   | lw     | nucleus paraventricularis thalami | T | 3v-E-T              | 3, 4, 5                  |
| 5                                   | lw     | nucleus ruber             | NR           | 3v-E-NR              | 1, 3, 8                  |
were designated as short. We also detected the presence of microvilli with a thickness of 0.1 μm and a length up to 1 μm. Based on the number and length of cilia present on the apical surface, we categorised the ECs into 13 different groups (Figs 3, 4, 5 and Tab. 4). We observed the surface area of 50 x 50 of neighbouring ECs. When we detected within this area any of 13 ependymal types, we marked its presence as positive (+) into the table. If a specific ependymal type was present 10 and more times within a cluster of 100 neighbouring ECs, we marked its presence as increased (++)..

Our findings are summarised in Tables 3 and 4.

6 distinct areas – results of SEM observations

**Area 3v-E-CF** was evaluated in patients 2, 3, and 5 (Tab. 1). The apical surface of ECs in 3v-E-CF displayed a hexagonal, slightly elongated shape with dimensions of 11 μm x 8 μm (Fig. 6A). A comparison of the apical portions of ECs among the cohort of brains of three patients did not reveal the predominance of any type of cell in this specific area. The number of secretory bodies in all three brains of patients indicates that the secretory activity of ECs in this area was low.

**Area 3v-E-CHO** was evaluated in patients 4, 7, and 8 (Tab. 1). The apical surface of ECs in this area had a hexagonal shape

---

**Tab. 3. Characteristics of apical surfaces of ECs in the examined patients (P) observed with a SEM in marked ependymal areas of the third human brain ventricle (+ positive presence, ++ elevated incidence).**

| Labelled ependymal area | Characteristics of apical surfaces of ependymal cells | Obvious secretory bodies (sb) on the apical cell surfaces |
|-------------------------|------------------------------------------------------|---------------------------------------------------------|
|                         | Shape of apical cell membrane                        | sb ≤ 2 μm                                                |
|                         | Dimensions of apical cell membrane                   | sb ≥ 2 μm                                                |
| 3v-E-CF                 | hexagonal                                             | P3+ P2+ P5+                                             |
| 3v-E-CHO                | hexagonal                                             | P8+ P4+ P7+                                             |
| 3v-E-I                  | hexagonal                                             | P8+ P9+ P7+                                             |
| 3v-E-CM                 | hexagonal                                             | P8+ P6+ P3+                                             |
| 3v-E-T                  | hexagonal                                             | P3+ P5+ P4+                                            |
| 3v-E-NR                 | hexagonal                                             | P8+ P1+ P3+                                            |

---

Fig. 3. A: ependymal cell (EC) without cilia (type 1), only microvilli are present B: EC with short cillum (<3 μm in length) (type 2) C: on the bottom of the image - EC with one long cillum (≥3 μm in length) (type 3) and on the top – EC with two short cilia (type 4) D: on the right side – EC with two long cilia (type 5), on the left side – EC with one short and one long cillum (type 6).

Fig. 4. A: on the right side of the image - ependymal cell (EC) with 3–7 cilia (type 7), on the left side - EC with 8–15 cilia (type 8) B: ECs with more than 15 cilia (type 9 – multi-ciliated ECs) C: on the right side EC with very short processes (type 11) D: on the right side – EC with short processes (<1 μm in length) in combination with short cilia (<3 μm in length) (type 12).

Fig. 5. A: ependymal cell (EC) with short processes (<1 μm in length) and long cilia (≥3 μm in length) (type 13) B, C, D: presence of secretory vacuoles of different sizes on the ependymal surface.
with dimensions of 10 μm x 8 μm. Based on the observations of brains of all three patients, we characterised the apical surface of ECs from this area as non-ciliated with short processes or containing few cilia (types 1–7, 11, and 12 in Table 4) (Fig. 6B). The secretory activity of these cells was minimal.

Tab. 4. Incidence of various types of ECs in the examined patients (P) according to the presence of cilia and various sized processes on the apical surface of ECs in the examined areas of the third human brain ventricle (+ positive presence, ++ elevated incidence).

| Labelled ependymal area | 3v-E-CF | 3v-E-CHO | 3v-E-I | 3v-E-CM | 3v-E-T | 3v-E-NR |
|-------------------------|---------|----------|--------|---------|--------|---------|
| without cilia, only microvilli are present (type 1) | P3 P2 P5 | P8 P4 P7 | P8 P9 P7 | P8 P6 P3 | P3 P5 P4 | P8 P1 P3 |
| 1 short cilium (type 2) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| 1 long cilium (type 3) | P3 P2 P5 | P8 P4 P7 | P8 P9 P7 | P8 P6 P3 | P3 P5 P4 | P8 P1 P3 |
| 2 short cilia (type 4) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| 2 long cilia (type 5) | P3 P2 P5 | P8 P4 P7 | P8 P9 P7 | P8 P6 P3 | P3 P5 P4 | P8 P1 P3 |
| 1 short and 1 long cilium (type 6) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| 3–7 cilia (type 7) | P3 P2 P5 | P8 P4 P7 | P8 P9 P7 | P8 P6 P3 | P3 P5 P4 | P8 P1 P3 |
| 8–15 cilia (type 8) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| more than 15 cilia (type 9) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| short and long cilia (type 10) | P3 P2 P5 | P8 P4 P7 | P8 P9 P7 | P8 P6 P3 | P3 P5 P4 | P8 P1 P3 |
| short processes only (type 11) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| short processes with short cilia (type 12) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |
| short processes with long cilia (type 13) | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ | ++ ++ ++ |

Fig. 6. A: hexagonal shape of ECs (3v-E-CF) B: area with rare incidence of cilia (3v-E-CHO) C: secretory body protruding from the EC lateral surface (3v-E-I) D: ependymal surface with obvious secretory activity (3v-E-I).

Fig. 7. A: suprapependymal cell in 3v-E-CM B: apical surfaces of ECs with little or no cilia and with the presence of secretory bodies in 3v-E-CHO C: presence of cilia, microvilli and secretory bodies on apical surface in 3v-E-I D: ciliated ependymal surface in 3v-E-NR.

Area 3v-E-I was evaluated in patients 7, 8, and 9 (Tab. 1). The apical membrane of ECs in this area was of a hexagonal, sometimes irregular shape with dimensions of 9 μm x 9 μm. Numerous secretory bodies were present on the surface of ECs coming from the depth or originating from the lateral or apical surface.
Based on the study of the apical surface of ECs in 3v-E-CM, the ECs in all three examined patients had the dimensions of apical surfaces of 5 μm x 6 μm, which is less in comparison to 3v-e-CF, 3v-e-CHO, 3v-e-I, 3v-e-T and 3v-e-NR. This seems to be due to the higher density of ECs in this area. In brains of all three patients, there were locations with ECs which showed a large number of cilia on its apical surface (EC types 8, 9, 10 according to Tab. 4) and locations with little or no cilia (ECs type 1, 2, 3, 4, 11). The ratio of more to less ciliated locations varied in individual patients. Secretory bodies were more or less present in brains of all three patients. In patient No. 3, the shape of the ventricular wall in 3v-E-CM was slightly folded, and ECs in elevated areas of ridges displayed less cilia, compared to deeper areas of grooves which possessed numerous cilia on the apical surfaces of the ECs. We also observed the presence of supraependymal cells (Fig. 7A).

The shape of the apical membranes of ECs in this area was regular and hexagonal with a diameter of approximately 8 μm. When considering the results of 3v-E-T from brains of all three patients, that was characterised by the presence of secretory bodies with a diameter as well as larger than 2 μm (Fig. 7B), which could indicate a high secretory activity. Although in general, ECs with a small number of cilia dominated in this area (types 1–7, 11 according to Tab. 4), the folded surface of patient No. 3 contained alternating locations of high number (especially in grooves) and low number of cilia.

The observed secretory activity of ECs varies in number and length (Tab. 4), had a smooth surface and the same diameter of 0.3 μm throughout their entire length. Mirzadeh et al (10) defined three basic groups of ECs according to their apical surface morphology: E1 cells – multi-ciliated ependymocytes with motile cilia (we labelled them as type 9), E2 cells – bi-ciliated ECs with one to two motile cilia (we divided these according to cilia length and labelled them as types 4, 5, 6), and E3 cells – uni-ciliated ECs with primary cilium (which we labelled according to the length of the cilium as types 2 and 3). E2 and E3 cells contain a long basal process thanks to which various signals reach more distant locations in the brain.

Using SEM, we examined the apical surface of ECs of the third human brain ventricle in six precisely designated locations, and examined them in brains of three patients (Tabs 1 and 2). We observed cilia, microvilli, very short processes and secretory bodies on the surface of ECs. We also occasionally observed supraependymal nerve fibres and cells which we did not specify.

The cilia of individual ECs varied in number and length (Tab. 4), had a smooth surface and the same diameter of 0.3 μm throughout their entire length. Scott et al (16) used SEM to analyse the third human brain ventricle based on the presence of periventricular structures (Fig. 1, 2, Tab. 2) (7, 21).

The cilia of individual ECs varied in number and length (Tab. 4), had a smooth surface and the same diameter of 0.3 μm throughout their entire length. Scott et al (16) used SEM to analyse the third human brain ventricle based on the presence of periventricular structures (Fig. 1, 2, Tab. 2) (7, 21).

Using SEM, we examined the apical surface of ECs in six precisely designated locations, and examined them in brains of three patients (Tabs 1 and 2). We observed cilia, microvilli, very short processes and secretory bodies on the surface of ECs. We also occasionally observed supraependymal nerve fibres and cells which we did not specify.
cline in cilia density and an appearance of microvilli and “rounded lucent excrescences”. They characterized the ventral third of the third ventricle with an infundibular recess as almost free of cilia but with the presence of numerous microvilli and “large lucent excrescences”. Our examined areas (3v-E-CF, 3v-E-T a 3v-E-NR) correspond with the central third of the third ventricle in the dorsal-ventral axis (Figs 1, 2). In terms of all three areas, locations with a low and high number of cilia alternated here, and in 3v-E-NR the transitions between multiciliated and little ciliated locations were the most obvious. The “rounded lucent excrescences” correspond to secretory bodies in the examined samples. According to our observations, 3v-E-CF was the least active in terms of secretion in contrast to 3v-E-NR, which showed a high secretory activity. In general, our results correspond to previous observations (15, 16), although we precisely specified and defined the observed regions. 3v-E-CHO, 3v-E-I and 3v-E-CM, which we examined and clearly defined, correspond to the ventral third of the brain ventricle. In general, we noticed the low number of cilia and higher number of microvilli in this area; however, if we take into consideration the individual examined regions separately, this statement does not equally apply for all locations and all patients. For example, we also found locations in 3v-E-CM and 3v-E-I with an abundant incidence of cilia. The secretory activity within the ventral third of the brain ventricle was most distinctive in 3v-E-I, which corresponds to previous observations in humans and animals (15, 16). However, according to our observations, the secretory activity derived from the number of observed secretory bodies in 3v-E-I was comparable with 3v-E-NR in the central third of the brain ventricle (Tab. 3).

When observing the folds of the ventricular surface we repeatedly noticed ECs with numerous cilia in grooves and ridges that were covered by ECs with little or no cilia. This finding was independent of the location of the examined ependymal area; we found it in 3v-E-I, 3v-E-CM, 3v-E-T, and 3v-E-NR. In our opinion, the numerous cilia in grooves could be related to the increased need to prevent liquor flow stagnation, while individual cilia on the surfaces of ridges could represent a sensory function. Mirzadeh et al (10) attribute such function to uni- and bi-ciliated ECs.

The apical surface of the ependymal lining of the third brain ventricle is extremely heterogeneous. Significant heterogeneity was also observed in individual evaluated ependymal regions, where we found all types of apical surfaces. The systematic mapping of the ventricular surface significantly contributes to the clarification of ECs functions in individual locations.

References

1. Sarnat HB. Role of human fetal ependyma. Pediatr Neurol 1992 b; 8 (3): 163–178.
2. Bruni JE, Del Bigio MR, Clattenburg RE. Ependyma: normal and pathological. A review of the literature. Brain Res 1985; 356 (1): 1–19.
3. Bruni JE. Ependymal development, proliferation, and functions: a review. Microsc Res Tech 1998; 41 (1): 2–13.
4. Coletti AM, Singh D, Kumar S et al. Characterization of the ventricular-subventricular stem cell niche during human brain development. Development 2018; 145 (20).
5. Del Bigio MR. Ependymal cells: biology and pathology. Acta Neuropathol 2010; 119 (1): 55–73.
6. Mathew TC. Scanning electron microscopic observations on the third ventricular floor of the rat following cervical sympathectomy. Folia Morphol (Warsz) 2007; 66 (2): 94–99.
7. Mitro A, Kiss A. Ependymal tables designated for differentiation of the ependyma based on the adjacent periventricular structures. Biologia 2016; 71 (5): 603–611.
8. Mitro A, Polák Š, Filipčík P. Ependyma of the human brain ventricles. Giza: El-Meleigy Press, 2008: 1–100.
9. Mirzadeh Z, Merkle FT, Soriano-Navarro M, Garcia-Verdugo JM, Alvarez-Buylla A. Neural stem cells confer unique pinwheel architecture to the ventricular surface in neurogenic regions of the adult brain. Cell Stem Cell 2008; 3: 265–278.
10. Mirzadeh Z, Kunse Y, Duran-Moreno M et al. Bi- and uniciliated ependymal cells define continuous floor-plate-derived tanycytic territories. Nat Commun 2017; 8: 13759.
11. Mathew TC. Regional analysis of the ependyma of the third ventricle of rat by light and electron microscopy. Anat Histol Embryol 2008; 37 (1): 9–18.
12. Ray PK, Choudhury SR. Changes in the surface fine structure of ependyma of the rat third ventricle following operative leakage of cerebrospinal fluid. J Anat 1985; 140 (1): 1–11.
13. Scott DE, Sladek JR Jr. Age related changes in the endocrine hypothalamus: I. Tanycytes and the blood-brain-cerebrospinal fluid barrier. Neurobiol Aging 1981; 2 (2): 89–94.
14. Haemmerle CA, Nogueira MI, Watanabe IS. The neural elements in the lining of the ventricular-subventricular zone: making an old story new by high-resolution scanning electron microscopy. Front Neuroanat 2015; 9 (134).
15. Bruni JE, Montemurro DG, Clattenburg RE, Singh RP. A scanning electron microscopic study of the ependymal surface of the third ventricle of the rabbit, rat, mouse and human brain. Anat Rec 1972; 174 (4): 407–420.
16. Scott DE, Paul W, Dudley GK. A comparative scanning electron microscopicanalysis of the human cerebral ventricular system. I. The third ventricle. Zellforsch Mikrosk Anat 1972; 132 (2): 203–215.
17. Choudhury SR, Azzam NA, Donohue JM. Changes in the surface fine structure of rat third ventricular ependyma following chronic acetazolamide treatment. J Anat 1979; 129 (1): 51–62.
18. Mitro A. Method of labelling of individual ependymal areas according to periventricular structures of the rat lateral brain ventricles. Biologia 2014; 69 (9): 1250–1254.
19. Mai JK, Assheuer J, Paxinos G. The neural elements in the lining of the ventricular-subventricular zone: making an old story new by high-resolution scanning electron microscopy. Front Neuroanat 2015; 9 (134).
20. Ding SL, Royall JJ, Sunkin SM et al. Comprehensive Cellular-Resolution /Atlas of the Adult Human Brain. J Comp Neurol 2016; 524: 3127–3481.
21. Mitro A, Lorencova M, Mikušová R, Gállová P, Káttna V, Polák Š. Labelling of individual ependymal areas in the third and fourth ventricle of the human brain: ependymal tables Biologia 2019; 74: 533–541.
22. Lee DA, Bedont JL, Pak T et al. Tanyocytes of the hypothalamic median eminence form a diet-responsive neurogenic niche. Nat Neurosci 2012; 15 (5): 700–702.

Received January 28, 2020. Accepted March 31, 2020.