Changes in thickness of each layer of developing chicken cornea after administration of caffeine

Monika Kujawa-Hadryś, Dariusz Tosik, Hieronim Bartel

Department of Histology and Tissue Ultrastructure, Medical University of Łódź, Poland

Abstract: The aim of the study was the presentation of changes in thickness of each layer of a developing cornea, that came into being under an influence of caffeine which was administered to chicken embryos. Research materials were 26 chicken embryos from breeding eggs that had been incubated. Breeding eggs were divided into two groups: control (n=30) in which Ringer liquid was given, and experimental (n=30) in which teratogenic dose of caffeine was administrated – 3.5 mg/egg. In 36th hour of incubation solutions were given with cannula through a hole in an egg shell directly onto amniotic membrane. After closing the hole with paraffin, eggs were put back into incubator. On 10th and 19th day of incubation corneas were taken for morphometric and morphological analysis. In experimental groups reduction of corneal thickness, thickening of corneal epithelium and corneal endothelium as well as Bowman's and Descemet's membranes, decrease of thickness of corneal stroma in comparison with the control group have been observed. Caffeine causes thickness changes of all layers and decreases the total thickness of a developing cornea.

Key words: caffeine, cornea development, corneal thickness, chicken embryos.

Introduction

An eye is a complex structure deriving from different sources: the wall of the midbrain, surface ectoderm of the head and mesoderm of neural crest. In an eye development two important processes can be distinguish. First of them is a series of inductive signals leading to origins of most of eye structures. Second process consists of coordinate differentiation of those structures. Influence of adjoining eye elements is vital for correct visual reception [1-4].

Due to the generality of appearance (coffee, tea, chocolate, medicines) caffeine is the most commonly consumed psychostimulators.

In humans caffeine penetrates through placenta getting to amniotic fluid and umbilical cord blood. Next it is present in serum and urine of infant. Human infants have lower level of enzymes that are necessary for caffeine metabolism. Moreover, caffeine dissmission from blood of pregnant woman is delayed and caffeine concentration in a blood of infant is in balance with its accumulation in mother's blood. Caffeine dissmission in infants and embryos is also slowed down which makes, that the given dose of caffeine causes much more intensified effects in embryos than in adults. [5]. Caffeine clearance stays fundamentally unchanged during the first trimester of pregnancy, whereas, in the second, and third trimester decreases significantly. That is why half-life of caffeine in blood of pregnant women is four times longer than in those who are non-pregnant and this period lasts around 2.5 hours. Finally, it leads to an increase of caffeine concentration in blood of pregnant women [6-8].

Mechanism of a destructive caffeine effect on developing embryos is still hypothetical. Caffeine causes an increase of cAMP concentration inside a cell through phosphodiesterases inhibition, what may have an influence on a growth and development of embryo cells [9]. So far only few studies suggest destructive effects of caffeine on some structures of an eye development, especially on a cornea. Changes was specialy intensified in corneal anterior epithelium, stroma and endothelium [10].

The aim of the study was the presentation of changes in thickness of each layer of a developing cornea that came into being under an influence of caffeine which was given to chicken embryos.

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Table 1. Characteristics of the effectiveness of incubation in particular groups.

<table>
<thead>
<tr>
<th>Eggs/group</th>
<th>Embryos/group</th>
<th>Lived embryos</th>
<th>Dead embryos</th>
<th>No embryos</th>
</tr>
</thead>
<tbody>
<tr>
<td>C.10</td>
<td>15</td>
<td>9</td>
<td>8</td>
<td>1</td>
</tr>
<tr>
<td>E.10</td>
<td>15</td>
<td>7</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>C.19</td>
<td>15</td>
<td>8</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>E.19</td>
<td>15</td>
<td>7</td>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

Materials and methods

Chicken embryos. The research material was 26 chicken embryos from breeding eggs of ROSS 308 strand kept in incubator at the temperature 37-38°C and humidity 50-60%. 60 breeding eggs were used in the research. Effectiveness of chicken hatching eggs was about 50-60%. Breeding eggs were randomly divided into two groups. The first group was the control group. In 36th hour of incubation (9/10th stage of development according to Hamburger-Hamilton) [11] Ringer solution was injected into 30 eggs from that group. Next 30 eggs of the second, experimental group were given a teratogenic dose of caffeine – 3.5 mg/egg [12]. Solutions were given with a cannula through holes in eggs’ shells directly onto amniotic membrane. After closing the hole with paraffin, eggs were put back to incubator. On 10th and 19th day of incubation (36th and 45th stage of development according to Hamburger-Hamilton) corneas were taken for morphometric and morphological analysis. For investigation both right and left corneas only from the living embryos have been used. Living embryos were killed by decapitation. Eyeballs were removed from embryos. Corneas were taken with a use of microsurgical tools and stereo microscope. Specimens were prepared with methods used in electron microscopy. Whole corneas were fixed in a 3.6% solution of glutaraldehyde for 3 hours, next rinsed in 0.13 molar solution of cacodylic buffer and contrasted in 2% solution of osmium tetroxide. Then the taken tissues were dehydrated in alcohols of increasing concentration and in propylene oxide and embedded in epoxy resin – Araldite.

So prepared blocks, were cut in diameter plane with a diamond knife into the semithin sections. Afterwards specimens were stained with a toluidin blue solution and subjected to evaluation with optical microscope.

Morphometric analysis. Measurements were performed on three semi-thin cross sections of diameter area of each cornea. They were photographed by use of OLYMPUS BX41 microscope with 40× lens connected to digital camera OLYMPUS C-5060.

Each section was photographed three times. Depending on a group abundance 45 to 72 photographs of left and right cornea were taken. For the measurements 30 pictures from each group were randomly chosen.

The thickness of following layers of cornea were measured:
- anterior epithelium (AE),
- Bowman’s membrane (BM),
- corneal stroma (M),
- Descemet’s membrane (DM),
- endothelium (PE),
- cornea thickness (CT).

Thickness of each layer of cornea was measured with use of original software KIDNEY v.1.1. (Fig. 1). Boundaries of each layer were marked with optical mouse. Software algorithm assumes that a measurement of thickness is made if the angle between lines that mark boundaries of layers and section line, which length represents the distance between the boundaries, is equal 90°±3° [13]. Obtained results were saved in Microsoft Excel spreadsheet to be used in a further statistic analysis.

Statistical analysis. Statistical analysis was carried out with use of Statgraphics Centurion XV software. It covered the calculation of positional parameters as well as the comparison of each group distribution.

Following positional parameters were calculated: average values, standard deviation, coefficient of variation, mode, median, randomness evaluation of the taken sample. Evaluation of normal distribution was performed by use of both χ² and Kolmogorow-Smirnow tests. Comparison of the distribution between the groups was carried out with use of non-parametric Mann-Whitney U-test for non-related samples. The differences were treated as significant at the level p<0.01.

Results

On 10th day of incubation in the control group C.10, 1 dead embryo was observed, while in the respective experimental group E.10 – 2. On 19th day of incubation the number of dead embryos in the control C.19 and experimental group E.19 was 1. Characteristic of examined groups is in table 1.

No changes of macroscopic structure and size of the living embryos in each group were recorded. Collected on the respective days of incubation corneas of the experimental groups did not show macroscopic differences in comparison to respective control groups. At the light microscopic level, marked reduction in total corneal thickness in the experimental group, compared with control groups, both on 10th and 19th day of incubation was noted (Fig. 2 A,B,C,D). This observation was confirmed with morphometric evaluation, made by the "KIDNEY" software. The total corneal thickness increased with the development. On the 10th day of incubation it was lower than in 19th day of incubation in all groups. General composition of corneal layers was similar in all groups in the respective days of incubation.
In all groups, the anterior epithelium was stratified squamous. Under it Bowman's membrane was present. The corneal stroma was the thickest layer of the cornea. Increased amount of collagen fibers and reduced number of cells in the corneal stroma during the development were observed. Between corneal stroma and the corneal endothelium was Descemet's membrane.

On 10th day an average thickness of epithelium in the control group was 13.00 μm (4.72% of entire cornea thickness), Bowman's membrane 2.54 μm (0.91%), stroma 250.06 μm (90.83%), Descemet's membrane 3.30 μm (1.21%), endothelium 5.10 μm (1.83%), entire 275.89 μm.

On 10th day an average thickness of epithelium in the experimental group was 14.33 μm (11.57% of entire cornea thickness), Bowman's membrane 3.87 μm (4.33%), stroma 101.38 μm (78.20%), Descemet's membrane 3.05 μm (2.39%), endothelium 4.46 μm (3.19%), entire 129.02 μm.

On 10th day an average thickness of epithelium in the experimental group was 14.33 μm (11.57% of entire cornea thickness), Bowman's membrane 3.87 μm (4.33%), stroma 101.38 μm (78.20%), Descemet's membrane 3.05 μm (2.39%), endothelium 4.46 μm (3.19%), entire 129.02 μm.

On 19th day an average thickness of epithelium in the control group was 42.36 μm (10.77% of entire cornea thickness), Bowman's membrane 7.33 μm (1.64%), stroma 334.97 μm (84.86%), Descemet's membrane 3.73 μm (1.01%), endothelium 6.49 μm (1.65%), entire 394.48 μm.

On 19th day an average thickness of epithelium in the experimental group was 47.91 μm (18.62% of entire cornea thickness), Bowman's membrane 6.21 μm (2.45%), stroma 185.12 μm (75.68%), Descemet's membrane 2.85 μm (1.22%), endothelium 7.14 μm (2.86%), entire 273.92 μm.

Showed no significant differences in the thickness of the layers of the cornea of the eye left and right, both from the same individual as well as within research groups.

Tables 2 and 3 present thickness of each layer and the entire thickness of cornea in both groups and percentage of the various elements of the cornea in relation to its thickness.

**Discussion**

Administration of caffeine may generate disorders both in an organ of vision and a whole organism. From several dozen years it has been known, that high doses of caffeine have a negative influence on both women and laboratory animal’s fertility. Caffeine is a factor of intrauterine growth retardation (IUGR). Relation between caffeine consumption and a risk of preterm birth and low birth weight is not still clear [14-16].
Nevertheless, caffeine consumption together with smoking and alcohol drinking may have an influence on birth weight [17]. Moreover, synergism between caffeine and smoking or alcohol drinking significantly increases the number of spontaneous abortion [18]. It has also been shown, that drinking of two cups of coffee may lead to decrease of blood flow through placenta. Vessels contraction induced by caffeine, leads to tissues necrosis, due to hypoxia and ischemia [19]. Caffeine may, as well, influence behavior of healthy embryos in third trimester, especially breathing activity but also sleep and motor abilities [20].

During prenatal period caffeine may lead to improper development of many organs causing for example delayed neural tube closure [21], craniofacial and limb malformations, cleft palate [22,23], cardiovascular system malformations [12], the thymus involu- lution and defects of skeletal system [24].

In organ of vision caffeine causes an increase of intraocular pressure not resulting in changes in out- flow through the trabecular meshwork [25], having, however, an influence on epithelial cells of ciliary body [26]. It also causes multiple decrease of ocular blood flow [27]. Moreover, it increases blood vessel resistance and decreases blood flow in the human optic nerve head and choroid-retina, what may lead to disorders of retina development [28]. Few research that have been made, indicate a damaging effects of caffeine on some of the structures of a developing eye, particularly on cornea. Caffeine administered to pregnant rats during organogenesis (9th – 21st day of pregnancy) does not cause macroscopic changes in embryos’ corneas, but morphological changes could have been observed in some of the corneas. Changes were specially intensified in the group which had been given the highest dose of caffeine (100 mg/kg/d) and they concerned multilayered endothelial cellular proliferation with hyperchromasia and polymorphism, increased stromal cells mitotic activity, focal morphological changes, and corneal swelling in the injured segments with irregular and widely separated destruc-
This is why, restricted caffeine consumption, to 300 mg/d which is an equivalent of 2-3 cups of coffee, is advised during pregnancy. Not only does it refer to pregnant women, but also breast-feeding mothers, since the caffeine is also present in milk [29,30].

The results of the following research indicate a negative influence of caffeine on corneal development. Conclusions from that experiment may give a new vision on patomechanism of innate opacification of the cornea.

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References

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