Investigation of the effects of fascia lata on neochondrogenesis in a rabbit model

Fasia latanın neokondrogenezis üzerinde olan etkilerinin tavşan modelinde araştırılması

Şengül Yılmaz¹, Özge Bilkay², İsmail Özdemir³, Arzu Avcı⁴, İslı Adadan Güvenç⁵

¹Department of Otorhinolaryngology, Denizli Government Hospital, Denizli, Turkey
²Department of Otorhinolaryngology, Alsancak Government Hospital, Izmir, Turkey
³Department of Otorhinolaryngology, Gazi Hospital, Izmir, Turkey
⁴Department of Pathology, İzmir Katip Çelebi University Medical Faculty, İzmir, Turkey
⁵Department of Otorhinolaryngology, Bağlent University Hospital, İzmir, Turkey

Abstract

Objective: The impact of fascia lata on neochondrogenesis placed between both leaflets of the perichondrium after full thickness submucoperichondrial cartilage resection was investigated in a rabbit septum model.

Methods: Twenty adult New Zealand rabbits were studied in two groups with equal number of animals in each group. Full thickness cartilage with an area of 1 cm² was resected to create a cartilage defect in all rabbits. Autogenous fascia lata was interposed between both leaflets of the perichondrium and the surgical field was closed (fascia lata group). Any material was not interposed between both leaflets of the perichondrium of the defective area (control group). Rabbits were sacrificed on postoperative 90th days and their septa were totally excised. Neochondrogenesis was investigated histopathologically with a light microscope. The preparations were evaluated as for thickness of newly formed cartilage (in mm), inflammatory reaction and presence of macroscopic perforation.

Results: Thickness of the cartilage in the fascia lata group was significantly greater than that of the control group (M-W-U=15,000, p<0.05).

Conclusion: Use of fascia lata in the acceleration of perichondrial neochondrogenesis can result in successful outcomes in the repair of cartilage defects.

Keywords: Neochondrogenesis, nasal septum, fascia lata.

Özet

Amaç: Submukoperikondrial tam kat kıkırdak rezeksyonu sonrası her iki perikondrum yaprak arasına konan fasia latanın neokondrogenezis üzerine olan etkilerinin araştırılması.

Yöntem: Yirmi adet genç erişkin Yeni Zelanda tavşan eşit sayıda iki grubu ayrıldı. Her tavşanın nazal septumlarında 1 cm²’lik alan kat kıkırdak rezeksyonu yapıldı. Olusan kıkırdak defektine ilk grupta tavşanın kendisinden alınan fasia lata submukoperikondrial olarak yerleştirildi (fasia lata grubu). Diğer grubu herhangi bir materyal konulmadı (kontrol grubu). Operasyon sonrası 90. günde defekt bölgesini içine alacak şekilde subtotal septum rezeksyonu yapıldı. Yeni kıkırdak oluşumu histopatolojik olarak, işık mikroskobuyla incelendi. Değerlendirmede perikondrum kökenli yeni kıkırdak oluşumu, ilihtabı reaksiyon ve nekroz, makroskopik olarak da perforasyon oluşumu göz önüne alındı.

Bulgular: Fasia lata yerleştirilen çalışma grubunda oluşan yeni kıkırdak doku kontrol grubuna oranla anlamlı derecede fazla bulundu (M-W-U=15.000, p<0.05).

Sonuç: Fasia latarın neokondrogenenezise olan katkıda, kıkırdak defektlerinin onarımına bu materyalin kullanılabilirliğini desteklemekte-dir.

Anahtar sözcükler: Neokondrogenesis, nazal septum, fasia lata.

Cartilage tissue has a mesenchymal origin and consists of chondrocytes and matrix. Basic function of this tissue is to protect and support soft tissues.¹² Growth and proliferation of chondrocytes are realized through cellular proliferation from perichondrial tissue of the young cartilage which has chondrogenic properties (appositional growth).
and via mitotic division of available young chondrocytes (interstitial growth). When cartilage is damaged chondroblasts migrate to the injured region and form a new cartilage tissue. If a larger damaged area is present and perichondrial regeneration is not fast enough to compensate the loss, then the defect cannot be repaired completely and heals with formation of a scar tissue.\(^2,3\)

Perichondrium wraps the cartilage and forms an interface between the cartilage and the surrounding tissues. Perichondrium provides support for the avascular cartilage.\(^1,2,4\) Perichondrium is a diffuse connective tissue layer required for the growth and maintenance of the cartilage. It is rich in collagen and contains numerous fibroblasts. Chondroblasts found in the inner layer of perichondrium differentiate and form chondrocytes.

Defects occurring in the cartilage tissue due to various reasons lead to significant functional and esthetic problems which require replacement of the cartilage.\(^6\) Unfortunately, it is very difficult for chondrocytes embedded in matrix to migrate to the defect site so as to start repair process, so cartilage defects hardly heal and regenerate.\(^6\) Therefore, various materials have been used to repair the cartilage defects. Nowadays, mostly autogenous cartilage grafts have been used.\(^5,6\) In 1959, Lester reported that perichondrium dissected away from the cartilage induced formation of new cartilage tissue. Since this discovery of chondrogenic characteristic of perichondrium, perichondrial grafts have been used with successful outcomes.\(^5,6\) Though in young patients these procedures have yielded good results, in some old patients frustrating outcomes have been obtained. These unfavorable results were associated with physiological changes in the cartilage and decrease in in the neocondrogenic potential with aging.\(^6\) If a larger area is damaged, then perichondrium cannot induce cartilage formation with due rapidity. In this case defect is closed with scar tissue which is formed by rapidly developing connective tissue.\(^6\) Therefore, a perichondrial neochondrogenesis faster than the developmental rate of scar tissue is critically important.\(^6\) Fascia lata has been used to enhance regeneration of the cartilage tissue with successful outcomes. The advantages of fascia lata include, its acceptance by the surrounding tissue without any complication, its biocompatibility and its collagenous and regenerative characteristics.\(^10-13\)

In this study, the impact of fascia lata on perichondrial neochondrogenesis developed in defective areas of cartilages was investigated in a rabbit septum model.

### Materials and Methods

#### Study Design

Twenty adult New Zealand rabbits were studied in two groups with equal number of animals in each group. Anaesthesia was applied using 2 mg/kg xylazine hydrochloride (IM) and ketamine hydrochloride (IM). A median vertical incision was made on nasal dorsum from frontal bone up to nose tip. Nasal bone was retracted laterally from the midline to approach to the septum. Perichondrium was elevated bilaterally beginning from dorsal edge of the septum down to its base. Then full thickness cartilage with an area of 1 cm\(^2\) (Fig. 1) was resected to create a cartilage defect.

In Group 1, any material was not interposed between both leaflets of the perichondrium of the defective area. Bony parts were closed. Subcutaneous tissue and skin were closed with sutures before termination of the surgical procedure. This group was termed as the “control group.”

Lateral aspects of the femoral region of ten rabbits were incised and after passing through skin and subcutaneous tissue, fascia lata was approached. A portion of the cartilage with the same size of the defect area was excised. Then this portion of the cartilage was interposed between both leaflets of the perichondrium and the surgical field was closed. This group was termed as “fascia lata” group.

#### Outcome Parameters

Rabbits were sacrificed on postoperative 90th days and their septa were totally excised. The harvested specimens were kept in 10% formaldehyde solution. They were then fixated and embedded in paraffin blocks. Afterwards, 5 μ sec-

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Fig. 1. Full thickness cartilage was resected with an area of 1 cm\(^2\) in size.
tions were cut using a microtome. The specimens were stained with hematoxylin-eosin and examined under ×1000 magnification of a light microscope. The preparations were evaluated as for thickness of newly formed cartilage (in mm), inflammatory reaction and presence of macroscopic perforation.

**Results**

During the study period, one rabbit from each group died and so excluded from the study. Areas of cartilage tissue were larger in the fascia lata group. In the middle of these areas basophilically stained chondroblasts and in the periphery acidophilic chondroblasts were observed. Newly formed cartilage and mature cartilage were distinctly separated from each other (Fig. 2). However, in the control group neochondrogenesis was restricted to small patchy areas. Relatively smaller number of chondrocytes was discerned around chondrocytes (Fig. 3). In both groups, chondroblasts were smaller than chondrocytes and also lacunes in mature cartilages were bigger than those in the areas of neochondrogenesis.

Median areas of neochondrogenesis in the control and fascia lata groups were 0.59 (range: 0.22-0.89) mm and 1.35 (range: 0.36-2.32) mm, respectively (Table 1). Thickness of the cartilage in the fascia lata group was significantly greater than that of the control group (M-W-U=15,000, p<0.05). Inflammatory reaction in the area of defect (Fig. 4) was detected in four (4/9; 44.4%) rabbits in the control and six (6/9; 66.6%) rabbits in the fascia lata group (Table 1) without any significant difference between groups (Fischer’s exact=0.637, p<0.05). Necrotic areas (Fig. 5) were observed in four (4/9; 44.4%) rabbits in the control and three (3/9; 33.3%) rabbits in the fascia lata group without any significant difference between both groups (Fischer’s exact=0.637, p<0.05) (Table 1). Percentages of perforation were 55.5% (5/9) and 44.4% (4/9) in the control and fascia lata groups, respectively (Table 1), without any significant difference between the groups (Fischer’s exact=0.637, p<0.05).

**Discussion**

Various studies have demonstrated the presence of perichondrial induction of neochondrogenesis.\textsuperscript{14} Perichondrium adhered to the cartilage is not active. However, when it is separated from the cartilage, it becomes highly chondrogenic.\textsuperscript{15} Perichondrium consists of two distinct layers as the outer thick fibrous layer and inner and thinner progenitor layer. It is a known fact that perichondrium also contains specific chondrocyte precursors.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Thickness of the cartilage (mm)</th>
<th>Inflammatory reaction</th>
<th>Necrotic areas</th>
<th>Percentages of perforation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fascia lata</td>
<td>1.35 (0.36-2.32)</td>
<td>66.6% (6/9)</td>
<td>33.3% (3/9)</td>
<td>44.4% (4/9)</td>
</tr>
<tr>
<td>Control</td>
<td>0.59 (0.22-0.89)</td>
<td>44.4% (4/9)</td>
<td>44.4% (4/9)</td>
<td>55.5% (5/9)</td>
</tr>
</tbody>
</table>

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Fig. 2. Neochondrogenesis in the fascia lata group (HE ×100).

Fig. 3. Neochondrogenesis in the control group (HE ×100).
chondrial grafts harvested from rabbits and examined histomorphological changes. On 7th day, they detected chondrocytic differentiation, mRNA 2 and related proteins in the inner layer of the perichondrium without any change in the fibrous tissue. Lopez Aguado et al. demonstrated neo-chondrogenesis and cartilage formation on the operation site following resection of subperichondrial layer from the septum. Neo-chondrogenesis and cartilage formation developed at regions far away from each other and transition zone between both tissues was made up of intermediate cells with chondrocytic and myofibroblastic characteristics. Since scar tissue starts to develop within the first 24 hours of the traumatic incident, it generally proceeds faster than neo-chondrogenetic repair. Therefore, in order to achieve maximum cartilage repair, it is mandatory to restrict bleeding and inflammation to minimize scar tissue formation. Proliferation and differentiation of chondrocytes, as seen in many cells, require an attachment site. Besides this attachment can facilitate the response of these cells to other regulatory factors.

Various studies have been performed to accelerate neo-chondrogenesis. One part of these studies is related to autologous mesenchymal cell implantation. The authors think that adequate number of repair cells should be available in the areas of defect for the repair of the cartilage tissue. Indeed, repair cells have the capability of repairing various cell types. When implanted in the defect, chondrogenic process rapidly starts to take effect. However this process may fail to induce development of complete cartilage tissue which might stem from inadequate amount of repair cells implanted in the area of defect.

In studies targeting tissue regeneration, various materials have been used as regenerative tissue. Among them fascia lata is widely used. In a study performed with knee joints of the sheep, fascia lata was placed on the defective cartilage tissue and remodeling characterized by the increase in the number of fibrils, thickness of the collagen fibers of the fascia lata and foci of neo-chondrogenesis on the area of defect was detected. As a regenerative membrane, fascia lata was interposed between endosseous implant and bone allograft as a regenerative barrier membrane with resultant successful outcomes.

Fascia lata is a partially acellular mesenchymal tissue containing longitudinal elastic and collagen fibrils with interspersed fibroblasts. Relative cellular deficit of fascia lata is its most important characteristic feature, which permits realization of successful tissue transplantation without any need for comparison and matching between donor and recipient sites. In a study on nasal septal perforation, fascia lata was used due to the requirement for a very low metabolic activity which allows tissue growth and vascularization on its surface. In studies performed, fascia lata has been recommended as an non-antigenic ideal donor graft.

In this study autologous fascia lata graft was used. Fascia lata is easily available with low donor area morbidity. New cartilage formation was observed both in the control and the fascia lata groups. As supported by the studies performed, the rate of neo-chondrogenesis was statistically significantly higher in the control group. (M-W-U=15,000, p<0.05). The difference between both groups as for the rates of inflammatory reaction, necrosis and proliferation was not statistically significant.

**Fig. 4.** Diffuse cartilage proliferation (short arrow) and inflammatory reaction in the surrounding tissue (long arrow) in the fascia lata group (HE ×100).

**Fig. 5.** Cartilage necrosis (long arrow) and new cartilage tissue (short arrow) in the fascia lata group (HE ×100).
When evaluated together with literature findings, the results of this study support the assertion that perichondrial neochondrogenesis can greatly contribute to the repair of cartilage defects and use of fascia lata in the acceleration of this process can result in successful outcomes.

**Conflict of Interest:** No conflicts declared.

**References**


