Lanosterol Eye Drops in a Human Juvenile Nuclear Cataract

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Abstract
A revolutionary non-surgical and low-cost treatment of the largest cause of world's blindness could have an immeasurable effect on global health, since surgical procedures are not accessible to all patients worldwide. In cataracts we still keep replacing a clouded lens without considering the pathogenetic mechanism of lens clouding, and without helping the lens epithelium to restore its homeostatic functions. In the light of recent scientific findings and considering the lack of studies on humans in this field we tested lanosterol eye drops on a human juvenile nuclear cataract with a lower lanosterol concentration and with a different vehicle compared to the eye drops recently tested on animals. Our experiment has demonstrated that there were no clinically relevant changes in intraocular pressure neither reversal of cataract using eye drops with a relatively low dose of lanosterol. These results encourage to test eye drops with higher lanosterol concentrations on human cataract cases.

Aim
The aim of this study was to demonstrate clinically relevant changes in intraocular pressure by the use of low concentration lanosterol eye drops.

Keywords: Blindness; Lens clouding; Human cataract; Lanosterol
1. **Introduction**

Cataract is considered responsible for the half the world's blindness and 33% of visual impairment worldwide [1, 2] and a nonsurgical treatment would potentially be able to prevent cataract formation and to treat cataracts without the surgical costs and risks also in developing countries, where cataracts contribute to more than 90% of the total disability-adjusted life year [2] as well as in countries where access to eye care is limited. Actually the protein aggregation amidst the regularly arranged crystallins of lens fibers is considered responsible of the pathogenesis of cataract [3], and the finding of mutations in lanosterol synthase as a cause for cataract formation in rats and subsequently in families with congenital cataract has encouraged in evaluating its role in the formation of senile cataract in rabbits and dogs, where Zhao et al. have shown that lanosterol acts by reversing the protein aggregation within the lens fibers both in vitro and in vivo [4]. In order to study the effects of lanosterol in human cataract we evaluated if a lanosterol 5 mM solution could reverse a unilateral juvenile nuclear cataract.

2. **Subject and Methods**

An evaluation request was asked to the Hospital Ethical Committee before preparing the eye drops and informed consent was obtained from the patient. Lanosterol 5mM solution was prepared by adding Ph. Eur. sterile olive oil to lanosterol 20 mg powder (Sigma Aldrich) at room temperature to a final volume of 8 ml. Because of the solubility grade of lanosterol in olive oil, the mixture needed more than 20 minutes of manual dissolution. After that we performed a filtration (with a 0.2 μm filter designed for oily solutions) to guarantee appropriated sterility to the eye drops. The patient with idiopathic unilateral juvenile nuclear cataract was treated with lanosterol 5 mM eye drops two times daily for the first week and three times daily for the next seven weeks. Visual acuity, intraocular pressure and slit lamp examination were repeatedly measured during this period.

3. **Results**

During the study period the patient showed a progressive worsening of visual acuity with an increasing of miopic shift (Table 1) as well as a slight worsening of the degree of lens opacity in slit lamp examination (See Figure 1 and 2), without clinically relevant effects on intra-ocular pressure during the whole observation period (Table 2).

4. **Conclusion**

Lanosterol 5 mM oily ophtalmic solution did not improve visual acuity in a single case of human idiopathic juvenile nuclear cataract.

<table>
<thead>
<tr>
<th>Time</th>
<th>Myopic Shift (Spherical Equivalents)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visit 1</td>
<td>-0.5</td>
</tr>
<tr>
<td>Visit 2</td>
<td>-3</td>
</tr>
<tr>
<td>Visit 3</td>
<td>-3.5</td>
</tr>
<tr>
<td>Visit 4</td>
<td>-4</td>
</tr>
<tr>
<td>Visit 5</td>
<td>-13</td>
</tr>
</tbody>
</table>

Table 1: The patient showed a progressive worsening of visual acuity with an increasing of miopic shift.
Figure 1 and 2: Slit lamp examination of slight worsening of the degree of lens opacity.

![Figure 1 and 2: Slit lamp examination of slight worsening of the degree of lens opacity.](image)

<table>
<thead>
<tr>
<th>Time</th>
<th>Affected Eye</th>
<th>Contralateral Eye</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visit 1</td>
<td>19</td>
<td>16</td>
</tr>
<tr>
<td>Visit 2</td>
<td>19</td>
<td>17</td>
</tr>
<tr>
<td>Visit 3</td>
<td>18</td>
<td>11</td>
</tr>
<tr>
<td>Visit 4</td>
<td>19</td>
<td>14</td>
</tr>
<tr>
<td>Visit 5</td>
<td>17</td>
<td>14</td>
</tr>
</tbody>
</table>

Table 2: Effects on intra-ocular pressure during the whole observation period.

5. Discussion

Before testing lanosterol eye drops on humans we reconsidered the eye drops composition of the lanosterol eye drops recently tested on animals in order to have a potentially more appropriate vehicle for human use. Sterile olive oil was the vehicle chosen because of the moderate solubility of lanosterol in it that permitted to avoid the use of ethanol (EtOH) in order to have an expected greater tolerability. A relatively low concentration of lanosterol was chosen because of the lack of studies on humans in this field. In our experiment five mM lanosterol solution failed to reverse nuclear opacity of human juvenile nuclear cataract after 8 weeks of treatment. During the study period no cataract regression, no visual acuity improvement and no visual acuity stabilization could be observed. However the concentration of lanosterol used by us may have been inadequate to reverse the nuclear cataract in human lenses. Shanmugam et al. found that Twenty-five mM lanosterol solution failed to reverse nuclear opacity of human age-related cataractous nuclei after 6 days of incubation [5]. Considering that it is also possible that molecules other than lanosterol could play a role in human nuclear cataract reversal, specially in age-related senile cataracts, where a variety of biochemical and physical insults lead to changes in the lens proteins, determining increased protein aggregation and cataract formation. Hejtmancik et al. have elucidated that congenital and senile cataracts show different molecular pathways [6, 7], in particular mutations in enzymes such as lanosterol synthase and or every mutation that destabilize the lens proteins lead to congenital cataract formation due to increased protein aggregation.
Furthermore, multiple authors have shown that congenital cataracts are usually secondary to missense mutations causing the accumulation of altered protein residues or, in specific cases, to ferritin levels, while age-related imbalance of chaperones and accumulation of degraded and denatured normal proteins was usually shown in senile cataract [8]. Therefore, future basic science research and clinical studies may provide us with more information concerning different pathogenesis and more specific treatments. It may be reasonable to verify if different lanosterol concentrations or the choice of a different solvent or treating cataract with different etiologies could improve clinical outcomes.

References