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Comparisons of Efficacy between Autograft and Allograft on Defect Repair *In Vivo* in Normal and Osteoporotic Rats

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**Introduction.** In the field of orthopaedic surgery, the use of osteogenic material in larger defects is essential. Autograft and allograft are both known methods, and autograft is believed to be the best choice. But autograft is associated with additional invasive procedures which can prove difficult in fragile patients and can cause local side effect after bone harvest. For feasible purposes, the use of allograft is hereby rising and comparing efficacies, and the differences between autograft and allograft are essential for the clinical outcome for the patients.

**Method.** 24 female Norwegian brown rats were included, 12 normal rats and 12 induced with osteoporosis (OP). OP inducement was verified in vivo by bone volume fraction (BV/TV) at 90 days after ovariectomy (OVX). The primary surgery in each rat consisted of a 2.5 × 3 mm hole in the proximal tibia, bilaterally. Autograft and allograft were randomly allocated in the right and left tibia. After an observation of 21 days, the rats were sacrificed. Tibia samples were harvested, micro-CT scanned for bone inducement and microarchitectural properties, and then embedded for histology.

**Results.** The OP induction was verified three months after the OVX by a reduction of 68.5% in the trabecular bone BV/TV compared to normal bone. Microarchitectural analysis and histology showed no significant differences in the bone-forming capabilities between autograft and allograft in normal or osteoporotic bone after 3 weeks.

**Conclusion.** This study did not demonstrate any difference between autograft and allograft in a normal or osteoporotic rat tibial defect model after 21 days, suggesting allograft is a good alternative to autograft.

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**1. Introduction**

Bone loss and defects can be caused by trauma, infections, or following arthroplasties. They are categorized as one of the biggest clinical challenges in orthopaedic surgery [1]. Implant surgery is the general term and includes a large portion of intervention surgeries like the repair of bone defects in orthopaedic surgery, neurosurgery, oral, and maxillofacial specialties. These surgeries are estimated to be performed over 2 million times a year and hereby have a high impact on the patients and the economic outcome [2].

Autograft bone is considered as a "living" material bearing osteogenic, osteoinductive, and osteoconductive properties [3] and should provide the best treatment, where allograft consists of inactive "dead" bone with mainly osteoconductive properties [4]. However, harvesting autograft bone is an additional invasive procedure, and the amount available is often insufficient. Autograft collected from the iliac crest bone is termed autogenous iliac crest bone graft (AICBG) [3]. This procedure can be associated with morbidities, such as blood loss, donor site pain, risk of infections, and nerve injuries. Failure rates in autograft bone graft surgeries have been shown to be 50% caused by different types of harvesting, handling, implantation method used, and differences between patient conditions and bone vitality [5]. Due to these complications and high costs, allogenous bone material is often used as an alternative graft material. Allogenic bone is gathered conveniently without side effect from other patients. But this graft material has mainly osteoconductive effect [6] and has the potential risks of disease transmission, bacterial infections, autoimmune host response, and graft host nonunions. These side effects are more severe but are however extremely rare [7]. Theoretically, the gentlest treatment is the allograft. With increasing
elderly and fragile patients, it is essential to acquire knowledge by comparing them in a relevant defect model, both in regular and fragile bone structures. An alternative to diminishing the need for harvesting live bone graft is substitute [3, 8]. The approach has been wide from the use of stem cells from different tissues [9, 10] to combinations with growth factors in animal and clinical models [10, 11]. Yet, no substitute has shown a stable effect to replace all the procedures using autograft and allograft in the clinic.

Osteoporosis is an age-related rising disease and is a major public health problem related to increasing age in the population and hereby an increasing prevalence and treatment cost globally by 20-30% in 2030 [12]. Osteoporosis is a bone disease with pronounced reduction of bone mineral density. This is caused by the osteoclasts (OC) that resorb more bone than the osteoblast (OB) can produce, disturbing the stability in the bone remodelling unit (BMU), which leads to an imbalance between the bone resorption and formation and eventually bone loss [13]. This states the importance of using verified models and testing both normal and fragile bone structures for the best clinical correlation to enhance the correlation to the relevant situation.

Thus far, there is limited knowledge on the differences between autograft and allograft in bone defect repair. This study is aimed at investigating the effects of autograft and allograft on tibial defect repair in vivo in both normal and osteoporotic rats. Particularly, the in vivo longitudinal microarchitectural changes postoperatively, at 7 days, 14 days, and 21 days. The evaluation was performed with micro-CT scans at all time points and histomorphometric evaluation after euthanization at 21 days. The primary objective was to evaluate the bone formation between autograft and allograft. The secondary objective was to verify the induction of osteoporosis 90 days after the removal of the ovaries. It was hypothesized that there were no differences in the defect repairs between autograft and allograft in both the normal and the osteoporotic rats, which can help reduce the use of autograft clinically in defect models in both normal and fragile bone structures and decrease the need for additional invasive harvesting procedures.

2. Materials and Methods

2.1. Animal Model. Twenty-four female brown Norwegian inbred (BN/SsNOlaHsd) rats were included in the study. Both the normal and osteoporotic rats were 4 months of age with an average weight of 190 ± 25 g. They were housed and acclimatized 2 months prior to surgery at the Biomedical Laboratory, University of Southern Denmark. The controlled environment had a temperature of 21-28°C, humidity of 40-60%, and lights on between 6 a.m. and 6 p.m, with access to sterile water and normal or Ca-deficient diet ad libitum, respectively. The cages had sawdust flooring and bedding material. The animals were observed every day for changes in behavior or signs of discomfort by either the animal technicians or the researcher.

2.2. Animal Approval. All the experimental procedures were performed in accordance with the Danish Animal Research guidelines. This experimental protocol was approved by the Danish Animal Experiments and Inspectorates (no. 2011/561-1959). This article follows the Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines.

2.3. Bone Graft Materials

2.3.1. Autograft. The autograft bone material was harvested from bilateral rat tibiae during the drilling process of the tibia defect surgery. The bone was treated sterile, and the chips were of approximately 0.5-1 mm. In the filling of the tibia defect, the normal bone had sufficient volume from the drilling procedure to fill the gap. However, for the osteoporotic rats, the defect needed additional filling by bone material from the tail vertebrae. For the osteoporotic rats, additional bone material was acquired from the tail vertebrae. With a small incision 2 mm distal from the attachment of the tail to the body, soft tissue was removed by surgical equipment and sectioned into chips. The need for additional bone was due to the pronounced bone loss of bone mineral density after osteoporotic inducement. This method is well known for the harvest of autologous bone use in the rat model [14].

2.3.2. Allograft. This was produced from one healthy brown Norwegian female rat. It was euthanized with pentobarbital according to the guidelines, and the condyles of femur, tibia, and humerus were harvested under sterile conditions. Additional soft tissue was removed. The bone was prepared by bone mill (Ossano Scandinavia ApS, Stockholm, Sweden). Trabecular bone structure was divided into chips with a diameter of 0.5-1 mm and stored at 80°C. Before use, the frozen bone was thawed for 30 minutes and added by surgical standards to the defect. All these procedures were performed under an aseptic condition in the small animal surgery room in the Biomedical Laboratory, University of Southern Denmark.

2.4. Study Design. A paired longitudinal study design was used, with one control group and one intervention group in each tibial bone. A total of 24 Norwegian inbred (BN/SsNOlaHsd) female rats were divided into two groups of 12 each in the normal and osteoporotic bones. Autograft and allograft were blinded and randomly allocated in the right and left tibial defect and furthermore blinded in the evaluation stage (Figure 1). At 13 weeks prior to tibial surgery, 12 rats received an ovariectomy (OVX) for the osteoporosis induction. The allograft material was thawed 30 minutes before usage, and autograft was prepared and collected under the anesthesia for the primary surgery of the proximal tibia defect. At day 0, day 7, day 14, and day 21 postoperatively, micro-CT scans were performed for ongoing evaluation. At sacrifice, bilateral proximal tibial defect samples were harvested for histology and histomorphometry.

2.5. Surgical Procedures

2.5.1. Ovariectomy (OVX). 12 rats were anesthetized with 0.3 ml/100 g body weight hypnorm (VetaPharma Ltd., Leeds, UK) and dormicum (B. Braun, Melsungen, Germany) mixture, subcutaneously (s.c.) until sedation. Before surgery, the rats were given buprenorphine (Temgesic,
The back of the rat was shaved and disinfected with iodine and ethanol (70%). A sharp incision of approximately 1 cm was made over the caudal part of the back and bluntly dissected until the fascia. By penetration of the cavity, the ovary was harvested with a tweezer, and ligation was made beside the ovary with 5.0 ethilon suture and removed. The wound was closed in two layers. All operations were performed at the same time of the day and on the same location. Postoperatively, the rats were analgesized with buprenorphine (Temgesic, RB Pharmaceuticals Limited, Berkshire, UK), 0.2 ml/100 g body weight s.c. in intervals of 8 hours for the following 3 days.

After ovariectomy, the rats were given a special diet with low calcium and water ad libitum. The inducement of the osteoporotic bone structure was verified after 12 weeks, and primary tibial defect surgery could be initiated.

2.5.2. Tibia Proximal Defect. Anesthetic protocol during surgery followed the methodology of the OVX surgery.

Bilateral proximal tibial defects were made in all rats using the standard surgical procedure. Both limbs were isolated, shaved, and disinfected with iodine vet (Kruuse Vet, Denmark) and 70% ethanol. Sharp incision and blunt exploration presented the medial side of the proximal tibia, where a cylindrical defect of 2.8 mm and depth of 3 mm till opposite cortical shell were created based on the results from a pilot study. The surgery for allograft group was performed firstly, due to the collection of autografts from both legs. In total, 48 cylindrical defects were created and filled with either autograft or allograft scheduled by randomization. The wound was closed in two layers with suture 4.0. Postoperatively, the rats were analgesized with buprenorphine (Temgesic, RB Pharmaceuticals Limited, Berkshire, UK), 0.2 ml/100 g body weight s.c. in intervals of 8 hours for 4 days.

2.6. Micro-CT Scanning and Microarchitectural Analysis. The bilateral proximal tibial regions were scanned in vivo with a high-resolution microtomographic system (vivaCT 40, Scanco Medical AG, Brütisellen, Switzerland).

In vivo micro-CT scanning for evaluation of bone growth development in normal and osteoporotic rats was performed at the following 4 time points: the day prior to surgery (day 0) and then at 7, 12, and 21 days postoperatively.

Furthermore, to verify the osteoporosis induction model, a longitudinal evaluation was conducted at 3 time points: (1) before OVX surgery to establish a baseline of bone mineral density (day 90), (2) before the tibia surgery (day 7), and (3) after tibial surgery (day 0).

During in vivo scanning, the rats were anesthetized by isoflurane in a closed box system, 1 L/min oxygen and 4 ml/min isoflurane (IsoFlo vet, Abbott Laboratories Ltd, Berkshire, England) for 6 minutes, with regulation according to the reflexes of the rat. After full sedation, the rats were placed in an animal holding bed covered with a mask with a running supply of oxygen and isoflurane according to protocol. The area of interest was fixated for accurate scans. The images were scanned in a high resolution resulting in a 3D reconstruction voxel sizes of $10.5 \times 10.5 \times 10.5 \, \mu m^3$ ($2048 \times 2048 \times 2048$ pixels) for 500 slices for representative evaluation of each defect area. Scanning time for each specimen was 30 minutes.

The parameters specified by this scan included microarchitectural properties of the trabecular bone for the confirmation of osteoporosis induction and bone enhancing effect of the grafts [15, 16]. This included bone volume/tissue volume (BV/TV), structure model index, connectivity density (CD), trabecular thickness (TbTh), trabecular separation (Tb.Sp), degree of anisotropy (DA), bone surface density, bone surface-to-volume ratio (BS/TV), apparent density, and material density.

Figure 1: Illustration of study design. 24 rats were included and divided into normal or osteoporotic groups. 12 rats each. Each rat had a group of autograft and allograft in the left and right tibia for a total of 24 samples from normal bone and 24 samples for the osteoporotic bone. T1 (week 0): day for surgery; T2 (weeks 0, 1, 2, 3): micro-CT scans; T3: histology section.
3. Results

3.1. Animal Observation. In total, four of the 24 rats died during the 3 postoperative weeks, consisting of 2 from each group. Three of them died as a response to the anesthesia during in vivo micro-CT scanning, and one died caused by an infection. The remaining rats were included into the study. During the observational period of the experiment, the animals were observed daily by an animal technician or the researcher for any signs of discomfort or violation of the animal license. No significant weight change was observed in either group after tibia defect surgery. The rats exposed to OVX gained weight during the first 12 weeks from $193 \pm 8$ g to $224 \pm 44$ g ($p < 0.001$).

3.2. Three-Dimensional Microarchitectural Properties

3.2.1. Induction of Osteoporosis in Rats. After 12 weeks, OVX-treated rats had a decrease in bone volume fraction, connectivity density, bone surface density, and apparent density ($p < 0.001$) compared to normal bone. Structure model index increased from typical plate, -1.4, to typical rod, 3.3. Trabecular separation, degree of anisotropy, material density, and bone surface-to-volume ratio increased ($p < 0.001$). Trabecular thickness did not have any significant change ($p < 0.097$) (Table 1).

3.2.2. Microarchitectural Changes with Autograft and Allograft Treatments

(1) Changes in Microarchitecture in Normal Bone. BV/TV was decreased in allograft defects compared to autograft on days 0, 7, and 14 ($p < 0.05$). However, on day 21, there was no statistical difference between the two groups (Figure 3). Representative 3D reconstructions of micro-CT images have been displayed in Figure 4.

The same trend applied to the connectivity tissue with decreased value on days 0, 7, and 14 ($p < 0.05$) but no difference on day 21. Trabecular thickness was increased at all time points in the allograft groups ($p < 0.05$) and no difference in the degree of anisotropy (Figure 3).

(2) Changes in Microarchitecture in Osteoporotic Bone. BV/TV of osteoporotic bone displayed decreased bone in the allograft group on days 0, 7, and 14 ($p < 0.05$) and no difference after 21 days (Figures 4 and 5). Trabecular thickness was significantly higher in the autograft group on days 0, 14, and 21 ($p < 0.05$) but no difference on day 7. Connectivity density and degree of anisotropy had no significant difference at any time point (Figure 5).
3.3. Histology and Histomorphometry

3.3.1. Histology. New bone formation was observed in the defect area in all samples. It was not possible to distinguish the difference between remnants of graft and new bone, whereas the total bone volume within the defect area was calculated as bone within the ROI (Figure 1, T3).

3.3.2. Histomorphometry. Bone volume showed no significant difference between the autograft and allograft groups within the normal and osteoporotic bone. When comparing autograft and allograft from the normal bone with autograft and allograft in the osteoporotic bone, there was significantly decreased amount of bone at 21 days of evaluation ($p < 0.001$).

The mean formation of new bone including graft material in the defect in normal bone was 53% for autograft, whereas 51% for allograft. In osteoporotic bone, allograft had the highest bone volume with a mean of 35%, whereas autograft had 33% (Figure 6). There was no significant difference between any other measured parameters within the defect ($p > 0.05$).

### Table 1: Assessments of changes in microarchitectural properties with osteoporosis induction after OVX performed on day 90. Day 0 is the day of the tibial surgery.

<table>
<thead>
<tr>
<th></th>
<th>Bone volume fraction (%)</th>
<th>Structure model index (-)</th>
<th>Connectivity density (mm$^3$)</th>
<th>Trabecular thickness (μm)</th>
<th>Trabecular separation (μm)</th>
<th>Degree of anisotropy (-)</th>
<th>Bone surface-to-volume ratio (mm$^3$)</th>
<th>Apparent density (mg/cm$^3$)</th>
<th>Material density (mg/cm$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 90</td>
<td>54 ± 8</td>
<td>-1.4 ± 1.6</td>
<td>1324 ± 282</td>
<td>63 ± 5</td>
<td>56 ± 9</td>
<td>1.4 ± 0.03</td>
<td>19.1 ± 1.4</td>
<td>34 ± 5</td>
<td>505 ± 52</td>
</tr>
<tr>
<td>Day 7</td>
<td>19 ± 12</td>
<td>3.3 ± 0.9</td>
<td>248 ± 230</td>
<td>62 ± 3</td>
<td>160 ± 50</td>
<td>1.4 ± 0.06</td>
<td>7.8 ± 4.3</td>
<td>44 ± 4</td>
<td>234 ± 100</td>
</tr>
<tr>
<td>Day 0</td>
<td>17 ± 12</td>
<td>3.3 ± 0.9</td>
<td>210 ± 198</td>
<td>59 ± 3</td>
<td>173 ± 55</td>
<td>1.4 ± 0.58</td>
<td>6.8 ± 4.6</td>
<td>46 ± 4</td>
<td>217 ± 105</td>
</tr>
</tbody>
</table>

ANOVA $p < 0.001$ $p < 0.001$ $p = 0.097$ $p < 0.001$ $p < 0.001$ $p < 0.001$ $p < 0.001$ $p < 0.001$ $p < 0.001$

**Figure 3:** Microarchitecture properties of defect treated with autograft vs. allograft in normal bone. BV/TV: bone volume/tissue volume; CD: connectivity density; TbTh: trabecular thickness; DA: anisotropy. $p < 0.05$ is considered significant.
Normal rat: 3D reconstruction of micro-CT images at termination (day 21)

Allograft group

Autograft group

(a)

Osteoporotic rat: 3D reconstruction of micro-CT images at termination (day 21)

Allograft group

Autograft group

(b)

Figure 4: Representative 3D reconstructions of micro-CT images at termination (day 21) for both autograft and allograft groups and in normal (a) and osteoporotic (b) rats are illustrated from the same tissue as Figure 2. Red circle indicates where original defect holes were created. At the right side of each image is the newly generated bone mass within the hole. All images are displayed as whole mass and thin layer of 10 slices (105 micrometers).

BV/TV changes in autograft vs. allograft

<table>
<thead>
<tr>
<th>Day 0</th>
<th>Day 7</th>
<th>Day 14</th>
<th>Day 21</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>40</td>
<td>60</td>
<td>80</td>
</tr>
</tbody>
</table>

$\text{p} = 0.047$

$\text{p} = 0.003$

$\text{p} = 0.020$

$\text{p} = 0.101$

CD changes in autograft vs. allograft

<table>
<thead>
<tr>
<th>Day 0</th>
<th>Day 7</th>
<th>Day 14</th>
<th>Day 21</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1000</td>
<td>2000</td>
<td>3000</td>
</tr>
</tbody>
</table>

$\text{p} = 0.709$

$\text{p} = 0.709$

$\text{p} = 0.709$

$\text{p} = 0.709$

TbTh changes in autograft vs. allograft

<table>
<thead>
<tr>
<th>Day 0</th>
<th>Day 7</th>
<th>Day 14</th>
<th>Day 21</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>100</td>
<td>250</td>
<td>0</td>
</tr>
</tbody>
</table>

$\text{p} = 0.029$

$\text{p} = 0.051$

$\text{p} = 0.011$

$\text{p} = 0.003$

DA changes in autograft vs. allograft

<table>
<thead>
<tr>
<th>Day 0</th>
<th>Day 7</th>
<th>Day 14</th>
<th>Day 21</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.5</td>
<td>1.0</td>
<td>1.5</td>
</tr>
</tbody>
</table>

$\text{p} = 0.149$

$\text{p} = 0.811$

$\text{p} = 0.290$

$\text{p} = 0.848$

Figure 5: Microarchitecture properties of defect treated with autograft vs. allograft in osteoporotic bone. BV/TV: bone volume/tissue volume; CD: connectivity density; TbTh: trabecular thickness; DA: anisotropy. $\text{p} < 0.05$ is considered significant.
showed no signi- 

bone. The results from the micro-CT or histomorphometry 

graft or allograft, whether it is used in normal or osteoporotic 

and osteoporotic bones. The hypothesis was that there 

a tibial defect rat model for the best defect repair in normal 

This study compared the e-

Figure 6: Histomorphometric evaluation of bone formation in 

Figure 6: Histomorphometric evaluation of bone formation in 

normal and osteoporotic bones: **p < 0.001.

4. Discussion

This study compared the effects of autograft and allograft in 

a tibial defect rat model for the best defect repair in normal 

and osteoporotic bones. The hypothesis was that there 

would be no difference in bone formation using either auto-

graft or allograft, whether it is used in normal or osteoporotic 

bone. The results from the micro-CT or histomorphometry 

showed no significant difference in the use of autograft and 

allograft in a tibial bone defect after an observation of 

21 days, in neither normal nor osteoporotic bone structures. 

However, micro-CT scans showed a decreased amount of 

new bone in the allograft group at 0, 7, and 14 days.

When studies compare their inventions, there should be 

some disagreement whether autograft or allograft should serve as 

the golden standard. The difference is primarily between the 

academic and clinical opinions and which kind of defect 

is used in the study [20, 21]. By making a comparison within 

the same model, a valuable information can be obtained for 

the academic purpose and for clinical application, especially, 

due to the increasing focus on substitute materials to over-

come the challenges in using graft materials and their limita-

ions [22].

When evaluating the results from this study, it should be 

noted that in both normal and osteoporotic bones, the 

BV/TV on the micro-CT scan is signi-

ficantly lower in the 

allograft group with 65.8% increase vs. only 16.4% 

increase in the autograft group. At day 21, there is no differ-

cence between autograft and allograft quantified by histomor-

phometry or microarchitectural analysis.

Histomorphometry revealed on day 21 a decreased gen-

eral bone formation when trying to use graft materials in 

osteoporotic bone (Figure 6). Again, it should be noted that 

the allograft is from a healthy donor, and yet it provides sig-

ificantly lower regeneration than it does in normal bone. 

This perfectly emphasizes the need to test graft materials or 

medical devices in the osteoporotic bone to be able to evalu-

ate their full efficacies. However, this test of graft material in 

both normal and osteoporotic bone tissues seems to be lack-

ing in regenerative potential.

Induction of osteoporosis in a rat model has previously 

been verified, with an illustration of the importance of obser-

vation until induction. Kinney et al. [23] investigated the 

changes in the trabecular bone due to OVX and found that 

the OVX procedure leads to an immediate and continuous 

decrease in trabecular bone, and after 50 days, the OVX rats 

had lost 50% of their bone volume with no rebound effect. 

Campbell et al. [24] established a detailed longitudinal time 

course of bone loss in OVX rat model in twelve weeks. The 

results of this study indicate that microarchitectural changes 

occur within the first 12 weeks after OVX in the rat model. 

Hereby, with the observation of 21 days, there are no expec-

tations of a rebound effect.

The current knowledge in the use of allograft and auto-

graft is that studies verify different outcomes dependent on 

location. In the cranioplasties, allograft has been shown to 

be superior [25]; in anterior crucial ligament (ACL), auto-

graft showed better effect on bone formation [26], whereas 

in the posterior crucial ligament (PCL), the results are equal 

between the 2 grafts [27, 28]. Hence, when comparing results 

for clinical implementation, using the correct graft is essen-

tial for the correct comparisons. Optimal results will be 

obtained by using both grafts for both positive and negative 

controls. This will make the clinical impact higher and will 

provide overall convincing results.

When facing a clinical implementation, it is furthermore 

required to focus on an economic and patient-related out-

come. The cost of using autograft is reported to be lower in, 

for example, ACL surgeries [26], but the possible side effect 

in harvesting autograft is associated with rather severe side 

effects [3]. This gives the dilemma for choosing methodology 

when harvesting autograft, but it even further requests the 

need for another substitute to replace both graft materials. 

Current tissue engineering and biomaterials with stem cells 

might provide new hope to bone regeneration.

The strengths of this study are the standardized tibia 

defect model in rats and the well-verified induction of osteo-

porosis. Yet, nothing in this paper relies on previous results, 

and the osteoporosis induction is verified by the same scan-

ner that evaluates the results in the intervention groups. This 

means that the reliability of the results of this study increases. 

Furthermore, results and same methods are tested in differ-
ent types of the bone structure allowing for the best compar-

ison of graft material, especially as the allograft in normal and 

osteoporotic bone is from the same donor.
Limitations are the lack of an empty defect that could demonstrate the baseline effectiveness of the bone regenerative potential without any bone grafts in both normal and osteoporotic bone. However, the focus of this study is to compare potential efficacies between the graft materials and not according to a specific baseline for the design model. Another limitation is the significant difference on day 0 between the groups. But when the evaluation is limited to 21 days and the allograft manages to provide the same results in such a short duration, and hence, the results are validated in this model.

5. Conclusion

This study concludes that autograft and allograft have similar bone-forming capabilities with an observation of 21 days in a rat tibial defect in a rat model, suggesting allograft could be a good alternative to autograft. Furthermore, OVX surgery for lasting osteoporosis induction in the rat model is a feasible method.

Data Availability

The data for this study were analyzed by histomorphometry and all data are stored at the Orthopaedic Research Laboratory, Department of Orthopaedics & Traumatology, Odense University Hospital, Institute of Clinical Research, University of Southern Denmark in datatables from Visio-Pharm, Denmark verifying every count and statistics made for the analyses incorporated in this study. Micro-CT scans are large files stored at TB tapes. All data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

Authors declare no conflict of interest.

Acknowledgments

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References


