Metabolic and morphological changes in rats’ eye after 7-days microgravity simulation

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Abstract: Microgravity simulation in rats resulted to changes in eye lavage content. Molecular subfraction analysis with laser correlation spectroscopy revealed significant differences in the pattern of particle distribution in five intervals: 1.9 -2.56 nm, 4.64-6.25 nm, 91.3 nm, 165-300 nm, and 400-734 nm. Microgravity modeling induced a series of interrelated processes in the retina of experimental rats manifesting in decreasing of total and layer-by-layer thickness of the retina, mild edema, and consequently, the increasing of specific density of neurons. In the inner nuclear layer, the cell density decreased because of the amacrine and bipolar cells. These changes could be explained by relative ischemia of the retinal layers, which are dependent on the retinal vessels circulation system.

Keywords: Amacrine Cells, Bipolar Cells, Laser Correlation Spectroscopy, Microgravity Simulation, Muller Cells, Retinal Vessels

1. Introduction

During the last years, following the increased intensity of space flights, the investigations of weightlessness influence on the human body in general as well as on the delicate and high developed body functions are also increasing.

It was published [1] that approximately 20% of astronauts on International Space Station (ISS) missions have developed measurable ophthalmic changes after flight. Authors found out the significant role of such nutrition factors as folate- and vitamin B-12-dependent 1-carbon and related biochemical indices and alteration of their metabolic pathway. The biochemical differences observed in crewmembers with vision issues strongly suggest that their folate- and vitamin B-12-dependent 1-carbon transfer metabolism was affected before and during flight. According to these changes were found out the hyperopic shift in cycloplegic refraction. It was suggested that polymorphisms in enzymes of this pathway may interact with microgravity to cause these pathophysiologic difference. It was also hypothesized that microgravity-induced cephalad fluid shifts and resultant increased intracranial pressure and/or localized intraorbital changes may be involved.

It was also found out that microgravity in humans cause decreasing of intraocular pressure (IOP) [2], as well as optical coherence tomography (OCT) documented an increase in retinal thickness and volume with following corresponding scotomas. It took almost 6 months until these changes came back to the baseline. Severe cases presented additional signs, such as globe flattening with hyperopic shifts, choroidal folds and cotton wool spots, resulting in some cases in permanent visual impairment. It is clear, that all these changes can lead to the changes of visual functions and influence the ability of the staff to make right and quick decisions.

Further investigations showed neuronal loss after a spaceflight [3], increasing the speed of cataract formation [4] and damage of retinal pigment epithelium [5]. Currently, the real reasons and pathogenesis of visual changes during and after the spaceflight are unknown, that is why we decided to
perform our investigation.

2. Materials and Methods

Experiments were performed on male Wistar rats weighing 220-270 g. The animals were divided into two groups: experimental group rats (n=186) served for microgravity modeling and controls (n=193) stayed under vivarium conditions. The rats of both groups received standard laboratory chow throughout the experiment and had free access to food and water. The studies were approved by the Ethical Committee and were performed with strict adherence to the corresponding rules of good laboratory practice.

The effects of microgravity were modeled by suspension of the experimental animals [6, 7] for 7 days in a special stand with a head-down tilt of 30°. The stand consisted of a frame with a beam rolling between the two guide rails and carrying a movable carriage. The carriage moved in all directions in the horizontal plane above a grid floor (40x40 cm), under which a tray for collection of excrements was placed. For uniform body weight distribution, a special system consisting of a Capron sheath with holes for limbs was used for suspension of the animal to the movable carriage. Two metal plates placed in the sheath along the animal back prevented sagging and ensured the chosen suspension angle for the animal body.

Lavage samples from both eyes were collected as follows: 200 µl physiological saline was applied to the eye with a disposable tip without touching the cornea and simultaneously collected with another pipette to avoid the contact of fluid excess with the hair. The samples were stored at +4°C for 3-4 h or at -20°C for no more than 5 days until assay. Immediately before the analysis, the samples were centrifuged for 15 min at 3000 rpm and the supernatant was added to the measuring cell of the instrument. Laser correlation spectroscopy (LCS) allows analysis of the subfraction composition of bioingredients in a wide range of their molecular sizes (from 1 to 10,000 nm). From the physical point of view, the method is based on principles of Doppler spectroscopy [8]. Detailed LC-histograms formed by the spectrometer operational program consist of 32 columns. The number of columns corresponds to the number of molecular subfraction analyzed during spectrum processing (minimization), while their height reflects the contribution of the corresponding fraction into light scatter [9].

After the experiment, the rats were sacrificed by hypnotic drug overdose, enucleated eyes were fixed in cold 2.5% glutaraldehyde for 2-8 h; penetrating circular incision in the prelimbal area were preliminary made. After washing in phosphate buffer, the anterior segment of the eye was separated along the projection of the pars plana of the ciliary body, the lens was removed, and a horizontal block was cut out. Fragments of eyeball wall (1.5×1.5 mm) containing all three layers corresponding to posterior pole, equatorial zone, and peripheral zone were isolated. The fragments were postfixed in 1% osmium tetroxyde (1 h), dehydrated in increasing concentrations of an organic solvent, and embedded in mix of epoxy resins epon-araldite (1:1). Semi-thin 0.5-1.5-µ sections were cut on an Ultratome-IV (LKB, Sweden) and stained with methylene blue and fuchsin (polychrom staining). The preparations were examined under a Leica DM-2500 light microscope, photographed using a Leica DFC320 camera, and images were analyzed using Image Scope Color software.

3. Results and Discussion

Two experimental series with 7-day simulation of microgravity were performed, the interval between the series was 9 months. First, it was necessary to ensure stability and reproducibility of the results. To this end, we compared the results obtained for the left and right eyes of experimental and control animals. As is seen from Figures 2 and 3, there were no significant differences in the particle distributions in the washout fluids from the left and right eyes in both the control and experimental groups. Minor differences between the series can be explained by seasonal fluctuations of serum homeostasis. These findings allow us pool the results obtained from both eyes within the same series.

Comparison of the results in the control and experimental groups obtained in the two experimental series is presented in Figure 4. The main significant differences in the pattern of particle distribution were found in five intervals: 1.9 -2.56 nm, 4.64-6.25 nm, 91.3 nm, 165-300 nm, and 400-734 nm. It should be noted that these shifts in both series had the same direction in all intervals, except the last one. In the experimental group, the contribution into light scatter was higher in zone 1 corresponding to albumins. Elevated content of albumins can be caused by dehydration caused by the experimental conditions. In addition, this zone can contain small proteins released as a result of degenerative processes in cells. Zone 2 (4.64-6.25 nm) contains globulins; their reduced content in rats of the experimental group can attest to vulnerability of the immune system caused by microgravity exposure [10 – 12].

Figure 1. Experimental animal in the stand for microgravity modeling.
**Figure 2.** Histogram of size distribution of particles in control rats’ eye lavages. Ordinate – contribution of particles of the corresponding zone into light scattering (%). Abscissa – particle size (nm).

**Figure 3.** Histogram of size distribution of particles in suspended rats’ eye lavages. Ordinate – contribution of particles of the corresponding zone into light scattering (%). Abscissa – particle size (nm).
The peak at 91 nm, as was demonstrated in experiments on lymphocytes [13], is determined by the release of nucleoproteins into the extracellular medium. In our experiments, the contribution of this zone into light scatter in the experimental group was higher than in the control. This can indicate enhanced cell destruction under modeled microgravity conditions. In the experimental group, the contribution of zone 165-300 nm particles into light scatter was lower than in the control. A similar decrease in this zone was observed in patients with keratoconjunctivitis [14]. Zone 400-734 nm contains large immune complexes and probably cell destruction products. Instability of the results in this zone in the two experimental series can be related to seasonal changes.

Normally on semi-thin sections, rat retina by its architectonics and vascularization pattern is very similar to human retina. An important difference is that it has thinner outer and thicker inner plexiform layers. Near the posterior pole, the rat retina has a total thickness of 256±18.4 µ; the thickness of different layers is shown in Table 1. The nerve fiber layer contains the major retinal blood vessels (Fig. 5). In the equatorial zone, the thickness of the retina is decreasing to 204.7±14.2 µ, mainly because of the outer nuclear and inner plexiform layers (IPL) (Fig. 6). At higher resolution, retinal blood vessels could be seen at the level of the outer plexiform layer (OPL) (Fig. 7). At the periphery, the retina thickness is decreasing up to 133.7±11.4 µ, mainly also because of the plexiform layers. The specific density of neurons and glial cells equally decreasing as well.

| Parameter                  | Central area       | Equatorial area   | Peripheral area |
|---------------------------|--------------------|-------------------|-----------------|
| Total thickness, µ        | 256±18.4           | 204±14            | 146±11          |
| Thickness of retina layers, µ | |                   |                  |
| RPE (retinal pigment epithelium) | 6.7±0.82            | 4.1±0.51          | 3.8±0.63        |
| PHR (photoreceptors layer) | 38.4±1.12           | 34.6±3.07         | 28.2±1.88       |
| ONL                        | 52.1±3.55          | 42.4±2.2          | 39.3±1.14       |
| OPL                        | 14.4±1.88          | 10.2±2.02         | 9.7±1.86        |
| INL                        | 39.3±1.97          | 34.1±1.36         | 25.6±2.79       |
| IPL                        | 68.46±5.22         | 59.5±3.34         | 49.3±3.54       |
| GCL (ganglion cells layer) | 20.1±1.67          | 14.7±3.04         | 10.7±2.42       |
| Cell density in ONL, cells/mm² | 41397              | 35916             | 31897           |
| Average area of nuclei in ONL, µ² | 13.2±1.89          | 11.1±2.04         | 10.4±2.18       |
| Cell density in INL, cells/mm² | 18506              | 13785             | 18506           |
| Average area of nuclei in INL, µ² | 34.7±10.1           | 32.5±9.2          | 28.4±8.7        |
| Density of ganglion cells, cells/mm² | 4501               | 3708              | 3308            |
| Average area of nuclei in ganglion cell layer, µ² | 52.6±12.5          | 42.1±12.1         | 38.3±18.4       |

Table 1. Morphometric parameters of rat retina (control group, n=5)

![Figure 4](image-url)  
* - comparison of control and suspended groups, p< 0.05; ** - comparison of control and suspended groups, p< 0.005; *** - comparison of control and suspended groups, p< 0.001 (Mann-Whitney test).
After the 7 day of experiment, the total thickness of the retina in the central zone (posterior pole) decreased a little (up to $218\pm3.95\ \mu m$) without specific predominance of any layer. In contrast to narrowed retinal arterioles, retinal veins looked dilated (Fig. 9), this change of vessel diameter can lead to hypoxia of the inner layers of the retina and alterations related to venous stasis. Also at this period we can see that neuronal density in the corresponding layers slightly decreased, but the size of neurocyte nuclei was increased, thus the difference between the layers thickness in the compared groups disappears (Table 2). Increased nuclei size can be a sign of over activation of the synthetic processes, which exhaust cell reserves.

**Table 2. Morphometric parameters of rat retina (experimental group, $n=8$)**

| Parameter                                      | Central area    | Equatorial area | Peripheral area |
|------------------------------------------------|-----------------|-----------------|-----------------|
| Total thickness, $\mu m$                      | $218\pm3.95$    | $182.4\pm14$    | $133 \pm11$     |
| Thickness of retina layers, $\mu m$           | $4.31\pm0.72$   | $4.18\pm0.51$   | $3.84\pm0.63$   |
| RPE (retinal pigment epithelium)               | $34.3\pm1.52$   | $34.6\pm3.07$   | $26.2\pm2.58$   |
| PHR (photoreceptors layer)                    | $49.9\pm1.19$   | $36.6\pm2.2$    | $30.3\pm1.14$   |
| ONL                                            | $10.2\pm2.1$    | $9.84\pm1.02$   | $5.11\pm1.58$   |
| OPL                                            | $32.5\pm1.87$   | $25.6\pm3.72$   | $17.7\pm1.79$   |
| INL                                            | $62.3\pm3.39$   | $59.5\pm3.34$   | $43.3\pm4.54$   |
| GCL (ganglion cells layer)                    | $15.8\pm2.68$   | $13.3\pm1.84$   | $8.97\pm1.41$   |
| Cell density in ONL, cells/mm$^2$             | $38710$         | $36581$         | $32369$         |
| Average area of nuclei in ONL, $\mu m^2$      | $14.9\pm2.15$   | $13.5\pm3.14$   | $12.8\pm3.74$   |
| Cell density in INL, cells/mm$^2$             | $15721$         | $13957$         | $12600$         |
| Average area of nuclei in INL, $\mu m^2$      | $33.4\pm5.7$    | $35.7\pm8.1$    | $39.7\pm2.6$    |
| Density of ganglion cells, cells/mm$^2$       | $3801$          | $3608$          | $3278$          |
| Average area of nuclei in ganglion cell layer, $\mu m^2$ | $51.6\pm12.5$   | $55.9\pm11.4$   | $61.7\pm16.5$   |
Figure 8. Control group. Peripheral area. Dilated vessels in the nerve fiber layer (arrows). Semi-thin section. Staining with methylene blue and fuchsin.

Figure 9. Experimental group. Posterior pole area. Dilated retinal vein (arrow). Semi-thin section. Staining with methylene blue and fuchsin.

In the equatorial area, rarefaction and enlargement of neurons in inner nuclear layer (INL) becomes more evident. Thinning of INL from the inner side visually appears like "movement" of the Muller cells nuclei closer to the boarder of IPL (Fig. 10).

Figure 10. Experimental group. Equatorial area. Thinning of inner nuclear layer “brings” nuclei of Muller cells (arrows) towards the inner plexiform layer. Semi-thin section. Staining with methylene blue and fuchsin.

Figure 11. Experimental group. Equatorial area. Aggregation of erythrocytes in deep retinal capillaries (arrow). Edema and rarefaction of amacrine (Am) and bipolar (BP) cells. M – Muller cells. Semi-thin section. Staining with methylene blue and fuchsin.

At higher magnification, we can see that the density of INL nuclei decreased because of the bipolar and horizontal cells (Fig. 11). Erythrocyte aggregation (sludging) in deep capillary loops can lead to worsening of nutrition for the dependent inner retinal layers.

Figure 12. Experimental group. Peripheral area of the retina. No dilated vessels. Reduced density of neurons and increase in their mean area. Semi-thin section. Staining with methylene blue and fuchsin.

The total density of the retina in peripheral zones decreased up to 133±11.2 µ (Fig. 8), as well as the tendency of contraction of choroidal and retinal vessels was also seen. Most of the capillaries were occluded with erythrocyte aggregates.

Thus, microgravity modeling induced a series of interrelated processes in the retina of experimental rats manifesting in decreasing of total and layer-by-layer thickness of the retina, mild edema, and consequently, the increasing of specific density of neurons. These changes were more visible in the inner nuclear and ganglion cell layers.

In the inner nuclear layer, the cell density decreased because of the amacrine and bipolar cells.

In opposite the INL thinning, the nuclei of Muller cells retained their position and were optically “brought” close to
the IPL. The size of ganglion cell layer neurons was increased due to the cell edema. These changes could be explained by relative ischemia of the retinal layers, which are dependent on the retinal vessels circulation system.

The latter is characterized by constriction of the large arterial vessels, dilation of veins, and erythrocyte aggregation in microvessels impairing transcapillary blood flow. It is evident that the retinal circulation system is less resistant to gravity factors than the choriocapillary bed. Long-term exposure to factors causing ischemia in the inner layers of the retina, i.e. regions supplied via retinal vessels, can disturb processing of the primary signal from photoreceptors and impair highly organized and sensitive functions of the visual analyzer provided by bipolar and ganglion cells. This fact should be taken into account when planning visual tasks for astronauts.

Comparative morphometric analysis revealed significant differences in the thickness of the retinal pigment epithelium and photoreceptors in the central zone of the retina supplied with blood EXCLUSIVELY from the choroidal vessels. Hence, these areas are more hemodynamically vulnerable than equatorial and peripheral regions of the retina, where significant changes at the chosen exposure were recorded only in the outer nuclear and plexiform layers. The most pronounced changes (reduced thickness) were observed in the inner nuclear layer of the central, equatorial, and peripheral regions of the retina.

All morphometric differences in the both groups were statistically insignificant, transient, and temporary. Longer experimental exposure to the gravity factors can help to identify most susceptible elements of the visual analyzer. We hope to obtain these data in further experiments at different terms after microgravity modeling.

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