Comparative studies on the corneal structural adaptation of two rodents inhabiting different environments

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Abstract

The corneas of Rattus rattus and Meriones shawi are composed of three main layers; an outer epithelium, a middle stroma (basement membrane, total stroma and Descemet’s membrane) and an inner endothelium. The mean thickness of the epithelium, total stroma, Descemet’s membrane, and endothelium was about 52 ± 7.3 μm, 275 ± 18.7 μm, 5.5 ± 0.7 μm, and 7.5 ± 0.23 μm in R. rattus, whereas it was 38 ± 5.8 μm, 124 ± 4.7 μm, 4 ± 0.21 μm, and 4.2 ± 0.17 μm in M. shawi.

In R. rattus and M. shawi, the outermost cells of the corneal epithelium are mostly polygonal and hexagonal in shape with nearly regular borders and show a dense pattern of microplicae with different scatter electron that exhibits three and two polymorphic appearances, respectively. Type A: numerous light cells with dense microplicae; type B: many dark cells with a moderate density of microplicae, and type C: few dark cells with a less density of microplicae are found in R. rattus; whereas, types A and B are found in M. shawi.

In both investigated species, the epithelial cells are characterized by some structural components, such as glycocalyx, fibrous components and tight junction between these cells, to resist the impact of the external stressed factors and to protect the underlying tissue, as well as to maintain an excellent transparency of the cornea. Among these structures, the cytokeratin filaments are the major components of the cytoplasm of the corneal epithelial cells (basal, polygonal, wing and squamous cells). Actin filaments are also found in the corneal epithelial cells, but they are prominent within the apical epithelial cells.

In R. rattus, the stroma is formed of an outer lamellar zone and an inner lamellar one; the latter is thicker and characterized by its interfibrillar spaces between the bundles of wavy dissociated collagen fibrils, which are arranged in an orthogonal manner. In M. shawi, however, the stroma is formed of one lamellar zone of flattened bundles of highly wavy and branched collagen fibrils, which are composed of perpendicular fibrillae alternating with longitudinal ones.

In R. rattus, the SEM showed that the endothelial cell surfaces are slightly bulging with many blebs, whereas in M. shawi, it showed that the surfaces of the endothelial cells are flattened and nearly smooth.

In conclusion, the transparency of the cornea, may be highly attributed to the increase in the thickness of the stroma, the presence of stromal interfibrillar spaces and the case of the stromal swelling. These aforementioned features are found in the corneal stroma of R. rattus, which live in different habitats of varying degrees of density such as water and dry or humid air, whereas these features are lacking in M. shawi, which live only in arid zones.

Key words: Cornea, mammals, rodents, adaptation, environment, LM, SEM, TEM

Introduction

The transparent nature of the cornea and its importance in the visual pathway as a major refracting lens of the eye have intrigued the investigators in many different disciplines, and their studies have added immeasurable knowledge to the understanding of the cornea in health and disease of human (Svedbergh and Bill, 1972;
Comparative studies on the corneal structural adaptation of two wild species of rodents, the black rat, Rattus rattus (Family: Muridae), and theshaw’s jird, Meriones shawi (Family: Cricetidae), were used in this study. Rattus rattus were collected from Abou Rawash at Giza, whereas M. shawi were collected from Borg Al-Arab near Alexandria. The average lengths for R. rattus and M. shawi were as follows: for the head and body, they were 165.4 mm and 140.2 mm, whereas for the tail, they were 205.3 mm and 135.7 mm, respectively.

The head of each species studied, was sacrificed and both eyes were carefully and immediately enucleated and cleansed of extraneous tissues with fine forceps and iris scissors. The eyes were then hemisected with a razor blade and the central corneal cups were taken out under a dissecting microscope and prepared for light, transmission and scanning electron microscopy.

**Light and transmission electron microscopy (LM & TEM):**

The dissected central corneas were cut into small pieces and fixed in cold (4°C) 2.5 % glutaraldehyde in 0.2 M phosphate buffer (pH 7.4) for 2 hours and post-fixed (2 hours at 4°C) in OsO₄ for electron microscopy. The samples were then dehydrated, treated with propylene oxide, infiltrated and embedded in Epon 812. Semithin sections were cut with the RMC-MT7 ultramicrotome and stained with toluidine blue. The corneas were then examined and photographed using a Joel 1010 Transmission Electron Microscope at the Regional Center for Mycology and Biotechnology, Al-Azhar University.

**Scanning electron microscopy (SEM):**

The dissected corneas were cut into small pieces and fixed in 2 % glutraldehyde in 0.2 M phosphate buffer (pH 7.4) for 2 hours and post-fixed (2 hours at 4°C) in OsO₄ for electron microscopy. The samples were then dehydrated, treated with propylene oxide, infiltrated and embedded in Epon 812. Semithin sections were cut with a diamond knife and stained with uranyl acetate and lead citrate. The sections were then examined and photographed using a Joel 1010 Transmission Electron Microscope at the Regional Center for Mycology and Biotechnology, Al-Azhar University.

**Materials And Methods**

**Specimen collection and dissection:**

Ten adult species of both the black rat, Rattus rattus (Family: Muridae), and the shaw’s jird, Meriones shawi (Family: Cricetidae), were used in this study. Rattus rattus were collected from Abou Rawash at Giza, whereas M. shawi were collected from Borg Al-Arab near Alexandria. The average lengths for R. rattus and M. shawi were as follows: for the head and body, they were 165.4 mm and 140.2 mm, whereas for the tail, they were 205.3 mm and 135.7 mm, respectively.

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Ultrathin sections (silvery) were cut, using a diamond knife and stained with uranyl acetate and lead citrate. The sections were then examined and photographed using a Joel 1010 Transmission Electron Microscope at the Regional Center for Mycology and Biotechnology, Al-Azhar University.

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dehydrate- 2% paraformaldehyde in 0.1M phosphate buffer, pH 7.4 at room temperature for 60 minutes. They were then washed twice in buffered sucrose (0.1M phosphate buffer, 5% sucrose solution) for 5 minutes. The tissues were post-fixed in 2% osmium tetraoxide (OsO₄) with 2% phosphate buffer for 2 hours at 4°C. After fixation, the tissues were dehydrated in upgraded series of ethanol. They were further dehydrated in absolute ethanol : acetone (1:1) solution for further 30 minutes, then in absolute acetone for 3 - 10 minutes. The tissues were dried in CO₂ drying apparatus (Critical-Point Dryer, CPD 030) and mounted on stubs, then coated with gold in Sputter Coater (SCD 005). They were examined and photographed using a Philips Scanning Electron Microscope XL30 at the Department of Anatomy, Faculty of Medicine, Ain Shams University.

Statistical Analyses:
The standard deviation, statistical analyses and significant levels (P) for all measurements were made by using Student’s t-test.

Results
The mean thickness of the total corneas of both Rattus rattus and Meriones shawi measures about 334.5 ± 21.9μm and 166.2 ± 14.5 μm, respectively and table (1), shows a highly significantly difference (p < 0.001) of them.

Light microscopy:
The corneas of both species are composed of three main layers: an outer epithelium, a middle stroma (basement membrane, total stroma and Descemet’s membrane) and an inner endothelium (Figs.1-6). The mean thickness of the epithelium, total stroma, Descemet’s membrane, and endothelium is about 52 ± 7.3 μm, 275 ±18.7 μm, 5.5 ± 0.7 μm and 7.5 ± 0.23 μm in R. rattus, whereas it is 38 ± 5.8 μm, 124 ± 4.7 μm, 4 ± 0.21 μm and 4.2 ± 0.17 μm in M. shawi.

Table (1), shows that, in R. rattus and M. shawi, there are very highly significantly differences (P < 0.0001) in the mean thickness of both the total stroma and endothelium, a highly significantly difference (P<0.001) in the mean thickness of the Descemet’s membrane, and a significantly difference (P<0.05) in the mean thickness of the epithelium.

Epithelium: The epithelial layer of R. rattus and M. shawi overlies a thin and distinct stromal basement membrane; it is composed of a stratified squamous, non-keratinized epithelium (about 9 rows of cells). These cells are arranged as a basal monolayer of short columnar cells, an intermediate polygonal cells (about 4 cells thick), an outer layer of flattened polygonal cells (wing cells) of about 3 cells thick and an outermost layer formed of flattened squamous cells (one cell thick) (Figs.2 & 5). It must be noted that some dark epithelial cells have been observed in between the epithelial cells near the basal layer (Figs. 2 &5).

Stroma: In R. rattus, it consists of an outer thin (about 72 ± 12.5 μm) lamellar zone of less distinguishable parallel collagen fibrils, and an inner thick (about 203 ±17.6μm) lamellar zone of dissociated ones by many large interfibrillar spaces (Figs.2 & 3). However, in M. shawi, it is composed of distinct and compact collagen lamellar fibrils, which are regularly arranged as parallel fibrils to each other in a wavy and branched manner (Figs. 5 & 6).

In R. rattus, few elongated keratocytes are observed scattering in the whole stroma (the outer and inner lamellar zones), however, in M. shawi, these cells are more distinguishable, great in number and scatter in between the lamellae of the stroma (Figs.2,3,5 & 6). The stroma of both species lies above a moderately thick and distinct Descemet’s membrane (Figs. 3 & 6).

Endothelium: In R. rattus and M. shawi, the endothelium is the innermost layer bordering the anterior chamber of the eye and is formed of a single layer of ill-defined low cuboidal cells underlying the Descemet’s membrane (Figs.3 & 6).

Scanning and transmission electron microscopy:
Epithelium: In R. rattus and M. shawi, the SEM shows that the outermost cells of the corneal epithelium are mostly polygonal and hexagonal in shape with nearly regular borders and show a dense pattern of microplicae with different scatter electron that exhibits three and two polymorphic
Comparative studies on the corneal structural appearances, respectively. Type A: numerous light cells with dense microvilli; type B: many dark cells with a moderate density of microvilli, and type C: few dark cells with a less density of microvilli are found in *R. rattus*; whereas, types A and B only are found in *M. shawi* (Figs. 7 & 8). It must be noted that minute microholes have been observed on the surfaces of the apical epithelial cells of *R. rattus* (Fig. 9), as well as on those of *M. shawi*.

The TEM shows that, in both investigated species, the outermost apical cells possess numerous and minute microvilli, which are coated with extensive glycocalyx to the extent that the microvilli can not be easily seen. Moreover, the so termed wing cells of the outer layer possess lateral thin wing-like extensions from the main bulks of the cell bodies (Figs. 10 a & b). The basal cell layer appeared to be the most active mitotically epithelial cells (Fig. 11). It must be noted that the major components of the cytoplasm of the corneal epithelial cells (basal, polygonal, wing and squamous cells), are the cytofilaments (Figs. 12 & 13). Actin filaments are also found in the corneal epithelial cells, but they are prominent within the apical epithelial cells (Figs. 10a & b). Moreover, microtubules are also present in the corneal epithelium, but they are prominent within the mitotic basal cells (Figs. 12 & 13). Besides, the corneal epithelial cells possess a sparse accumulation of mitochondria, Golgi bodies, endoplasmic reticulum and small vesicles. The membranes of the epithelial cells are highly interdigitated. Desmosomes are prominent cell-to-cell adhesion junctions along the cell borders. Gap junctions are also found between the lateral membranes of the wing cells (Figs. 10 a & b). The basal cells of the epithelium adhere to their basement membrane and the underlying fibrils of the stroma through an adhesion complex. A foam layer of newly synthesized collagen fibrils appeared in the interface of the epithelium and the upper part of the stroma (Fig. 13). In *R. rattus* and *M. shawi*, no Bowman’s layer was observed under the basement membrane.

Stroma: The stromal layers, of both investigated species, are nearly similar in their general arrangement, but differ in their thickness and composition.

In *R. rattus*, the stroma is formed of outer and inner lamellar zones which differ in their thickness (Table 1). The outer lamellar zone occupies about the anterior third of the stroma and has wavy and branched collagen fibrils. On the other hand, the inner lamellar zone occupies the main posterior part of the stroma and is characterized by its interfibrillar spaces between the more wavy and dissociated bundles of collagen fibrils, which are arranged in an orthogonal manner i.e. at right angle to another (Figs. 13 and 14 a & b).

In *M. shawi*, the stroma is formed of one lamellar zone. It is composed of flattened bundles of highly wavy and branched collagen fibrils, especially those near the epithelium. The lamellar stroma is composed of perpendicular fibrillae alternating with longitudinal ones (Figs. 5, 6, 15 & 16).

The lamellar stroma, of both studied species, is secreted and maintained by the keratocytes, which are resided between the fibrils and each possesses a large elongated nucleus, Golgi bodies and endoplasmic reticulum. These keratocytes are great in number and size in *M. shawi* than those of *R. rattus*, while they are highly vesiculated in *R. rattus*, than those of *M. shawi*, which possess prominent granular endoplasmic reticulum (Figs. 14a & 16).

The Descemet’s membrane, of both investigated species, is a regular, dark and exaggerated layer, which is synthesized by the endothelium and serves as a substratum for the endothelial cells. It is largely formed of parallel and very fine collagen fibrils (Figs. 14b & 15).

Endothelium: In *R. rattus* and *M. shawi*, the corneal endothelium is a single layer of low cuboidal cells, which appear by the SEM as irregular and flattened polygonal cells. In *R. rattus*, the surfaces of the endothelial cells appear slightly bulging with many blebs, however in *M. shawi*, their surfaces appear more flat and nearly smooth (Figs. 17 & 18). In both investigated species, the endothelial cells have large and flattened nuclei, a number of mitochondria, endoplasmic reticula and Golgi bodies. In *R. rattus*, the endothelial cells are characterized by the presence of a small
number but prominent large vacuoles, as well as great numbers of vesiculated structures of the granular endoplasmic reticulum, however in \textit{M. shawi}, the cytoplasm of the endothelial cells is highly granulated with some scattered vesiculated structures of the endoplasmic reticulum (Figs. 19 & 20).

Table 1: The mean thickness of the different corneal layers of the black rat, \textit{Rattus rattus}, and the shaw’s jird, \textit{Meriones shawi}.

<table>
<thead>
<tr>
<th>Corneal layers</th>
<th>\textit{Rattus rattus} Mean thickness (μm) ± SD</th>
<th>\textit{Meriones shawi} Mean thickness (μm) ± SD</th>
<th>\textit{P}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total cornea</td>
<td>334.5 ± 21.9</td>
<td>166.2 ± 14.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Epithelium</td>
<td>52 ± 7.3</td>
<td>38 ± 5.8</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Total stroma</td>
<td>275 ± 18.7</td>
<td>124 ± 4.7</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>O L Z</td>
<td>72 ± 12.5</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>I L Z</td>
<td>203 ± 17.6</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Descemet’s membrane</td>
<td>5.5 ± 0.7</td>
<td>4 ± 0.21</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Endothelium</td>
<td>7.5 ± 0.23</td>
<td>4.2 ± 0.17</td>
<td>&lt; 0.0001</td>
</tr>
</tbody>
</table>

N.B. \textit{P} < 0.05 (significant)  
\textit{P} < 0.001 (highly significant)  
\textit{P} < 0.0001 (very highly significant)  
O L Z = Outer lamellar zone of the stroma  
I L Z = Inner lamellar zone of the stroma

Explanation Of Figures

Figs. 1 - 3 & 4 - 6: Photomicrographs of vertical semithin sections of the corneas of \textit{Rattus rattus}, and \textit{Meriones shawi}, respectively, showing the general structure of the cornea (the epithelium: Ep; basement membrane: B; stroma: S; keratocytes: K; Desemet’s membrane: D; and endothelium: En). Toluidine blue.

Figs. 1 & 4: Show the structure of the cornea and the thickness of its different layers. Notice the interfibrillar spaces (asterisks) and the outer and inner lamellar zones (OLZ and I LZ) in figure 1.(X 138).

Figs. 2 & 5: Show the structure of the epithelial layer (Ep), the dark cells (arrows), the wing cells (asterisks), the basement membrane (B) and the underlying stroma (S). Notice also the differences between the stroma of both species, the keratocytes (K), and the branched collagen fibrils in figure 5 (white arrows). (X 435).

Figs. 3 & 6: Show the structure of the stroma (S), the distinct Descemet’s membrane (D) and the endothelium (En). Notice many and large interfibrillar spaces (asterisks) in between the lamellae of the inner lamellar zone (ILZ) and the keratocytes (K) of the stroma of \textit{R. rattus} (Fig. 3). Also notice the distinct lamellar stroma with distinct keratocytes (K) of \textit{M. shawi} (Fig. 6). (X 435).

Figs. 7 & 8: Scanning electron micrographs of the corneas of \textit{R. rattus} and \textit{M. shawi}, respectively, showing the different types of the apical epithelial cells.

Fig. 7: Shows type A, B and C cells. Notice the nearly regular borders of the cells. (Scale bar = 20 μm)

Fig. 8: Shows type A and B. Notice the slightly regular borders of the cells. (Scale bar = 10 μm)

Fig. 9: Scanning electron micrograph of the cornea of \textit{R. rattus}, showing the microholes (arrows) on the surfaces of the apical epithelial cells (Scale bar = 2 μm)

Figs. 10a &b: Transmission electron micrographs of the corneas of \textit{R. rattus} and \textit{M. shawi}, respectively, showing the structure of the epithelial cells.

Fig. 10a: Notice the surface of the apical cell is coated with extensive glycocalyx (arrows). Notice also the wing cells (asterisks), the tight adhesion between
the epithelial cells (short arrows) and the gap junctions (arrowheads).
  (Scale bar = 2 μm)

**Fig. 10b**: Notice the microplicae (Mp), extensive glycocalyx (arrows) and extensive dark fibrous filamentous structures (white arrows) in the outer cells. Notice also the wing cells (star) and the tight adhesion (short arrows).
  (Scale bar = 2 μm)

**Fig. 11**: Transmission electron micrograph of the cornea of *R. rattus*, showing the structure of the epithelial cells. Notice the mitotically active nuclei (N) of the basal cells that rest on a very thin basement membrane (B) and overlying less distinguishable fibrils of the outer lamellar zone (OLZ) of the stroma (S). Notice also the dark cells (asterisks) and many cytoplasmic vacuoles (V).
  (Scale bar = 10 μm)

**Fig. 12**: Transmission electron micrograph of the cornea of *M. shawi*, showing the microtubules (arrows) in an epithelial cell. Notice also extensive fine cytokeratin filaments (short arrows).
  (Scale bar = 500 nm)

**Fig. 13**: Transmission electron micrograph of the cornea of *R. rattus*, showing the basal cells that rest on very thin basement membrane (B) and overlying less distinguishable fibrils of the outer lamellar zone (OLZ) of the stroma (S) and the foam appearance (star). Notice the microtubules (arrows), fine cytokeratin filaments (short arrows). Notice also the adhesion complex (white arrows).
  (Scale bar = 2 μm)

**Figs. 14a & b**: Transmission electron micrographs of the cornea of *R. rattus*, showing the orthogonal arrangement of the collagen fibrils (arrows) of the inner lamellar zone (ILZ) and an elongated keratocyte (K) with extensive vacuoles (V) and a large nucleus (N). Notice the interfibrillar spaces (asterisks).
  (Scale bar = 10 μm)

**Figs. 15 & 16**: Transmission electron micrograph of the cornea of *M. shawi*, showing the structure of the lamellar stroma.

**Fig. 15**: Shows the wavy appearance of the lamellar stroma (S) and an elongated keratocyte (K), with a large nucleus (N). Notice the structure and relation of the Descemet’s membrane (D) with the stroma and endothelium (En).
  (Scale bar = 2 μm)

**Fig. 16**: Shows the arrangement of the perpendicular (dark star) and longitudinal (white star) collagen fibrillae. Notice the structure of the keratocyte (K) with its prominent granular endoplasmic reticulum (ER).
  (Scale bar = 500 nm)

**Figs. 17 & 18**: Scanning electron micrographs of the corneas of *R. rattus* and *M. shawi*, respectively, showing the irregular polygonal endothelial cells, and the nature of their surfaces.

**Fig. 17**: Shows the irregular cell borders and the slightly bulging surfaces. Notice the small blebs on the surfaces.
  (Scale bar = 20 μm)

**Fig. 18**: Shows the irregular cell borders and the smooth surfaces of the endothelial cells. (Scale bar = 20 μm).

**Figs. 19 & 20**: Transmission electron micrographs of the corneas of *R. rattus* and *M. shawi*, respectively, showing the structure of the endothelium (En) and its relation to the Descemet’s membrane (D).

**Fig. 19**: Shows the highly vesiculated structure of the endothelial cells. Notice a large vacuole (asterisk) and the nucleus (N).
  (Scale bar = 2 μm)

**Fig. 20**: Shows the highly granulated cytoplasm of the endothelial cells, scattered vacuoles (V) and mitochondria (M).
  (Scale bar = 2 μm).
Discussion

The black rat, *Rattus rattus*, and the shaw’s jird, *Meriones shawi*, are two rodents living in two different habitats; the first species inhabits the Nile Valley, countries, houses, tunnels, stores...etc, whereas the second species inhabits the rocky desert (Wassif, 1995). These habitats have characteristic climatic factors affecting the structure of the different organs of their animals. Temperature is considered as one of the main factors affecting the structure of the skin and cornea. Consequently, the cornea of the desert animals (*M. shawi*) is subjected to a great effect of the high temperature rather than that of *R. rattus* inhabiting the Nile Valley of the moderate temperature. In spite of these two species are related to order Rodentia and suborder Myomorpha, they are apparently have some morphological and structural differences of their corneas according to the mode of life and type of habitat.

The general corneal structure of both, *R. rattus* and *M. shawi*, is nearly similar to each other, as it is composed mainly of three layers: the epithelium, stroma (basement membrane, total stroma and Descemet’s membrane) and endothelium. Such general structure is somewhat similar to what has been described for the corneas of most vertebrate classes including human and other mammals (Smolin and Thoft, 1994; Beuerman and Pedrosa,1996; Krachmer *et al.*, 1997; Chakravarti, 2001; El-Dawi, 2002 & 2004; Svaldeniene, *et al.*, 2003).

Epithelium

The epithelial layer of *R. rattus* and *M. shawi* is composed of a stratified squamous, non-keratinized epithelium (about 9 rows of cells), which are arranged as a basal monolayer of short columnar cells, an intermediate polygonal cells (about 4 cells thick), an outer layer of wing cells (about 3 cells thick), and an outermost layer of flattened squamous cells (one cell thick). Similar arrangement and number have been described in the knockout mice (Chakravarti, 2001). However, Gibson (1994) stated that the mammalian corneal epithelium is composed of 5 - 7 cell layers arranged as basal columnar cells, one to three cell layers of intermediate wing cells and an outer cell layers of three to four flattened squamous cells. Also, Krachmer *et al.* (1997) described 5 – 6 layers of three corneal epithelial cell types of mammals including human; these types were monolayer basal columnar cells, two to three layers of wing cells and two to three layers of superficial cells. In the present investigation, the increase in the number of the epithelial layers and types of cells, most probably, for protection the underlying stroma from the extreme external factors which are highly correlated to the variable and hard habitats in which they live.

In *R. rattus* and *M. shawi*, the SEM shows that the outermost corneal epithelial cells are mostly polygonal and hexagonal in shape with nearly regular borders. They possess microholes and show a dense pattern of microriplicae. Such appearance simulates that found in human and most studied mammals (Gibson, 1994; Doughty, 1996; Krachmer *et al.*, 1997; Collin and Collin, 2000). In the rhino mouse (Amemiya, *et al.*, 1980) and in the rat (Takami *et al.*, 2004), the surfaces of the outermost corneal epithelial cells possess microvilli, microplicae and microholes which possess elevated rims formed by large folds of the apical epithelial cells. In human (Gibson, 1994) the apical epithelial cells possess numerous microplicae and microvilli, which are suggestive of the epithelium normal function.

In *R. rattus* and *M. shawi*, the dense pattern of microplicae shows different scatter electron that exhibits three and two polymorphic appearances, respectively; type A: light cells with dense microplicae; type B: dark cells with a moderate density of microplicae, and type C: dark cells with a less density of microplicae. In the white rabbit (Doughty, 1996) three cell types of the corneal epithelium (light, medium and dark), were distinguished. Also, Gibson (1994) demonstrated that the mammalian apical epithelial cells show three types of
cells (light, moderate and dark) according to the degree of the scatter electron of the microvilli. However, two types of cells, dark and light were observed in the corneal epithelium of the piglets (young pigs). These two types not observed in young dogs and adult dogs and pigs (Svaldeniene, *et al.*, 2003). Moreover, microvilli and microvilli and microvilli have been observed on the rat corneal epithelium (Takami *et al.*, 2004). On the other hand, in different mammals, Krachmer *et al.* (1997) distinguished two types of corneal epithelial cells, large dark cells and small light ones. The dark cells are covered with dense microvilli, whereas the light ones have fewer microvilli. Most probably, the presence of various types of the apical epithelial cells according to their microprocesses represents stages of developmental processes, as the light cells are the most recent, whereas the dark cells are the oldest ones. Moreover, these microprocesses, microvilli, micro in microvilli or even microridges, most probably, are highly correlated to the external environments and their demands, as they enlarge the total cell surface area to allow the active exchange of oxygen and nutrients between the cells and tear fluid.

In the present investigation, the epithelial cells of both investigated species are characterized by some structural components, such as glycocalyx, fibrous components and tight junction between these cells, to resist the impact of the external stressed factors and to protect the underlying tissue, as well as to maintain an excellent transparency of the cornea.

In *R. rattus* and *M. shawi*, the microvilli of the outermost apical cells were coated with extensive glycocalyx to the extent that they can not be easily seen. This is also observed in the guinea pigs and rats (Gibson, 1994; Svaldeniene, *et al.*, 2003). Gibson (1994) described that the glycocalyx is loosely associated with the tear film layer, as well as with the mucin and tear film spreading over the surface of the eye. Moreover, Nishida (1997) stated that glycocalyx is structurally formed of floating particles of glycolipid and glycoprotein molecules in the cell membrane and covered by oligosaccharides. He suggested that this structure maintains the hydrophilic properties of the epithelial cells. Most probably, the presence of extensive glycocalyx, in the species of the present investigation, is apparently highly correlated to retain the tear fluid for long periods as possible for protection against microbial or fungal infections and the different habitats of *R. rattus* or maintain the stability of the tear fluid under the stress of high temperature and preventing their eyes from drying as in the case of *M. shawi* which inhabits the rocky desert.

In *R. rattus* and *M. shawi*, the presence of the cytokeratin filaments in the cytoplasm of the corneal epithelial cells, as well as the actin filaments especially within the outermost epithelial cells was greatly similar to what have been reported in human and many mammalian species by Gibson (1994) and Nishida (1997). The latter author stated that the wing cells are rich in K45 keratins which are specific to non-keratinized corneal epithelial cells. In the knockout mice, Chakravarti (2001) stated that keratins (K1–K20) form the intermediate filaments of the epithelial cells. Most probably, the extensive appearance of such fibrous components in the corneal epithelial cells of *R. rattus* and *M. shawi*, reflects their stiffness as a supportive cells against external stress. On the other hand, the highly interdigitation, the presence of extensive desmosomes and gap junctions between the corneal epithelium, especially the wing and intermediate cells, of the present investigation, have been also described in some mammalian species (Gibson, 1994; Smolin and Thoft, 1994; Nishida, 1997; Svaldeniene, *et al.*, 2003). Most probably, the presence of these junctional complexes reflects the tight junctions of these cells. Such tight junctions have a mechanical barrier function and prevents the penetration of the exterior aqueous material easily to the corneal stroma, as in *R. rattus* that live in various habitats including the tunnels or from the stroma to exterior as in *M. shawi* that live in rocky desert of hot climate.

### Stroma

The presence or absence of the Bowman’s layer is still a matter of debate.
In *R. rattus* and *M. shawi*, no Bowman’s layer was observed under the basement membrane. It has been found that most species of mammals do not have the Bowman’s layer (Gibson, 1994; Smolin and Thoft, 1994; Krachmer et al., 1997). However, Nishida (1997) stated that this layer is observed in human and certain other mammals including the primates. Moreover, Hayashi et al. (2002) and Svaldeniene, et al. (2003) found that the Bowman’s layer is thick and well developed in higher mammals such as dogs, cattle and human; thin and ill-developed in the mouse, rat, guinea pig, rabbit and cat. The functional significance of the Bowman’s layer remains a matter of discussion. Many authors suggested that it plays an important role in the maintenance of the epithelial structure, for maintaining epithelial uniformity, thus forming an appropriate refractive power or preventing the contact between the epithelium and stroma (Gibson, 1994; Smolin and Thoft, 1994; Krachmer et al., 1997; Nishida, 1997). In fact, the presence or absence of the Bowman’s layer is greatly dependent on the epithelial structure in its ability to allow or prevent the passage of aqueous material to or from the stroma. This is apparently clear in *R. rattus* and *M. shawi*, who lack the Bowman’s layer as their epithelial cells are formed of many layers and contain prominent cytokeratin and actin filaments, as well as the presence of tight junctions between these cells.

The transparency of the cornea is still subjected to different hypotheses. Nishida (1997) stated that the regular arrangement of the collagen fibrils, in mammals and human, contributes to the corneal transparency and any disturbance in the uniformity of the interfibrillar spaces cause loss of transparency. Freud et al. (1995) suggested that, in human and rabbit, the corneal transparency is the result of the small size of the constituent collagen fibrils, and the lamellae of the posterior stroma are more regular in arrangement than those of the anterior stroma. Most probably, the transparency of the cornea is highly attributed to the increase in the thickness of stroma, presence of interfibrillar spaces and the case of the stromal swelling. The aforementioned features are found in the corneal stroma of *R. rattus*, which live in different habitats of varying degrees of density such as water and dry or humid air as compared with *M. shawi*, which live only in arid zones and lack these features.

**Endothelium**

In *R. rattus* and *M. shawi*, the SEM showed that the corneal endothelial cells were flattened polygonal cells with irregular borders. In most vertebrates including human and other mammalian animals such as rabbits, rats, mice, Guinea pigs and dogs, the corneal endothelial cells are a mixture of hexagonal and pentagonal shaped cells in which their borders are irregular and interdigitating (Gibson, 1994; Smolin and Thoft, 1994; Krachmer et al., 1997; Nishida, 1997; Collin and Collin, 1998). However, Svaldeniene, et al. (2003) stated that the endothelial cells of the adult pig were square shaped; whereas in aging human they are not consistently hexagonal in shape, but rather pleomorphic in shape. In *R. rattus*, the SEM showed that the endothelial cell surfaces were slightly bulging with many blebs, however in *M. shawi*, their surfaces were flattened and nearly smooth. Collin and Collin (1998) observed microvilli on many endothelial cells of human, rat, Guinia pig, dog and rabbit. However, a single primary cilium was a normal component of the rabbit endothelial cells (Gallagher, 1980) and was occasionally found in the endothelium of human (Beuerman and Pedroza,1996). Gordon et al. (1983) stated that, the endothelial surfaces in quiescent rat and rabbit were devoid of microvilli but display globular projections, however, the regenerating endothelial cells possess microvilli and filopodia. Yamaguchi et al. (1992) revealed several changes during the development of the rat corneal endothelial surfaces. These changes appeared as mushroom-like microvilli at 1 month-old, numerous filamentous microvilli at 3 months-old and numerous slender microvilli at 6 months-old.

Most probably, such controversies in the surface structures either microvilli, cilia, globular projections, blebs and filopodia in the rats and rabbits were related to the mitotic activity of the endothelial cells.
which are highly affected by their direct exposure to the surrounding aqueous humor of the eye. The TEM showed the presence of a small number but prominent large vacuoles, great numbers of vesiculated structures of the granular endoplasmic reticulum and mitochondria in the endothelial cells of R. rattus; however, in M. shawi, highly granulated cytoplasm with some scattered vesiculated structures of the endoplasmic reticulum were observed. Such vesiculated structures of the endoplasmic reticulum were reported in many experimental mammals (Gibson, 1994; Nishida, 1997; Svaldeniene, et al., 2003). However, the presence of large vacuoles, highly vesiculated structures and mitochondria in R. rattus than those of M. shawi, most probably, is related to the high activity of these cells in the former species and their involvement in secretion, as well as active transport. Thus, it is not surprising that their primary function is to nourish and hydrate the cornea. This interpretation is highly logic in species inhabiting areas with continuous changeable environments as in R. rattus of the present investigation.

References


دراسات مقارنة للتكيف التركبي للقرنية لاثنين من القوارض في بيئات مختلفة

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في هذا البحث وجد أن القرنية في كل من الجرذ الأسود ومورينز شاوي تتكون من عدة طبقات وهي الطالنلخارية والإستروما الوسطى والطلانية الداخلية. وقد تم حساب متوسط سمك هذه الطبقات في كل منها.

وقد أوضحت الدراسة الميكروسكوب الماجس في كل من الجرذ الأسود ومورينز شاوي أن الخلايا السطحية للطبقة الطلانية الخارجية تكون عددية الأضلاع و سساسية من الشكل ومكتظة بالتحفيز. وتتميز هذه الخلايا ويوجد نظام تكثيف من البروتيك الديفقة (ميكروبلوريا) ذات كثافات مختلفة. وقد أوضحت الميكروسكوب الماجس أيضاً عن وجود ثلاثة أنواع من الخلايا طبقًا لكثافة هذه الدوائر في الجرذ الأسود: نوع (أ) شديد الكثافة الفاتح ويتمثل النوع السائد، (ب) متوسط الكثافة الداكن و(ج) قليل الكثافة الداكن واقلهم.

عادة، بينما يوجد النوع (أ) و (ب) فقط في مورينز شاوي.

وقد وجد أن الخلايا الطلانية في كل من هذين القوارض تتميز وجود تراكيب معينة مثل جليكوكلابس، مكونات ليفية، خيوط سيتوكريناتينية وأكينية وتماسك شديد بين الخلايا لمقاومة تأثير العوامل الخارجية وحماية الأنسجة الداخلية بالإضافة إلى الاحتفاظ بشفافية ممتازة للقرنية.

وقد أظهرت الدراسة أن طبقة (غدراو) في الجرذ الأسود تتكون من منطقتي الصالح الخارجي والداخلي وتميز الأخيرة وجود انتفاخات وفراغات بين ليفات الكولاجينية متموجة، ولعل هذا التركيب يساعد معيشة الجرذ الأسود في بيئات مختلفة الكثافة كالحمر أو الظفر أو الحرف. أما في مورينز شاوي والذي يعيش في المناطق الجذب والتي تميز بالحرارة الشديدة، فإن طبقة ستروما تتكون من منطقة صائحة واحدة من حزم شديدة التموج وتتفرع إلى ليفات كولاجينية والتي تتكون بدورها من ليفات متعامدة على أطراف طولية.

أما الطلانية الداخلية في الجرذ الأسود فإن أسحة خلاياها يكون ذات إنبعاثات خفيفة ومموجة بعدة انتفاخات زوينية، أما في مورينز شاوي فإن أسحة الخلايا الطلانية الداخلية تكون مسطحة وملساء.

وقد أُنتجت من هذا البحث أن ترتيب القرنية في الجرذ الأسود يمكنه من المعيشة والتكيف في بيئات متباينة الحرارة ومختلفة الكثافة كالحمر الظفر أو الظفر. أما في مورينز شاوي فإن ترتيب القرنية تلائم الحياة في الصحراء والتي تميز بالجفاف وشدة الحرارة.