Role of high concentrations of mannitol on the stability of hyaluronic acid in an oxidative stress model induced by xanthine/xanthine oxidase

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Background

Osteoarthritis (OA) is a degenerative joint disease associated with harmful action of reactive oxygen species (ROS).

ROS are involved in the degradation of both aggrecan and high-molecular-weight hyaluronic acid (HMW-HA). The later plays a key-role in the joint lubrication and the visco-elastic and shock absorbing properties of the synovial fluid (SF).

Viscosupplementation consists in injecting intra-articularly exogenous HMW-HA to restore the SF rheological properties, that are dramatically decreased in OA.

However the injected HA is also rapidly degraded by ROS, decreasing its effectiveness and duration of action.

Objective

To evaluate the ability of mannitol, a powerful oxygen free radical scavenger, to reduce exogenous HMW-HA degradation using a model of oxidative stress induced by xanthine (X) + xanthine oxidase (XOD).

Methods

Hyaluronan (MW# 0.8 mDa) was submitted to an oxidative stress generated by the addition of X + XOD.

Then solution of the same HA + 35 g/L of mannitol in PBS buffer was studied.

Different enzyme concentrations (XOD 109 mIU/mL and 218 mIU/mL) were used and the HA properties were studied after 24 hours of contact at ambient temperature.

Changes of the viscosity of the solution were assessed by rheometry (rheology was determined using a rheometer at 25°C using a cone and plate geometry, steady-state viscosity was determined in Pa.s, as a function of the shear rate).

Results

The presence of X/XOD degraded HA in the conditions tested:

- HA viscosity decreased as a function of XOD concentration,
- HA MW decreased dramatically by 36.6%.

On the opposite, in presence of high concentration of Mannitol:
- HA viscosity was stable,
- HA MW decreased only slightly (-11%).

<table>
<thead>
<tr>
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<th>Initial</th>
<th>+10 microl. enz.</th>
<th>+32 microl. enz.</th>
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<tbody>
<tr>
<td>HA</td>
<td>798000 ±77600</td>
<td>62500 ±67300</td>
<td>503800 ±49890</td>
</tr>
<tr>
<td>HA Mannitol</td>
<td>781200 ±75400</td>
<td>562200 ±67300</td>
<td>674000 ±80030</td>
</tr>
</tbody>
</table>

Conclusion

High concentrations (3.5%) of mannitol protect HA from ROS-mediated degradation.

These in vitro data suggest that mannitol may increase the intra-articular residence time of HA and consequently may improve clinical efficacy of viscosupplementation.