Swelling studies of camel and bovine corneal stroma

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Abstract: In the present study we investigated the swelling characteristics of fresh camel and bovine cornea in sodium salt solutions. Swelling studies were carried out at 20 minutes, 14 hours, and 46 hours on five fresh camel and five fresh bovine corneas. During the 20-minute hydration of fresh corneal stroma was investigated using sodium chloride (NaCl), sodium bicarbonate (NaHCO3), sodium acetate (CH3COONa), sodium thiocyanate (NaSCN), and sodium fluoride (NaF) at 2-minute time intervals. During a 46-hour time period, the hydration study was carried out using NaCl (150, 300 mM) and NaF (150 mM) at random intervals. The 14-hour study was carried out to assess the rehydration of corneal stroma after 6 hours of drying. During the 20-minute swelling studies in the first 2 minutes the rate of hydration in both camel and bovine corneas was high but gradually reduced in the 2 –20-minute period. The rates and levels of hydration of camel and bovine cornea were not significantly different from each other in all the strengths of solutions. During the 46-hour swelling studies, the initial rate of hydration (0–2 hours) of camel and bovine stroma, in all solutions was significantly higher (Z = 0.056) compared to hydration during later hours (2–46 hours). Camel stromal hydration (high) in 150 mM NaCl was significantly higher compared to bovine stromal hydration in the same solution during the 10–24, and 24–46-hour time periods. Rehydration in camel stroma was significantly lower than bovine in 150 mM NaF. The 20-minute study showed that there was no selective affinity for particular ions in camel or bovine corneal stroma. Initial swelling in both corneal and bovine stroma is faster and more prominent compared to later swelling. The swelling in camel cornea is more prominent compared to bovine corneal stroma. This could be due to higher negatively charged keratin sulfate–proteoglycans in the stroma. Lower rehydration in camel cornea suggests stronger leaching of proteoglycans from stroma in NaF.

Keywords: camel, swelling, sodium thiocyanate, sodium bicarbonate, proteoglycans

The cornea and sclera together form the outermost covering of the eye and withstand both internal and external forces on the eye to maintain the shape of the eyeball and to protect the contents from mechanical injury.1,2 Approximately 90% of the cornea consists of stroma which contains collagen fibrils, proteoglycans (PGs) and keratocytes. The collagen fibrils have a uniform diameter and spacing and are organized in tightly packed, parallel-running lamellae. This specific arrangement of collagen fibrils is essential for corneal transparency.3

In birds and primates (including humans) corneal collagen fibrils (CFs) are heterotypic, comprising chiefly type I and type V collagen molecules.4,5 The cornea also contains large amounts of type VI collagen, which forms microfibrillar structures by lateral aggregation.6 PGs are macromolecular glycoconjugates consisting of specialized polysaccharide chains, glycosaminoglycans (GAGs), which are covalently attached to a protein core.6 In the cornea the extracellular matrix contains a class of small interstitial PGs known
as collagen-binding small leucine-rich repeat proteoglycans (SLRRPs), which carry two types of GAG side chains. The family members lumican, keratocan, and mimecan carry keratan sulfate chains, whereas decorin, biglycan, and versican carry chondroitin sulfate/dermatan sulfate chains. It is believed that these small leucine-rich repeat proteoglycans (SLRRPs) maintain the uniform organization of collagen fibrils, which is responsible for transparency of the cornea. Swelling of the cornea is believed to be due to an excess of water absorbed by the negatively charged proteoglycans, which results in nonuniform distribution of collagen fibrils.

Unlike other connective tissues, including sclera, the cornea is transparent. It is believed that the endothelium normally acts as a barrier to maintain the ionic composition of the stromal matrix and, more importantly, to maintain optimum hydration. The swelling of corneal stroma in vivo is not manifested because of the presence of a balancing outward directed bicarbonate ion pump located in the corneal endothelial lining at the posterior surface of the stroma.

When the stroma swells it loses its transparency. This is due to increased light scattering, which is thought to be caused by a nonuniform distribution of water and disruption of the collagen packing. Corneal swelling has been studied by placing the corneal stroma in distilled water and bathing solutions with different ionic strengths and pH levels.

Previous swelling studies have been carried out on bovine cornea, however, there has been no study carried out on the camel cornea. Camels are native to hot, sunny, and sandy hot air-blowing climates. Even though these climatic conditions cause dry eyes in most animals, camels have wet eyes. In the present paper we investigate the swelling characteristics of camel cornea and compare this with those of bovine cornea. To achieve our objective, we studied the free swelling characteristics between camel and bovine cornea. Additionally, a 14-hour study was carried out to assess leaching out of proteoglycans from the stroma to the bathing solutions. The difference between hydration and rehydration of the corneal stroma will be the leaching out of PGs from the stroma to the bathing solutions.

### 20-minute swelling study of stroma obtained from fresh eyes

Square pieces (8 × 8 mm side dimensions) of corneal stroma were air dried for 3–4 hours and weighed. Immediately after weighing, the samples were kept in 50 mL of bathing solution (Table 1) of NaCl (pH 7.4), NaCl (pH 8-8.4), CH₃COONa (pH 7.4), NaSCN (pH 7.4), or NaHCO₃ (pH 8-8.4) at 37.5 mM, 75 mM, 150 mM, 300 mM, or 600 mM concentrations separately. The wet weight of the sample was recorded at 2-minute intervals for a 20-minute time period. The samples were removed from the bathing solution and washed with distilled water for a period of 2–3 hours (to remove excess swelling) before drying and weighing.

<table>
<thead>
<tr>
<th>Solution</th>
<th>pH</th>
<th>Ionic concentration</th>
<th>Buffer</th>
<th>Study period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium chloride (NaCl)</td>
<td>7.4</td>
<td>37.5 mM, 75 mM,</td>
<td>HEPES</td>
<td>20 minutes;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>150 mM, 300 mM,</td>
<td></td>
<td>46 hours;</td>
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<td></td>
<td></td>
<td>and 600 mM</td>
<td></td>
<td>14 hours</td>
</tr>
<tr>
<td>Sodium bicarbonate (NaHCO₃)</td>
<td>8-8.4</td>
<td>37.5 mM, 75 mM,</td>
<td>none</td>
<td>20 minutes;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>150 mM, 300 mM,</td>
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<tr>
<td></td>
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<td>and 600 mM</td>
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</tr>
<tr>
<td>Sodium acetate (CH₃COONa)</td>
<td>7.4</td>
<td>37.5 mM, 75 mM,</td>
<td>HEPES</td>
<td>20 minutes;</td>
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<td>150 mM, 300 mM,</td>
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<td></td>
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<td>and 600 mM</td>
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<tr>
<td>Sodium thiocyanate (NaSCN)</td>
<td>7.4</td>
<td>150 mM and 300 mM</td>
<td>HEPES</td>
<td>20 minutes;</td>
</tr>
<tr>
<td>Sodium Floride (NaF)</td>
<td>7.4</td>
<td>150 mM and 300 mM</td>
<td>HEPES</td>
<td>20 minutes;</td>
</tr>
</tbody>
</table>
salt) and oven dried at 90–100°C for 24 hours. The dry weight of the sample was recorded and hydration calculated using the following formula:

\[
\text{Hydration} = \frac{\text{Wet weight} \ - \ \text{Dry weight}}{\text{Dry weight}}
\]

46-hour swelling study of stroma obtained from fresh eyes

Square pieces (8 x 8 mm side dimensions) of corneal stroma were air dried for 3–4 hours, and their weight was recorded. Immediately after weighing the samples were hydrated in NaCl (150 mM, pH 7.4), NaF (150 mM, pH 7.4), or NaF (300 mM, pH 7.4). The wet weight of the sample was recorded at irregular intervals. The samples were removed from the bathing solution and washed with distilled water for a period of 2–3 hours to remove excess salt and then oven dried at 90–100°C for 24 hours. The dry weight of the sample was recorded and hydration calculated by the aforementioned formula.

14-hour swelling studies of stroma obtained from fresh eyes

Square pieces (8 x 8 mm side dimensions) of corneal stroma were air dried for 3–4 hours, and their weight was recorded. Immediately after weighing the samples were hydrated for 14 hours in NaCl (150 mM, pH 7.4), NaCl (300 mM, pH 8-8.4), or NaF (150 mM, pH 7.4) separately and their wet weight monitored at 2-hour intervals. After 14 hours of swelling, the samples were air dried for 6 hours, weighed, then rehydrated in the freshly made bathing solution, as previously described. The samples were weighed at intervals of 2 hours for a total period of 14 hours. After that the samples were placed in water for 2–3 hours to remove excess salt and dried in an oven at 90–100°C for 24 hours. The dry weight of the sample was recorded and hydration and rehydration calculated by the formula described earlier.

In the present study, various concentrations of salt solutions were used to assess the variation in swelling of cornea. It has been reported that ‘Q’ (fixed charge of proteoglycans) is a function of pH which is dependent on the concentration of the bathing medium.26 The higher the concentration of bathing solutions, the lower the swelling of the cornea.

In previous studies it has been shown that swelling is rapid in the initial hour and slower in the later hours. Considering this information, intervals in the initial hours were chosen rather than later hours to observe the swelling trend in both camel and bovine cornea.

Swelling of the fresh sample is considered as hydration, whereas swelling of the air-dried sample after hydration is considered as rehydration. Nonparametric statistical tests were used to perform statistical analysis of the data and Excel was used for drawing graphs.

Results

Change in hydration over 20 minutes at 2-minute intervals

In both camel and bovine samples, the level of hydration was measured for 20 minutes in NaCl (pH 7.4, 8.4) (Figures 1A and 1B), NaHCO₃ (pH 8) (Figure 1C), and NaSCN (pH 7.4) (Figure 1D), CH₃COONa (pH 7.4) (Figure 1F) (monitored at 2-minute intervals). In general in all solutions, the swelling of both camel and bovine samples was initially fast but gradually slowed down. The level of hydration was also inversely proportional to the concentration of the solutions ie, the lower the concentration of bathing solution the higher the rate of hydration (Figures 1A to 5), but was overall not significant. Rates and level of hydration of camel and bovine samples were also not significantly different from each other in all the solutions (Table 1). The results showed that there was no selective affinity to any particular ionic solution.

Changes in camel and bovine fresh stromal hydration over a 46-hour period

Hydration in camel stroma

During the first 2 hours, the level of hydration in 150 mM NaCl (12.20), 150 mM NaF (10.31) and 300 mM NaF (9.31) was significantly higher (Z = 0.056) compared to hydration during later hours (2–46 hours) in these solutions (Figures 2A and 2B). There was no significant difference in hydration among the solutions from each other during the initial hours (0–2 hours). There was a significant difference in hydration between NaCl 150 mM and NaF 300 mM between 2–10 hours. The level of hydration of the corneal stroma increases gradually with increase in time (Figure 2A).

The total hydration of 150 mM NaF at 24 hours and 46 hours was significantly higher compared to 150 mM NaCl. The hydration of 150 mM NaF at 10 hours, 24 hours, and 46 hours was also higher compared to NaF 300 mM (Figure 2C).

Hydration in bovine

In bovine, as with camel corneal stroma, hydration was also high during the first 2 hours compared to hydration during later hours (2–46 hours) in all solutions (Figure 2A). The rate of hydration gradually reduced as time increased. There was no significant difference in change of hydration among the solutions (Figure 2B). Total hydration at 2 hours, 10 hours, 24 hours and 46 hours was also not significantly different (Figure 2C).
Comparison between camel and bovine stroma hydration

There was a significant difference in change in hydration observed between camel and bovine for some solutions. Camel stromal hydration (high) in 150 mM NaCl was significantly higher compared to bovine stromal hydration in the same solution at 10–24 hours, and 24–46 hours (Figure 2B). At 2–10 and 24 hours, change in hydration was also significantly different between camel stroma hydration in NaCl and bovine stromal hydration in NaF 300 mM (Figure 2B).

With respect to total hydration, camel corneal stroma imbibes more fluid over 2, 10, 24, and 46 hours compared to bovine stroma. In NaF 300, camel stroma hydration was significantly higher at 10 hours and 24 hours compared to bovine stromal hydration (Figure 2C).

Hydration and rehydration of fresh corneal stroma over a 14-hour period at 2-hour intervals

Hydration (H) and rehydration (RH) in camel stroma

The hydration of corneal stromas from fresh eyes was carried out in the above bathing solutions for 14 hours. At 2-hour intervals hydration levels were recorded as described earlier in the Method section. The initial mean hydration at 0–2 hrs, in 150 mM NaCl (H = 8.173), 300 mM NaCl (H = 8.325), and 150 mM NaF (H = 7.937) was significantly higher (Z > 0.000)
compared to later hydration (2–14 hours) in 150 mM NaCl ($H = 3.25 - 0.96$), 300 mM NaCl ($H = 2.94 - 1.21$), and 150 mM NaF ($H = 2.83 - 0.75$). The level of hydration among the solutions was not significantly different from each other (Figure 3A).

Following hydration, the samples were air dried for 6 hours and then samples were rehydrated in NaCl (150 mM and 300 mM) and NaF (150 mM) over a 14-hour period at 2-hour intervals. The initial mean rehydration of the samples during 0–2 hours in 150 mM NaCl ($RH = 7.452$) and 150 mM NaF ($RH = 5.940$) was significantly higher ($Z > 0.000$) compared to later hydration (2–14 hours) in 150 mM NaCl ($RH = 2.49 - 1.09$), 300 mM NaCl ($RH = 2.74 - 1.09$), and 150 mM NaF ($RH = 0.65 - 0.21$) (Figure 3A). The level of rehydration among the solutions at 0–2 hours was not significantly different from each other. The level of hydration in NaF 150 mM, was significantly less compared to NaCl 150 mM and 300 mM at 2–4 hours ($Z = 0.016$) and 4–14 hours ($Z = 0.016$).
The comparison between hydration and rehydration of camel corneal stroma showed that hydration in 150 mM and 300 mM NaCl was not different from the rehydration in these solutions. But rehydration in NaF (150 mM) was significantly less than hydration in the same solution. This suggests that PGs might have leached out in NaF during hydration.

Hydration and rehydration in bovine stroma
Similar to camel, in bovine the initial hydration at 0–2 hours, in 150 mM NaCl (H = 8.228), 300 mM NaCl (H = 8.578), and 150 mM NaF (H = 7.788) was significantly higher (Z > 0.000) compared to later hydration (2–14 hours) in 150 mM NaCl, 300 mM NaCl, and 150 mM NaF (Figure 3B). The level of hydration among the solutions was not significantly different from each other.

The initial rehydration (0–2 hours) of bovine corneal stroma was also higher compared to the later regular 2-hour hydration (2–14 hours) in all solutions (Figure 3C). During the 2–4-hour period, the rehydration in NaCl 150 mM was significantly higher than rehydration in NaF 150 mM. The rehydration in NaCl 150 and NaCl 300 mM was not significantly different from the hydration in the same solutions. But the rehydration in NaF 150 mM was significantly reduced compared to hydration in NaF 150 mM (Figure 3C).

Comparison of camel and bovine hydration and rehydration of stroma
Comparison of hydration and rehydration of camel and bovine stroma showed that there was no significant difference between them in NaCl (150 mM, 300 mM). The hydration and rehydration of camel stroma in NaF was significantly less (Z = 0.016) than bovine in the same solution at 4–14 hours. This suggests that leaching of PGs is taking place in both corneas but it is occurring more quickly in camel than bovine cornea.
Discussion

In the present study we concentrated on passive swelling, because it is driven by gel pressure which is, in turn, generated by electrostatic repulsion between the net fixed negative charges in the stroma. To study passive swelling, we removed the epithelium and endothelium. The epithelium was removed to avoid interference with the determination of stromal sodium concentration because it has a very slow exchanging sodium pool.27 The endothelium was removed to inhibit stromal swelling due to the presence of a balancing outward-directed bicarbonate ion pump located in the corneal endothelium.28

Swelling studies have been carried out on rabbit, bovine, and human corneal stroma.29–31 Corneal stroma has an innate tendency to imbibe fluid and swell; when it swells it loses its transparency.17 The ability to swell and the transparency of fresh corneal stroma are both unusual properties for a connective tissue.22,32 The corneal stroma has swelling properties in isotonic saline27,22 which can be quantitatively explained by the Donnan theory of corneal swelling.33 The single variable which regulates all the phenomena associated with the Hodson–Donnan theory of corneal swelling is the cation exchange capacity, Q, of the matrix molecule. Part of the cation exchange capacity of the corneal stroma is donated by the carboxylic and sulphonic acid groups of the glycosaminoglycans (GAGs)33 which are associated with collagen fibrils.34

Bron35 reported that GAGs of the corneal stroma are keratan sulfate (KS; a component, for instance, of the proteoglycans lumican) dermatan sulfate (DS) and chondroitin sulfate (CS) (components of small proteoglycans, decorin). In bovine corneal stroma, the keratan sulfate/chondroitin-4-sulfate ratio is higher posteriorly than anteriorly.36,37 If this is the case for human cornea, then since keratan sulfate has a higher water affinity than chondroitin-4-sulfate, this could explain, in part, the greater degree of posterior stromal swelling on immersion.35,38 This could be the case in our 46-hour comparative study of swelling of camel and bovine corneal stroma. The level of hydration in camel cornea in all solutions was found to be significantly higher compared to bovine cornea. We presume that camel cornea could have a higher KS/CS ratio compared to bovine cornea, and swell more than bovine cornea. If there is a higher concentration of KS, then there will be a higher amount of KS-PGs such as lumican, keratocan, and mimican compared to CS-PGs such as decorin.

Results of our 20-minute study suggested that there was no selective affinity for a particular ionic solution during the process of swelling. The results of our 46-hour study showed that camel stroma has a high level of hydration in all solutions and there is no significant difference in levels of hydration among the solutions from each other during the initial period (0–2 hours). This is consistent with the studies of Kinsey and Cogan39 and Hedbys,40 in which it was found that at high hydration (H > 10) swelling does not reduce with increases in concentration of bathing solutions. In this condition, a fixed charge of GAGs also increases. It is believed that in camel cornea, high levels of hydration are not affected by the concentration of bathing solution due to the presence of high fixed charge on GAGs.

Our 14-hour study suggested that leaching of PGs takes place in both corneas but occurs at a higher rate in camel than bovine cornea. It has been suggested that there is the possibility of a differential leaching of GAGs from the stroma during prolonged immersion.35 Although only around 1% of keratan sulfate is lost from corneas held in closed culture41 a significant loss of proteoglycans from swollen corneas has been recorded by some researchers,42,43 with a preferential loss of keratan sulfate from edematous rabbit corneas.44 Differential loss has not been studied, but a greater loss of GAGs from the anterior stroma could reduce its “swellability” in human cornea.35 There may be a high loss of GAGs from the camel cornea which inhibits “swellability”.

Further studies are required to investigate the structural and biochemical analysis of proteoglycans in bovine and camel cornea. These studies could involve biochemical, immuno-histochemical, and electron microscopic techniques.

Disclosure

The authors report no conflicts of interest in this work.

References


