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Effect of 3 Preservation Methods (Freezing, Cryopreservation, and Freezing 1 Irradiation) on Human Menisci Ultrastructure

An Ex Vivo Comparative Study With Fresh Tissue as a Gold Standard

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Background: Three main meniscus preservation methods have been advocated: freezing (-80°C), freezing with gamma irradiation (-80°C 1 25 kGy), and cryopreservation (-140°C).

Hypothesis: All preservation methods will result in structural and architectural properties similar to those of fresh meniscus, defined as the gold standard.

Study Design: Controlled laboratory study.

Methods: Five human intact menisci were collected from 5 patients undergoing total knee arthroplasty. The inclusion criteria were patients \( \geq 70 \) years old with primary unilateral (medial) femorotibial knee osteoarthritis and without surgical or traumatic history on the operated knee. Four cubes (9 mm\(^3\)) were cut inside of the white, or avascular, area of each specimen’s middle horn and divided into 4 groups: “fresh” control, frozen (-80°C), cryopreserved (-140°C), and frozen 1 irradiated (-80°C 1 25 kGy). Specimens of the control group were evaluated at day 1, and specimens from the frozen, cryopreserved, and frozen 1 irradiated groups were evaluated after 1 month of storage. Evaluation was performed with electron microscopy according a validated protocol to analyze (1) mean diameters of the collagen fibers in longitudinal and transverse sections in 5 points per section and (2) validated architectural scores.

Results: No significant difference was found between the control and cryopreserved groups regarding mean transverse and longitudinal diameters (transverse: 95.39 ± 15.87 nm vs 99.62 ± 19.23 nm, \( P = .1 \); longitudinal: 96.31 ± 13.96 nm vs 94.57 ± 16.42 nm, \( P = .1 \)). Significant differences were found between the control and frozen groups (transverse: 95.39 ± 15.87 nm vs 70.20 ± 13.94 nm, \( P \leq .001 \); longitudinal: 96.31 ± 13.96 nm vs 71.28 ± 16.64 nm, \( P \leq .001 \)) and the control and frozen 1 irradiated groups (transverse: 95.39 ± 15.87 nm vs 63.16 ± 15.57 nm, \( P \leq .001 \); longitudinal: 96.31 ± 13.96 nm vs 60.96 ± 14.8 nm, \( P \leq .001 \)). Regarding architectural score calculation, there were significant differences between the control and frozen groups (4.56 ± 1.3 vs 2.36 ± 1.4, \( P = .02 \)) and the control and frozen 1 irradiated groups (4.56 ± 1.3 vs 1.4 ± 0.9, \( P = .02 \)).

Conclusion: Cryopreservation is the only method that preserves fresh meniscus architectural specificities. Freezing and freezing 1 irradiation methods modify histologic properties of meniscal allograft. Irradiation deeply alters diameters and the organization of collagen fibers, and this method should be used with caution to preserve and sterilize meniscus tissue.

Clinical Relevance: The results of our study exhibited detrimental effects of simple freezing and freezing 1 irradiation on the collagen network of sample meniscus. If those effects occur in menisci prepared for allograft procedures, important differences could appear on the basis of the preservation procedure in terms of the graft’s mechanical properties and, thus, the patient’s outcomes.

Keywords: meniscus allograft; irradiation; cryopreservation; freezing; histologic; collagen

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to restore joint knee mechanics and potentially slow the onset of osteoarthritis. Verdonk et al. described midterm results showing a significant improvement in patients' pain associated with 5-year allograft survival of 85%, as other investigators. As these results imply a meniscal allograft equivalent to the native fibrocartilage, graft preservation methods play a vital role in the clinical and biological success of meniscal allograft techniques. Three main meniscus preservation methods were advocated: freezing, freezing with gamma irradiation, and cryopreservation. Simple freezing brings the material to 280°C, but deep modification of the collagen network and global architecture was reported with this preservation. Freezing associated with irradiation allows a deep sterilization of the graft but also compromises allograft microarchitectural and biomechanical properties. Cryopreservation techniques require cryoprotectants (dimethylsulfoxide) and bring the tissue to 2145°C. Gelber et al. demonstrated that cryopreservation was superior to freezing to maintain the meniscus ultrastructure during the storage process. To date, studies investigating human meniscus preservation had 2 main limitations: first, they did not compare preservation processes with the same meniscus sample; second, conserved tissues are not compared with fresh meniscus samples. As meniscus ultrastructure depends on various demographic and pathologic confounding factors, comparing potential detrimental effects of various conservation processes should be performed on "identical" tissue samples. Therefore, we aimed to analyze and compare 3 preservation processes with a "fresh" tissue control group using samples harvested from the same human meniscus. We hypothesized that all preservation methods would result in structural and architectural properties similar to those of fresh meniscus, defined as the gold standard. We aimed to estimate the effects of these methods on meniscus ultrastructure by using electronic microscopy to compare collagen fiber diameters in longitudinal and transverse sections and by calculating a validated architectural score per sample.

METHODS

After local review board approval, 5 human lateral menisci were collected from patients who had total knee arthroplasty between September and October 2017. All patients signed an informed consent form before their inclusion in our study. Inclusion criteria were as follows: patient aged ≤70 years undergoing total knee arthroplasty because of isolated internal femorotibial arthritis or femoropatellar and internal femorotibial joint degeneration (but with an external femorotibial compartment graded Kellgren and Lawrence ≥2) and no surgery, trauma, or developmental disease of the operated knee. Table 1 summarizes patients' characteristics.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age, y</th>
<th>Sex</th>
<th>Weight, kg</th>
<th>Size, cm</th>
<th>BMI, kg.m²</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>61</td>
<td>M</td>
<td>76</td>
<td>181</td>
<td>22.6</td>
</tr>
<tr>
<td>2</td>
<td>67</td>
<td>M</td>
<td>82</td>
<td>184</td>
<td>24.2</td>
</tr>
<tr>
<td>3</td>
<td>62</td>
<td>F</td>
<td>67</td>
<td>172</td>
<td>22.1</td>
</tr>
<tr>
<td>4</td>
<td>60</td>
<td>F</td>
<td>60</td>
<td>162</td>
<td>22.9</td>
</tr>
<tr>
<td>5</td>
<td>69</td>
<td>M</td>
<td>77</td>
<td>178</td>
<td>24.3</td>
</tr>
</tbody>
</table>

*BMI, body mass index.

Sample Creation

The anterior and posterior horns were sectioned to retain only the median horn of the meniscus. Four cubes (9 mm³) were cut inside of the white, or avascular, area of each specimen and divided into 4 groups: "fresh" control, freezing, cryopreserved, and freezing 1 irradiation. Histologic fixation of the control specimens was performed directly in the operating theater under a hood, which consisted of the following steps based on a previously validated protocol:

1. Five-minute rinsing with 0.1M cacodylate buffer.
2. Fixation in a solution of 2.5% glutaraldehyde in cacodylate buffer for 1 hour.
3. Rinsing 3 times for 10 minutes with 0.1M cacodylate buffer.
4. Postfixation with a solution of 2% osmium tetroxide in 0.1M cacodylate buffer for 30 minutes.
5. Rinsing 3 times for 15 minutes with 0.1M cacodylate buffer.
6. Progressive dehydration of samples ranging from 50% to 100% ethanol before inclusion in Spurr.
7. Resin fixation for transmission electron microscopy

Control specimens were analyzed immediately after those steps in a delay of 6 hours. The other 3 fragments were placed in physiologic saline in a cryokit and kept at 8°C until their transport to our local tissue bank (Etablissement Francais du Sang) (6 hours). The 3 samples were then prepared with the following steps: (1) graft reception in clean room (controlled atmosphere zone), (2) decontamination of the graft with an antibiotic solution (rifampicin 1 thiampenicol), (3) rinsing with 0.1M cacodylate buffer for 5 minutes, and (4) bacteriologic sampling. Then, the 3 conservation methods were applied (1 sample for each). For the cryopreservation group, cryoprotective solution (10% of DMSO 1 SCOT 30) was added, and the bag was vacuumed to extract the residual air and the temperature progressively decreased (staring at 24°C and then decreasing at 22°C per minute to 240°C and then 25°C per minute to 2140°C). Samples were stored in a nitrogen tank in vapor phase at 2145°C. For the freezing group, a simple freezing

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process was used by progressively decreasing the temperature (starting at 24°C and then decreasing at 22°C per minute to 240°C and then 25°C per minute to 280°C). For the freezing 1 irradiation group, a simple freezing process with a progressive decrease in temperature was performed (starting at 24°C and then decreasing at 22°C per minute to 240°C and then 25°C per minute to 280°C). The grafts were then transported in a dry ice–controlled container (stored at 280°C) to be irradiated by gamma ray by IONISOS. The doses received ranged between 23.9 and 26.5 kGy (2.4–2.6 Mrad). After this treatment, the samples were again stored at 280°C until analysis.

After 1 month, the samples were transported to the electronic microscopy laboratory to be analyzed. The fixing steps (steps 1–7) were the same as those described for the control group. For all samples, ultrafine 60-nm sections were made with an Ultracut ultramicrotome (Reichert-Jung); contrast of the sections was made with uranyl acetate and lead citrate. Ultrastructure pictures were obtained with a transmission electron microscope (JEM 1400; JEOL) at 80 kV with a Megaview III camera and iTEM Five software (SIS Imaging). For each sample, we took 10 pictures with a magnification of 60003 and 40,0003. The longitudinal (Figure 1) and transverse diameters of the collagen fibers were measured at 70 points on a picture taken at 40,0003 magnification. The collagen meniscal architecture scoring system was calculated for each sample on 5 pictures with 60003 magnification (Table 2).7

Statistics

Before the initiation of the study, a sample analysis estimated that 5 samples for each group was necessary to be

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**Table 2**

<table>
<thead>
<tr>
<th>Collagen Meniscal Architecture Scoring System</th>
<th>0 Point</th>
<th>1 Point</th>
<th>2 Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disrupt/periodicity</td>
<td>Mild</td>
<td>Moderate</td>
<td>Severe</td>
</tr>
<tr>
<td>Intrafibrillary edema</td>
<td>No</td>
<td>Yes</td>
<td>—</td>
</tr>
<tr>
<td>Packing</td>
<td>High density</td>
<td>Intermediate</td>
<td>Low density</td>
</tr>
<tr>
<td>Banding</td>
<td>Yes</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Fibril size variability</td>
<td>Low</td>
<td>High</td>
<td>—</td>
</tr>
</tbody>
</table>

powered to distinguish 10 6 10 nm regarding collagen transverse or longitudinal diameters.4,5,7

Patient characteristics were expressed with the appropriate descriptive statistics for the type of variables. Descriptive statistics included mean with SD or median with interquartile range as appropriate for continuous variables. The intraclass correlation coefficient with 95% CIs was calculated to assess intra- and interobserver reproducibility for the transverse and longitudinal diameter values. Student t tests were used to compare the distribution of continuous parameters between groups (or the Mann-Whitney test when the data were not normally distributed or when the homoscedasticity assumption was rejected). All reported P values were 2-sided, with a significance threshold of .05. Statistical analyses were performed with SPSS/JMP software (v 13; Microsoft).

**RESULTS**

**Transverse Diameter**

No significant difference was found between the control and cryopreserved groups regarding mean transverse
Collagen Meniscal Scoring System

The mean values of the collagen meniscal architecture scoring system are summarized in Table 5. No difference was found between the control and cryopreserved groups (4.5 6 1.3 vs 4.3 6 1.6 points, P = .9). There were significant differences between the control and frozen groups (4.5 6 1.3 vs 2.3 6 1.4 points, P = .02) and between the control and frozen 1 irradiated groups (4.5 6 1.3 vs 1.4 6 0.9, P = .02).

DISCUSSION

The main finding of the current study was that cryopreservation preserves meniscus histologic ultrastructure, unlike simple freezing or freezing 1 irradiation. We rejected our hypothesis that all preservation methods will result in structural and architectural properties similar to those of fresh meniscus. Cryopreservation does not entail a significant modification in terms of collagen fiber diameter or architectural organization as compared with fresh tissue. We did find, however, significant differences regarding those 2 measurements when we compared freezing and freezing 1 irradiation processes with fresh tissue or cryopreserved samples. Irradiation was the more detrimental process in terms of tissue preservation, as all of our measurements were inferior to those taken for fresh tissue, cryopreservation, and freezing samples.

Our study is limited in that our patients are older than usual donors (mean age: 63.8 years in our study vs 53.5 in the register). Because of this, menisci evaluated during our study could be considered a limitation. We supposed that most of the ultrastructure alteration related to the preservation method occurred during the decreasing of temperature steps, including direct chemical effects, ice formation, and dehydration, as described by Pegg. Our evaluation method involved manual measurement of
TABLE 5
Collagen Meniscal Architecture Scoring Comparison Between Preserved and Control Samples

<table>
<thead>
<tr>
<th>Group</th>
<th>Score, Mean 6 SD</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control vs</td>
<td>4.5 ± 6.1.3</td>
<td>0.89</td>
</tr>
<tr>
<td>Cryopreservation</td>
<td>4.3 ± 6.1.6</td>
<td>0.9</td>
</tr>
<tr>
<td>Frozen</td>
<td>4.3 ± 6.1.4</td>
<td>0.02</td>
</tr>
<tr>
<td>Irradiation</td>
<td>2.9 ± 6.1.9</td>
<td>0.02</td>
</tr>
</tbody>
</table>

collagen fiber diameters; as such, 1 senior technician trained in transmission electron microscopy carried out the entire collection of data and was blinded regarding the preservation process of the specimens. The measurements were always made at the same magnification (40,000x) and at 70 random points of the picture, with consideration to only the most circular fibers for the transverse diameter and the most linear for the longitudinal diameter. Our protocol did not aim to investigate cell viability. It was impossible for us to perform both ultrastructure analysis and flow cytometric cell counts on the same sample. Gelber et al were the first to demonstrate that cell survival was possible in cryopreserved samples, but in a study of 15 meniscal goat allografts, Fabbriani et al showed that even if cryopreservation made it possible to maintain partial cell viability in the tissue, the morphologic and biochemical characteristics of the graft were not improved. Finally, our sample size might seem low, but the numbers of specimens included was decided before the initiation of the study to compare groups upon collagen thickness evaluation.

Despite these limitations, our study is the first to directly compare the influence of preservation processes on 4 samples prepared from the same meniscus, permitting us to assert that the differences observed are mostly related to preservation methods and their specificities more than demographic confounding factors.

We used the collagen meniscal scoring system by Gelber et al., to compare our results with the existing literature. Gelber et al. advocated the superiority of cryopreservation on meniscus ultrastructure as compared with freezing, with 2 studies corroborating our results, even if they did not use a control group. In another study, Vangsen et al. demonstrated that the use of gamma irradiation caused clear alteration of musculoskeletal tissues mechanical properties.

Bone sample load-to-failure behavior was significantly lower when exposed to 0.3 Mrad of irradiation. Fideler et al. determined that 0.2 Mrad of irradiation of bone-patellar tendon–bone allografts adversely affected 4 of the 7 structural properties that were analyzed. Thus, they found that all structural parameters were deeply affected at irradiation levels of 3.0 and 4.0 Mrad. This conclusion is supported by our data: we found the highest rate of collagen disorganization for samples exposed to 2.5 Mrad (25 kGy: freezing 1 irradiation group). Advantages in terms of sterilization allowed by gamma irradiation should be balanced with those deep architectural and potentially mechanical consequences.

With regard to the meniscus, Lewis et al. studied 7 human menisci and demonstrated that samples that underwent a single freeze-thaw cycle had a significantly higher Young modulus than did those undergoing multiple freeze-thaw cycles (Young modulus: 1.2310^5 for 1 cycle vs 8.5310^4 for multiple cycles).

Our histologic results must be confirmed by mechanical trials to better understand potential consequences of ultrastructure alteration on meniscal graft mechanical properties. In the same manner, only in vivo studies will be able to assess the real clinical relevance of our ex vivo conclusion.

CONCLUSION
Cryopreservation is the only method that preserves fresh meniscus architectural specificities. Freezing and freezing 1 irradiation methods modify histologic properties of meniscal allograft. Irradiation deeply alters diameters and organization of collagen fibers, and this method should be used with caution to preserve and sterilize menisci tissue.

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