8-1-2011

Lubricin Immunohistochemical Expression in Human Temporomandibular Joint Disc with Internal Derangement

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Lubricin immunohistochemical expression in human temporomandibular joint disc with internal derangement

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Abstract
Lubricin is a chondroprotective, mucinous glycoprotein which contribute to joint lubrication, especially to boundary lubrication and maintains joint integrity. The present investigation aimed to study the immunolocalization of lubricin in TMJ discs from patients affected by anterior disc displacement with reduction (ADDwR) ADDwoR. Eighteen TMJ displaced disc affected by ADDwoR were processed immunohistochemically, with a polyclonal anti-lubricin antibody, used at 1:50 working dilution. The percentage of lubricin immunopositive cells (extent score = ES) and the extent of lubricin staining of the disc extracellular matrix
(ECM), were evaluated. Each sample was scored for histopathological changes. Percentage of immunostained surface disc cells was the same (ES = 4) in both control and ADDwOR cells, being this data not statistically significant ($P < 0.05$). In pathological specimens the percentages of lubricin-stained cells was very high with an ES of 4 respect to control specimen, and this difference was statistically significant different ($P > 0.05$). The extracellular matrix (ECM) of discs at the disc surfaces of both pathological and normal specimens was very heavily stained (++++). Both the ES and ECM staining were not statistically correlated to the TMJ degeneration score according to the Spearman’s rank correlation coefficient. According to our findings, a longstanding TMJ disc injury, affects lubricin expression in the TMJ disc tissue and not its surfaces, moreover, lubricin immunostaining is not correlated to TMJ disc histopathological changes.

**Keywords**

Derangement, disc, lubrification, temporomandibular joint disorders

**Introduction**

Disc displacement or internal derangement (ID) of the temporomandibular joint (TMJ) is defined as an abnormal positional relationship of the disc relative to the mandibular condyle and the articular eminence (1). There are two common types of ID: as an anterior disc displacement with reduction (ADDwR) and an anterior disc displacement without reduction (ADDwoR) (2, 3). ID is a frequent condition that has been documented by clinical, anatomical, radiographical and surgical observations (4, 5). It should be noted that ID frequently precedes the onset of TMJ osteoarthrosis (OA). The process of TMJ–OA is characterized by degenerative joint changes, such as deterioration and abrasion of articular cartilage and disc surfaces, as well as occurrence of thickening and remodelling of the underlying bone. Nevertheless, little is known about the aetiopathogenic mechanisms leading to disc displacement as investigations did not reach consistent results (6-10).

Trauma, functional overloading, joint laxity, lack of lubrification, hyperactivity of the lateral pterygoid muscle as well as occlusal factors have also been mentioned in this respect (4). Actually, it is believed that the most significant aberration of all suggested processes, terminating in disc displacement, is an increase in joint friction (6), because of a reduction in boundary lubrication. As a consequence, joint lubrication can be considered of great importance for a better understanding of the pathophysiology of the TMJ diseases (11).

Several molecules present in synovial fluid have been reported to contribute to joint lubrication, especially to boundary lubrication (1, 6). Among them, a mucinous glycoprotein called lubricin. Lubricin is a chondroprotective, mucinous glycoprotein, which is a product of the proteoglycan 4 (PRG4) gene, and is homologous to molecules referred to as superficial zone protein (SZP). Lubricin, also known as SZP, megakaryocyte stimulating factor, hemangiopoietin or PRG4 has been found in a number of tissues, including synovial membrane and fluid (12-14), superficial zone of cartilage (15), tendon (16) and meniscus (17, 18). The essential role for lubricin, in maintaining joint integrity, has been demonstrated through the identification of disease-causing mutations in patients with the autosomal recessive disorder camptodactyly-arthropathy-coxa vara-pericarditis syndrome (19).

As far as the TMJ is concerned, several reviews (6, 11, 20) have underlined the importance of lubricin as a biochemical factor associated with the regulation of TMJ lubrication. Nonetheless, data from these investigations were based on joints other than TMJ. In fact, except for one study which demonstrated the importance of lubricin in bovine mandibular condyle cartilage (15), no data are available for TMJ disc. Accordingly, the present investigation aimed to study the immunolocalization of lubricin in TMJ discs from patients affected by ADDwoR vs. normal TMJ discs, to determine any difference between these two samples in the percentage of disc cells that contained lubricin and the degree of extracellular matrix (ECM) staining.
Materials and methods

Patients and tissues
Eighteen TMJ-displaced disc specimens were collected from consecutive patients treated between 2002 and 2009 at the University of Pontifical Catholic University of Paraná, Brazil. Sixteen virtually unaffected human TMJ discs were studied along with the pathological material. After project approval by the ethic committee of the same University, informed consent was obtained from each patient before tissue collection.

The discs (10 from the right TMJ and 8 from the left) were obtained from 14 female and 4 male patients, affected by ADDwoR, as confirmed by their clinical history, examination and magnetic resonance imaging studies. The mean age of all patients was 43.2 ± 5.4 years. The diagnosis that led to surgery was disc derangement with associated pain and impairment of function. These patients had been unsuccessfully non-surgical treated before surgery and had TMJ pain or interference with mandibular movements. TMJ discs from patients with other TMJ diseases, dentofacial deformity, major jaw trauma, previous TMJ surgery and previous steroid injections in the TMJ were not included in this study.

All the surgically removed patients’ discs were macroscopically deformed, none of the discs having a normal biconcave shape.

The previously mentioned sixteen autopsy TMJ disc specimens were obtained bilaterally from eight cadavers (three males and five females). The mean period (and standard deviation) between the death of the donor and tissue harvest was 14 ± 2 h. The cadaver donors’ mean age was 44 ± 6 years, at the time of death. The causes of death of the donors were cardiovascular disease and cancer.

None of the donors’ clinical history revealed generalized joint disease or TMJ arthropathy. Macroscopic examination of the discs showed no signs of degenerative or inflammatory joint disease nor disc displacement and no pathological changes at microscopic examination. The original sample collection, previously used in former studies (21-25), was from the Institute of Anatomy at the University of Catania, Catania, Italy. Both patients and autopsy specimens included the anterior band, intermediate band and the posterior band.

Immunohistochemistry
The discs were fixed overnight in 10% neutral-buffered formalin (Bio-Optica, Milan, Italy). After fixation and overnight washing, each disc was sectioned through its centre along a parasagittal plane, perpendicular to its long axis. Each tissue block was dehydrated in graded ethanol and embedded in paraffin, preserving the longitudinal anatomical orientation. Specimens were sectioned at a thickness of 5 μm using a microtome, placed on silanized glass slides (the anterior band being always on the right side of the slides), and warmed at 60°C for a minimum of 2 h to ensure proper tissue adhesion to the slides prior to immunohistochemical staining. The TMJ disc sections were deparaffinized in xylene and rehydrated in reagent alcohol, and then, they were treated with hydrogen peroxide for 10 min to quench endogenous peroxidase activity. Then the sections were heated (5 min × 3) in capped polypropylene slide holders with citrate buffer (pH 6), using a microwave oven (750 W) to unmask antigenic sites. Non-specific binding site blocking was achieved by treating the specimens with 5% horse serum for 30 min.

For immunohistochemical staining, we used a rabbit polyclonal anti-lubricin antibody (Novus Biologicals, LLC, Littleton, CO, USA), used at 1:50 working dilution. The antibody was applied onto sections, and afterwards, they were incubated overnight at 4°C in a moist chamber. The secondary antibody, biotinylated anti-mouse/anti-rabbit IgG was applied (for 30 min, at RT), followed by the avidin-biotin-peroxidase complex (Vector Elite Kit Abbott, Chicago, IL, USA) for 30 min, at room temperature. The immunoreaction was visualized by incubating the sections for 4 min in a 0.1% 3,3′-diaminobenzidine and 0.02% hydrogen peroxide solution (DAB substrate kit; Vector Laboratories, Burlingame, CA, USA). The sections were lightly counterstained with Mayer’s...
haematoxylin (Histolab Products AB, Goteborg, Sweden) and finally mounted in GVA mount (Zymed Laboratories Inc., San Francisco, CA, USA).

**Evaluation of immunohistochemistry**

The lubricin-staining status was identified as either negative or positive. The percentage of lubricin immunopositive cells (extent score = ES) was independently evaluated and scored as a percentage of the final number of 100 cells in four categories: 0 = <5%; 1 = 6–30%; 2 = 31–50%; 3 = >51–75% and 4 = >76–100%. Counting was performed at ×200 magnification. Positive staining was defined as the presence of a brown detection chromogen on the edge of the haematoxylin-stained cell nucleus, distributed within the cytoplasm, or in the immediate lacunar/pericellular space. We also graded the lubricin staining of the disc ECM according to 0 to ++++ scale with the arbitrary cut-offs: 0 = none; + = up to 25% of the area; ++ = 26–50% of the area; +++ = 51–75% of the area and ++++ = 76–100% of the area at ×200 magnification. Three different TMJ disc tissue compartments were analysed, namely the central region (i.e. the inner portion) of the disc and the temporal (superior) and condylar (inferior) disc surfaces.

**Positive and negative controls**

Positive and negative controls were performed to test the specific reaction of primary antibodies used in this study at a protein level. For positive control testing, the sections from bovine calf stifle joint underwent an immunoperoxidase process (17). The positive immunolabelling for lubricin was both perinuclear and cytoplasmic. For negative control testing, sections of TMJ displaced discs were randomly drawn from degenerated disc samples. These were then treated with normal rabbit serum instead of the specific antibodies.

**Histopathological degeneration grading score**

One of three sections for each disc was stained with haematoxylin & eosin (H&E), to assign a histopathological degeneration score. Briefly, this score takes into account pathological disc tissue transformation, i.e. collagen bundles, non-specific degenerative changes and the presence of blood vessels. This grading system results in a score ranging from 0 up to 8 for heavily degenerated disc tissue (26).

**Statistical analysis**

Mean values and standard deviations were obtained for the ES and ECM staining of three disc portions examined in each sample. Data were further analysed using Mann–Whitney U-tests to evaluate any significant statistical difference, both for pathological and for normal discs. Comparisons were made based on the anatomical region of the disc, i.e. central portion, as well as superior and inferior disc surfaces. The non-parametric Spearman rank correlation test was used to obtain the correlation coefficient between the lubricin staining of the extracellular matrix and the percentage of cells staining for lubricin. P-values of <0.05 were considered statistically significant. All data were analysed with the SPSS program (SPSS® release 16.0; SPSS, Chicago, IL, USA).

**Results**

All TMJ patients had been unsuccessfully treated with other, various, interventions, for at least 6 months before surgery. Diseased TMJ disc sections, stained with haematoxylin (routine staining), showed a different severity of morphological damage, ranging from an altered collagen bundles architecture to the fragmentation of collagen bundles. A decrease in cellularity was appreciated too, in the more damaged discs. In contrast, the control discs showed a preservation of multidirectional collagen bundles; predominately fibrocyte-like cell populations with few chondrocyte-like cell aggregates; absence of clefts and/or fraying.

The average TMJ disc degeneration score of our TMJ ADDwoR disc sample was 3.52 ± 1.17.
Every TMJ disc sections, from pathological and control samples, displayed lubricin-positive immunohistochemical staining, although to a different extent. Lubricin staining within the TMJ disc cells was ubiquitous and appreciated in fibroblast-like cells, fibrochondrocytes or chondrocyte-like cells. In almost all cells, immunostaining was cytoplasmic or pericellular in nature (Fig. 1). Percentage of immunostained surface disc cells was the same (ES = 4) in both control and ADDwoR cells, being this data not statistically significant (P > 0.05). In the two samples, no statistically significant difference (P > 0.05) was obtained when comparing cell immunolabelling at the superior and inferior disc surfaces between. On the other hand, in the inner portion of the disc, the percentage of immunolabelled cells was statistically significant different (P < 0.05) between the two samples. In pathological specimens, the percentage of lubricin-stained cells was very high with an ES of 4 being cells immunostained in almost all of the cases (Fig. 2). In control specimens, this percentage was lower with an ES of 2, thus no more of 50% of cells were immunolabelled in each field. In fact, not every cell was immunolabelled by lubricin antibody, in the inner portion of the discs (Table 1).

Figure 1
Control temporomandibular joint (TMJ) disc specimen. Lubricin staining evidenced by brown chromogen in the different cell phenotypes of TMJ disc, fibroblast-like cells (f) and chondrocyte-like cells (c). Chromogen deposition appears granular (empty arrow) or band-like (arrow) at and/or between adjacent collagen bundles. Bar 50×.
Figure 2
Immunohistochemical staining of human control temporomandibular joint (TMJ) disc (A) and TMJ disc affected by ADDwoR (B). Extracellular matrix is much more heavily stained in diseased disc specimen. Bar 100×.

Table 1. Extent score and extracellular matrix immunostaining for lubricin in normal temporomandibular joint (TMJ) disc and anterior disc displacement without reduction (ADDwoR)

<table>
<thead>
<tr>
<th></th>
<th>Number of specimens</th>
<th>ES at the disc surfaces</th>
<th>ES at the disc inner portion</th>
<th>ECM staining at the disc surfaces</th>
<th>ECM staining at the disc inner portion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal TMJ discs</td>
<td>16</td>
<td>4</td>
<td>2</td>
<td>++++</td>
<td>+</td>
</tr>
<tr>
<td>ADDwoR TMJ disc</td>
<td>18</td>
<td>4</td>
<td>4</td>
<td>+++</td>
<td>+++ (12 discs of 18) ++ (6 discs of 18)</td>
</tr>
</tbody>
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Extent score (ES) = percentage of lubricin immunolabelled cells: 0 = <5%; 1 = 6–30%; 2 = 31–50%; 3 = >51–75% and 4 = >76–100%. Lubricin staining of the disc extracellular matrix (ECM): 0 = none; + = up to 25% of the area; ++ = 26–50% of the area; +++ = 51–75% of the area and ++++ = 76–100%.

Within the ECM of discs, lubricin immunolabelling was observed in every disc specimen, although to a different extent, according to the disc region (Figs 1–3). ECM at the disc surfaces of both pathological (Fig. 3) and normal specimens was very heavily stained (++++) by lubricin antibody. On the contrary in the inner portion, ADDwoR discs were moderately immunolabelled in some instances, but a faint (+) or high immunostaining (++++) was also appreciated in few other specimens (Fig. 2). In control disc specimens, at the same topographical area, it was always observed a faint staining (+). This ECM staining was particularly noted at the interface of collagen fibre bundles, and it was characterized by intense chromogen deposition that appeared granular or band-like at and/or between adjacent collagen bundles (Fig. 1) (Table 1). Newly formed vessels and areas of chondroid metaplasia were heavily immunostained by lubricin antibody (Fig. 4A,B).
Figure 3
Lubricin staining on disc surface. On the disc surface, both cells and extracellular matrix are immunolabelled by lubricin antibodies. Bar 200×.

Figure 4
Diseased disc specimen from temporomandibular joint. Lubricin is immunohistochemically detected in newly formed blood vessels (V) inside the disc (A) and in area of chondroid (C) metaplasia (B). Bar 100×.

Both the ES and ECM staining were not statistically correlated to the TMJ degeneration score according to the Spearman’s rank correlation coefficient.
No chromogen was detected in any of the 20 negative control disc sections.

Discussion

In fact, optimal functionality of synovial joints depends on maintaining extremely low coefficients of friction. In this respect, lubricin acts as a vital counteragent against aberrant protein and/or cellular adhesion, infiltration and overproliferation and serves as a critical boundary lubricant between opposing articular surfaces (12, 13, 27, 28), in such a way that alterations in its metabolism may profoundly impact joint function. A recent study has shown that lubricin expression may vary with health and disease, and it is involved in joint osteoarthritis (29).

Several investigations have been carried out on joints (12, 16, 18, 30) other than TMJ, and to the authors’ knowledge, this is the first report regarding the presence of lubricin in human TMJ discs affected by ADDwoR.

Data from this study demonstrated that lubricin is significantly overexpressed at a protein level in cells of the inner portion of the discs affected by ADDwoR respect to the control specimen. Differences in lubricin immunostaining, although not statistically significant, were detected also for ECM; in fact, lubricin immunolabelling in some instances was more heavily expressed in TMJ with ADDwoR. The presence of lubricin in the extracellular matrix suggests a lubricin release from disc cells, diffusing through the tissue. This overexpression of lubricin in macroscopically deformed TMJ disc affected by ADDwoR may be explained both by mechanical and by biochemical factors. In ID with ADDwoR, the disc is permanently displaced or dislocated remaining in an incorrect position, and the jaw’s range of motion is limited (2, 3), meanwhile several cytokines are released in TMJ disc synovial fluid (SF) and tissue. In this respect, numerous investigations have demonstrated that different load conditions may alter disc cell biosynthesis of a variety of molecules, and several studies in other musculoskeletal tissues support that lubricin expression can be regulated by mechanical stimuli (31). Accordingly, it has been reported that lubricin is expressed by chondrocytes from the superficial zone of cartilage, because the cells are subjected to more shear force than those in the deeper zones. Furthermore, lubricin has been detected, also, in load-bearing regions of meniscus (17).

Lubricin biosynthesis varies also in response to growth factors and cytokines. It has been assumed that the proinflammatory cytokines initiate a cascade of events that lead to a decrease in joint lubrication and an increase in joint damage. Although, later the lubricin levels eventually return to normal or they are even upregulated, ongoing damage has been initiated (32). However, it has also been demonstrated that lubricin expression is downregulated by pro-inflammatory cytokines, such as interleukin-1 beta, interleukin-6 and tumour necrosis factor alpha (32). On the contrary, transforming growth factor beta increased lubricin synthesis, secretion and cartilage boundary association. Therefore, a dysregulation of lubricin metabolism, under the influence of elevated cytokine concentrations in diseased or damaged joints, might thereby lead to lubrication changes and loss-of-function (27). The overexpression of lubricin at an immunohistochemical level, detected in our sample of ADDwoR TMJ discs, corroborates previous findings obtained in long-standing joint injury. In fact, it has been reported that following injury (32) in the SF of injured joints, lubricin levels were significantly lower compared with the contralateral joints at the early stage following injury, but normal lubricin levels were approached within 12 months, and a significant increase in the lubricin concentration with time post-injury was recorded later (32). The lack of any correlation between the extent of lubricin staining of the TMJ disc cell and extracellular matrix with ADDwoR and the histopathological score obtained in our sample is in accordance with previous findings, which reported that the extent of lubricin present in the intervertebral disc extracellular matrix and native disc cells was independent of the Thompson grade (33).

Thus, according to our findings, a long-standing TMJ disc injury affects lubricin expression in the TMJ disc tissue and not its surfaces; moreover, lubricin immunostaining is not correlated to TMJ disc histopathological changes.

Understanding the factors that may lead to, or result from, disc displacement is essential not only to permit the choice of optimal treatment but also to develop preventive treatment modalities (6). The fact that lubricin is constitutively expressed in TMJ disc and it is overexpressed during long-standing disease deservers further
studies to correlate its expression to the time of disease onset and cytokine expression to choose the proper kind of treatment.

New therapeutic perspectives, in fact, are going to be opened, for joints other than TMJ, that could be beneficial in the treatment of TMJ diseases, at least at the early beginning of disease (15, 34-36). These strategies consist of rescuing lubricin dysfunction. Moreover, the capacity for recombinant lubricin to effectively bind to and lubricate articular cartilage surfaces suggests therapeutic implications for the delivery of applicable biolubricant formulations.

References


