Topographic Analysis of the Distal Femoral Condyle Articular Cartilage Surface: Adequacy of the Graft from Opposite Condyles of the Same or Different Size for the Osteochondral Allograft Transplantation

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Abstract
Objective. To analyze the topography of the opposite condyle to treat focal femoral condyle articular defects with an osteochondral allograft (OCA). Design. Three groups were created: Group 1, same condyle with same width; Group 2, opposite condyle with same width; Group 3, opposite condyle with different width. Computed tomography (CT) of 22 cadaveric femoral hemi-condyles was used to create 3-dimensional CT models that were exported into point-cloud models. Three zones of the donor condyle (anterior, middle, and posterior) were quantified. Four defect sizes were created (15, 18, 23, 25 mm) at the weight-bearing region. The defect was moved throughout each donor condyle zone and the least distance was calculated, defined as the shortest distance between the defect and the donor condyle. Results. The mean least distance increased with larger defect size in all groups, yet there was a less than 0.2 mm difference in the least distance among defect sizes. The 15, 18, and 23 mm defect models in group 1 exhibited greater least distances at the anterior than middle and posterior zones. The 15 mm defect model exhibited greater least distance at the anterior zone than posterior zone in group 3. However, there was a less than 0.05 mm difference in the mean least distance between zones. There was no significant difference in the least distance between groups. Conclusion. OCAs from opposite condyles yield similar topographic matching to OCAs from the same condyles, suggesting that opposite condyles can be utilized. Clinical correlation and outcomes are necessary.

Keywords
distal femoral condyle, osteochondral allograft, topography

Introduction
Focal chondral defects of the knee are prevalent and are a significant source of pain and morbidity in the young, active population. Studies have reported localized, full-thickness chondral lesions in 5% to 20% of all patients undergoing arthroscopic evaluation.¹,² Treatment options for focal chondral defects in younger patients include debridement, microfracture, osteochondral autograft transplantation, autologous chondrocyte implantation, and osteochondral allograft (OCA) transplantation.³⁻⁸ OCA transplantation has been performed for the past few decades, with a goal of restoring the articular surface topography. The grafts can be fashioned in one or more plugs to cover the entire defect using a press fit technique or a shell allograft can be employed. Its popularity has increased as a result of its clinical success and expanding indications for the procedure. However, limited tissue availability has slowed the widespread use of fresh OCA.⁹,¹⁰

When OCAs are indicated, size-matched donor condyles are ordered based on posteroanterior flexion weight-bearing radiographs of the knee. Other measurement methods exist as well including magnetic resonance imaging (MRI) data

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in the sagittal plane. This graft is typically from the same side (right or left), same condyle (medial or lateral), and similar size (width of the affected condyle). As obtaining a donor can take from months to even a year, OCA availability is a major limitation to more widespread utilization of this technique. Therefore, allowing for the donor condyle to be from the opposite condyle and vary in width may allow increased access for patients.

Matching native topographic anatomy is paramount when performing osteochondral graft transplantation. Failure to anatomically match the articular surfaces may lead to poor clinical results. While work has been done to analyze the best location of osteochondral autograft harvests based on topographic matching, few studies have been applied to OCAs. Mologne et al. investigated surface matching of opposite condyle OCAs for a 20 mm condylar defect utilizing micro-computed tomography. However, no study to date investigated the topographic matching of OCAs based on the condylar size and the graft location. The purpose of this study was to analyze the topographic matching of the opposite condyle to treat focal distal femoral condyle articular defects with a circular OCA plug. We hypothesized that OCA grafts obtained from opposite condyles would maintain precise surface topographic matching despite different sizes and locations.

**Materials and Methods**

**Specimen Preparation**

Cadaveric tissue was procured from a donor tissue bank (AlloSource, Denver, Colorado) after institutional review board exemption was approved. Twenty-two femoral hemi-condyles with intact articular cartilage surfaces from 22 donors were prepared according to the company’s standard protocol for planned implantation (Table 1). Of these, 11 were medial femoral hemi-condyles (MFCs) and 11 were lateral femoral hemi-condyles (LFCs). Condylar width was measured with a digital micrometer 10 mm distal to the most superior aspect of the notch, which is the same method used by donor tissue suppliers (Fig. 1). Samples that were within 0.5 mm in condylar width were considered to be same size. Three groups were created from combinations of the 22 hemi-condyles on the basis of the difference between condylar width: Group 1 (n = 12), combinations of same condyle with same size (MFC donor matched to MFC recipient and LFC donor matched to LFC recipient); Group 2 (n = 8), combinations of opposite condyle with same size (LFC donor matched to MFC recipient and MFC donor matched to LFC recipient); and Group 3 (n = 8), combinations of opposite condyle with different size (Table 2). The recipient and donor condyles were matched by ipsilateral side (right and left).

![Figure 1](image1.png) **Figure 1.** Condylar width was measured with a digital micrometer 10 mm distal of the most superior aspect of the notch.

**Three-Dimensional CT Computer Model Creation of the Femoral Articular Surfaces**

Computed tomography (CT) (BrightSpeed, GE Healthcare, Wauwatosa, Wisconsin) images were acquired in the coronal or sagittal planes using 0.625 mm contiguous slices (120 kV, 437 mA, 96 mm field of view, 512 × 512 matrices) (Fig. 2). Three-dimensional (3D) CT models of the femoral hemi-condyles were then created and exported into point-cloud models using a 3D reconstruction software program (Mimics, Materialise Inc., Leuven, Belgium). The femoral hemi-condyle articular surface model was created by segmentation of the area covered by the articular cartilage. A local coordinate system of the articular surface was created as follows. A para-coronal plane including the most distal point was defined as a coronal plane (blue plane in Fig. 3). A para-transverse plane including the most posterior point was defined as a transverse plane (red plane in Fig. 3). A para-sagittal plane including the centroid of the articular surface mode was defined as a sagittal plane (green plane in Fig. 3). An intersection of these planes was defined as an origin of the local coordinate system. The Cartesian coordinate system was further transferred to a spherical coordinate system with the most distal point as the “South Pole.”

**Three-Dimensional CT Computer Model Creation of the Femoral Hemi-Condyle Articular Surface Defect**

Circular articular surface defect models were created in each recipient condyle with 4 different diameters (15.0 mm, 18.0 mm, 23.0 mm, and 25.0 mm) at the most distal region in each femoral hemi-condyle (Fig. 4). The point-cloud data within a distance of radius of each defect (i.e., 7.5 mm, 9.0 mm,
Zoning of the Femoral Condyle Articular Surfaces

The articular surface was first divided into the peripheral zone and the central zone. The margin of the central zone was defined such that its shape was concentric to the outer margin. The distance between the central margin and the outer margin was the radius of the defect model to be analyzed (Fig. 5A). Therefore, a defect centered beyond the central zone that is at least partially outside the articular cartilage surface was not included in this study. The central zone was further divided into 3 zones defined by an angular parameter in the spherical coordinate of each point of the articular surface model (Fig. 5A and B): anterior zone (blue zone), over 45° anteriorly; middle zone, within 45°; and posterior zone (red zone), over 45° posteriorly.
Three-dimensional surface topography was compared between the defect model and the femoral hemi-condyle articular surface for 112 defect-femoral hemi-condyle comparative combinations (4 sizes of defect models × 28 defect-femoral condyle models combinations). The defect articular surface model was virtually placed on the articular cartilage surface of the donor condyle so that the centroid of the defect merged at a point in the central zone (anterior, middle, and posterior zones) of the femoral condyle (Fig. 7). Defect model-donor condyle 3D articular cartilage surface topography matching was performed using the previous procedures. The “footprint” underneath the defect model was defined as a circle with the radius of the defect. Orientation of the defect model was adjusted so that its axis matched that of the footprint area on the femoral condyle. Distances between the defect model and the donor surface were calculated in 3D space. Least distance was defined as the shortest distance from the point in question to the footprint, where a perfect congruent match would equal a least distance of 0 mm for the given data points on the simulated articular cartilage surfaces. A mean value of the least distances was calculated for each position and each orientation of the defect model (Fig. 6).

The defect model was then rotated 360° around the axis perpendicular to the articular cartilage surface in 1° increments and the least distance was calculated at each rotating angle. The position of the defect model was then moved throughout the entire femoral condyle articular cartilage surface model and the least distance was calculated at each position (Fig. 7). In this procedure, the centroid of the defect model was located within the central zone. For each position that the simulated graft was moved, the articular surfaces were compared for match with respect to all data points.

Statistical Analysis

The data were analyzed using Excel 2010 (Microsoft Corp, Redmond, Washington) and JMP software (v12.0, SAS
Institute, Cary, NC). Statistical analysis proceeded with analysis of variance (ANOVA) to compare the condylar width among groups and the effect of defect size within each group followed by Fisher post hoc analysis. Group 1 was then used as a control group to compare mismatch when looking at the opposite condyle (Group 2) or different size of condyle (Group 3) as the variable. This analysis was performed with unpaired Student’s t test. The data were presented as mean ± standard deviation, and the level of significance for all analysis was set at $P < 0.05$.

### Results

The mean condylar width of the MFCs measured 24.8 ± 1.5 mm (range = 22.5-27.0 mm; mean age = 24.4 ± 5.3 years), whereas the LFCs measured 28.9 ± 2.5 mm (range = 24.1-33.0 mm; mean age, 23.6 ± 8.1 years) (Table 1).

![Figure 7](image)

**Figure 7.** A 3-dimensional representation of the distance distribution of the 18 mm defect model in Group 3. Gradient color code represents from 0.4 mm to 0.8 mm. Blue represents penetration into the femoral condyle surface, whereas red indicates prominence with respect to the femoral condyle surface.

<table>
<thead>
<tr>
<th>Specimens</th>
<th>MFC</th>
<th>LFC</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>Age, years</td>
<td>24.4 ± 5.3 (14-33)</td>
<td>23.6 ± 8.1 (12-33)</td>
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<td>5/6</td>
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<tr>
<td>Condylar width, mm</td>
<td>24.8 ± 1.5 (22.5-27.0)</td>
<td>28.9 ± 2.5 (24.1-33.0)</td>
</tr>
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MFC = medial distal femoral condyle; LFC = lateral distal femoral condyle. Results are expressed as mean ± standard deviation.

Table 1. Demographic Data and Characteristics of Specimens.

Within Group 1, the mean mean difference in the condylar width in Group 3 was significantly greater than that in Group 1 ($P < 0.01$) and Group 2 ($P < 0.01$).

The mean least distances in the topographic articular cartilage surface matching are summarized in Table 3. Overall, the mean least distance of each combination was less than 1 mm in all groups. However, the mean least distance increased with increased defect size within each group. Within Group 1, the mean least distance significantly increased with increased defect size when comparing defect size 15 mm to 18 mm to 23 and 25 mm ($F[3, 44] = 50.011, P < 0.001$), while there was no significant difference between defects sized 23 mm and 25 mm ($P = 0.696$). Within Groups 2 and 3, similarly, the mean least distance significantly increased with increased defect size (Group 2: $F[3, 28] = 54.048, P < 0.001$; Group 3: $F[3, 28] = 25.617, P < 0.001$).

The mean least distance was compared among zones of donor condyles (anterior, middle, and posterior zones) within each group. Within Group 1, the 15 mm, 18 mm, and 23 mm defect models exhibited significantly greater mean least distance at the anterior zone than the posterior zone (15 mm, $P < 0.01$; 18 mm, $P < 0.01$; 23 mm, $P < 0.01$) and the middle zone (15 mm, $P = 0.031$; 18 mm, $P < 0.01$; 23 mm, $P = 0.049$). However, there was less than 0.05 mm difference among zones of donor condyles. Within Groups 2 and 3, similarly, there was less than 0.05 mm difference among zones of donor condyles. However, the 15 mm defect model exhibited significantly greater mean least distance at the anterior zone than the posterior zone within Group 3 ($P = 0.02$).

When comparing same and opposite donor hemi-condyles (Group 1 and Group 2), there was no significant
difference in the mean least distance for any defect sizes or donor zones. When comparing same and different sized condyles (Group 1 and Group 3), there was no significant difference in the mean least distance for any defect sizes or donor zones.

**Discussion**

The findings of this study are that OCAs from opposite condyles provide similar topographic matching at the most distal region of the recipient condyle regardless of different sizes and different locations. Data from this study demonstrates that our “best efforts” when using the same condyle yields a fit of 0.45 to 0.62 mm. This mismatch increased with increasing defect size up to 25 mm. In terms of the graft harvest location, the best-fit hierarchy was as follows: the posterior zone was better than the middle zone, which was better than the anterior zone. However, the largest mismatch difference was between the posterior 15 mm defect (0.45 mm) and the anterior 25 mm defect (0.62 mm, difference 0.17 mm). The opposite condyle of the same or different size showed similar tendency of surface matching when comparing defect sizes or locations. However, most of the mismatch was not significantly different among locations. Importantly, there were no significant differences of mismatch in any sizes and any locations between the same condyle and the opposite condyle with the same or different size.

The goal of OCA transplantation is to restore normal biologic and biomechanical properties of intact articular cartilage with normal joint function. Many factors, such as patient age, chondrocyte viability of the recipient and the graft, and surgical procedures affect the outcome after OCA transplantation. The surface matching between the graft and the recipient is an important factor in achieving good clinical results after OCA transplantation. An ideal match for resurfacing of the articular cartilage would be a perfect congruous relationship between the graft and the recipient. Failure to anatomically match surfaces can lead to poor clinical results, which are caused by eccentric loading and altered contact pressure leading to early failure of the graft.

Topographic analysis is a crucial method to elucidate the surface geometry of the human body. Furthermore,
topographic anatomy can help surgeons achieve successful osteochondral graft transplantation. Previous studies mostly focused on topographic anatomy for osteochondral autografting. However, there are few studies that evaluate topographic matching of the distal femoral condyle to find the optimal donor site for OCA transplantation. Typical clinical practice involves obtaining OCAs with the same side, condyle, and size, which can result in limited graft availability and increased patient wait time to find a matching graft. Opposite condyles for the affected lesions may be a good graft source, allowing for significantly increased graft availability for OCAs. However, previous studies showed that the anatomical morphologies of MFCs and LFCs differ in shape, curvature, and size. Therefore, topographic analysis of the distal femoral condyle needs to be clarified before potentially utilizing the opposite condyle as OCAs. Mologne et al. previously investigated OCA surface matching with an MFC or LFC graft into a 20 mm MFC defect model. This study showed that the overall articular cartilage mismatch of 0.63 mm for area and 0.47 mm for step-off, with no significant differences between the MFC and LFC graft. In addition, the original articular cartilage surface contour matched within ±1 mm in 87.4% of the MFC grafts and 87.7% of the LFC grafts. These results were consistent with our study. Furthermore, our study evaluated topography matching using different size of donor condyle because it is difficult to adjust the size of the opposite condyle. Our results suggest that OCAs from the opposite condyle can yield an acceptable articular cartilage surface matching regardless of the size of donor condyle.

The clinical relevance of the mismatch, which affects clinical results for OCA transplantation, might be analyzed in the setting of previous works. Nakagawa et al. evaluated clinical results and second look arthroscopy after transplantation. They showed that the proud plugs gave poor clinical results but recessing less than 1 mm promoted acceptable cartilage healing and lead to good clinical results. Therefore, while recessing plugs up to 1 mm may be tolerated, proud plugs are likely not as well tolerated. On the other hand, a study using a sheep model showed that the flush and 1 mm recessing of osteochondral grafting obtained a good smooth surface compared with 2 mm recessing. In addition, another study using a rabbit model showed that grafts sunk by 2 mm lead to cartilage necrosis and fibrous overgrowth. Therefore, incongruities ranging from flush to proud or recessing of less than 1 mm can be tolerated for grafting. Our results showed that OCAs from opposite condyle had less than 1 mm of mismatch in any defect sizes and any graft locations.

With regard to biomechanical properties of graft incongruity surrounding the recipient, D’Lima et al. showed that grafts proud by 0.5 mm increase peak contact stress by 2 times compared to intact cartilage. This study showed that a graft that is even 0.25 mm proud over the recipient surface increased stress and strain. In addition, recessing of the graft increased the stress and strain in the recipient area surrounding the graft. Koh et al. demonstrated that peak contact pressure significantly increased with plugs elevated 1.0 and 0.5 mm above the surrounding surface, and that plugs sunk 0.5 and 1.0 mm significantly increased the peak contact pressure in the intact area. The peak contact stress and peak compression strain approached levels that have been shown to induce cartilage damage and cell death. Our results therefore suggest that the contact pressure may increase the same as the mismatch with increasing defect size, though a direct comparison as such is subject to speculation. However, the contact pressure could be unchanged whether grafts are harvested from the same or opposite condyles.

The strengths of this study include the computer-based computational analysis, which is based on an established 3D point-cloud creation instrument, determination of the centroid of the femoral condyle articular cartilage surface, matching of the orientation of the articular cartilage surface point clouds, and cross-referencing of each femoral condyle cloud with each defect cloud to account for any variability that may be present between combinations. Another strength is that the specimens used in this study were actual materials for OCA transplantation. So, these specimens could retain the articular cartilage surface and provide actual results of topography mapping relevant to performing OCA transplantation.

There were several limitations in the current study. First, his study focused strictly on the articular cartilage surface congruence. However, a surgeon should consider several situations of chondral defects such as different size, depth, and locations. Second, the thickness of cartilage was not considered for this analysis. The thickness of cartilage is different for MFCs and LFCs, and the cartilage in contact regions is thicker than noncontact regions. From a biomechanical point of view, the thickness of graft cartilage should be equal to that of recipient cartilage. Based on these limitations, further studies are needed to increase the graft availability for OCA transplantation.

In conclusion, our results indicate that OCAs from opposite condyles can yield precise surface topographic matching in the direct weight-bearing region regardless of the different size and different location of the donor condyle. These findings suggest that the utilization of the opposite condyle can increase the graft availability for OCAs’ transplantation while maintaining precise surface topographic. This is in agreement with recently published results reporting no significant difference in clinical outcomes when utilizing the opposite condyle for OCA transplantation. Future randomized trials and long-term studies are necessary to further understand the clinical effect of utilizing the opposite condyle.
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Declaration of Conflicting Interests

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Ethical Approval

Ethical approval for this study was waived by Rush University Institutional Review Board because it did not involve human subjects, only non-identifiable hemi-condyle allografts.

Informed Consent

Informed consent was not sought for the present study because it did not involve human subjects.

Trial Registration

Not applicable.

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