Functional Outcome of Human Adipose Stem Cell Injections in Rat Anal Sphincter Acute Injury Model

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ABSTRACT

Anal incontinence is a devastating condition that significantly reduces the quality of life. Our aim was to evaluate the effect of human adipose stem cell (hASC) injections in a rat model for anal sphincter injury, which is the main cause of anal incontinence in humans. Furthermore, we tested if the efficacy of hASCs could be improved by combining them with polyacrylamide hydrogel carrier, Bulkamid. Human ASCs derived from a female donor were culture expanded in DMEM/F12 supplemented with human platelet lysate. Female virgin Sprague-Dawley rats were randomized into four groups (n = 14–15/group): hASCs in saline or Bulkamid (3 × 10⁷/60 μl) and saline or Bulkamid without cells. Anorectal manometry (ARM) was performed before anal sphincter injury, at two (n = 58) and at four weeks after (n = 33). Additionally, the anal sphincter tissue was examined by micro-computed tomography (μCT) and the histological parameters were compared between the groups. The median resting and peak pressure during spontaneous contraction measured by ARM were significantly higher in hASC treatment groups compared with the control groups without hASCs. There was no statistical difference in functional results between the hASC-carrier groups (saline vs. Bulkamid). No difference was detected in the sphincter muscle continuation between the groups in the histology and μCT analysis. More inflammation was discovered in the group receiving saline with hASC. The hASC injection therapy with both saline and Bulkamid is a promising nonsurgical treatment for acute anal sphincter injury. Traditional histology combined with the 3D μCT image data lends greater confidence in assessing muscle healing and continence.

SIGNIFICANCE STATEMENT

The increasing awareness of good quality of life sets demands for better and less invasive treatment methods of the pelvic floor disorders, for example, anal incontinence. Human adipose stem cells (hASCs) are readily available, easily obtained, have low immunogenicity and high multilineage differentiation and are, therefore, an ideal cell source. In this study, an animal model was used to develop a mini-invasive injection treatment method using hASCs. The functional measurements showed significant improvement in hASC-treatment groups compared with the controls. A biocompatible carrier polyacrylamide hydrogel Bulkamid was found to be a suitable carrier for the stem cells, and a novel method of micro-computed tomography was found useful for targeting the histological slides.

INTRODUCTION

Anal incontinence (AI) is a devastating condition that significantly reduces the quality of life. Especially in women, the fecal incontinence symptoms cause depression, embarrassment and lifestyle changes that have a negative effect their everyday life [1]. The primary management of AI is conservative treatment including dietary, medical, and psychological interventions, as well as physiotherapy. A common operation for persistent AI has been secondary sphincteroplasty which may not have the desired long-term results, especially with denervated sphincters [2]. Sacral nerve modulation and transcutaneous posterior tibial neuromodulation have increasingly been used for fecal incontinence. However, the cure rates vary depending on the outcome measure in use with long-term results reaching up to 54% for sacral neuromodulation [3, 4]. Surgical methods often carry a risk of complications and are both demanding and expensive. AI is defined as an involuntary loss of flatus, liquid, or solid stool that is a social or hygienic
A study to use Bulkamid as a stem cell carrier, to study in vivo the hASCs in an animal functional model and to use micro-computed tomography (μCT) as a tool to view the treated anal sphincter area.

**MATERIALS AND METHODS**

The animal study protocol was approved by the Regional State Administrative Agency (AVI/Ella no ESAVI/2828/04.10.07/2015). Adipose tissue sample was obtained under the approval of the Ethics Committee of the Pirkanmaa Hospital District (Tampere, Finland, R15161). The hASCs were isolated from adipose tissue sample from a female donor undergoing elective plastic surgery at Tampere University Hospital (Tampere, Finland) with the patient’s written consent. The final study was designed after a pilot study of 24 Sprague-Dawley rats (results not included in the analysis). In the pilot study, the rat anorectal manometry (ARM) technique, anal sphincter cutting and repairing, the injection technique, the amount of the gel, and the amount of stem cells per injection were tested. The cell amounts of $5 \times 10^5/100 \mu l$ and $5 \times 10^6/100 \mu l$ were compared, and the lower cell count was chosen due to better cell viability before injection.

**Treatment Protocol**

Sixty (60) Sprague-Dawley female virgin rats (Janvier Laboratoires), weight 220–300 g, age 14 weeks were randomly selected, then anesthetized with intraperitoneal injections of medetomidine 0.25 mg/kg and ketamine 32.5 mg/kg, and buprenorfin 0.05 mg/kg s.c. and carprofen 0.05 mg/kg s.c. were used for postoperative pain. After the anal manometry and the sphincter operation, the anesthesia was reversed using atipamezole 1 mg/kg.

The ARM was performed using Polygraf ID manometry system with ERCP manometry triple lumen catheters (Medtronic, Polygram NET, computer unit Windows XP, Minneapolis, MN, USA). The measurements were conducted preoperatively and at 2 and 4 weeks after the operation. (Fig. 1, Supporting Information Fig. S1).
The duration of ARM procedure was approximately 30 minutes, during which resting anal sphincter pressures and peak pressures during spontaneous contraction were measured (resting pressure 3–7 times [mean 5.64] and contraction pressure 7–12 times [mean 10.4]). Mean values of these measurements were used in further analysis.

The sphincter was cut from all animals to mimic an acute fourth grade anal sphincter tear (injury of the external and internal sphincter muscle and anal mucosa) and sewed back with 6-0 poliglecaprone (Ethicon, Johnson & Johnson, Monocryl, Somerville, NJ, USA) continuous stitches utilizing magnifying loupes. First, the anal mucosa and internal sphincter were repaired, and then, the injections were delivered after which the perineal skin was closed.

The rats were divided into four groups for the injections: injection of \( 3 \times 10^5 \) hASCs in saline (0.9% NaCl) solution (\( n = 15 \)), hASCs in 2.5% polyacrylamide hydrogel (Bulkamid, Contura International A/S, Denmark) (\( n = 15 \)). The control groups consisted of rats with only saline (\( n = 16 \), two died within the first postoperative day presumably because of anesthetic reaction and were excluded from the analysis) and polyacrylamide hydrogel injections (\( n = 14 \)). There were two injections with a 25-gauge needle on both ends of the external sphincter (degrees 30° and 330° on superimposed clock face, Supporting Information Fig. S3); the total injection volume of the four injections being 60 μl per animal for all Bulkamid and 0.9% NaCl-groups.

The rats were again anesthetized for the anal manometry control with medetomidine and ketamine combination as described above. After the last anal manometry examination, the rats were euthanized using carbon monoxide. The anal manometry results were analyzed by a blinded researcher (HT) to exclude observer bias.

### ASC Isolation, Cell Culture, Cell Viability, and Phenotype

The hASCs were isolated as previously described [31]. The hASCs were expanded in DMEM/F12 (1:1) (Thermo Fisher Scientific Inc., Carlsbad, CA, USA, https://www.thermofisher.com) supplemented with 5% Pooled Human Platelet Lysate (Stemulate; Cook General Bio-Technology, Indianapolis, IN, USA, http://www.stemtrec.com), 1% antibiotics (100 U/ml penicillin and 0.1 mg/ml streptomycin; Thermo Fisher Scientific Inc.), and 1% l-glutamine (Thermo Fisher Scientific Inc, GlutaMAX-100, Indianapolis, IN, USA). The medium was changed twice a week, and the cells were divided upon reaching confluency. The cells were detached using TrypLE Select (Thermo Fisher Scientific Inc., Carlsbad, CA, USA). From 4 to 5 days prior to cell injections, the ASCs were labeled with 200 μg/ml magnetizable nanoparticles (PMP-50; Kisker Biotech GmbH & Co., Steinfurt, Germany, https://kisker-biotech.com) for 48 hours for cell detection. For cell injections, \( 5 \times 10^5 \) hASCs (passage between 4 to 7) were blended with 100 μl of 0.9% NaCl (Baxter Healthcare SA, Zurich, Switzerland, http://Baxter.com) or Bulkamid hydrogel (Contura International A/S, Soeborg, Denmark, https://bulkamid.com).

For the cell viability, \( 5 \times 10^5 \) hASCs were blended with 100 μl Bulkamid, incubated at room temperature for 3 hours (the maximum delay between the cell preparation and the injections), and the cell viability was assessed with LIVE/DEAD Viability/Cytotoxicity kit for mammalian cells (Thermo Fisher Scientific Inc., Carlsbad, CA, USA). The hASCs in Bulkamid were stained with 1 μM Calcein AM and 0.8 μM Ethidium homodimer-1 (EthD-1) for 45 minutes.

### μCT Imaging

The analysis of the anal sphincter was performed after sacrificing the animals. The excised samples were imaged in a μCT instrument (Kraxia MicroXCT-400, Carl Zeiss, Pleasanton, CA, USA). To increase the contrast between the soft tissues, the samples were put through a staining regime. The tissue samples were stored in 4% paraformaldehyde (PFA)-solution (Sigma). The PFA was then changed to 70% ethanol solution, then to 95% ethanol, >99.5% ethanol, and 10 mg/ml iodine in ethanol. The samples were transferred into sample holders and were kept at room temperature for a minimum of 24 hours prior to imaging in order to avoid imaging artifacts resulting from thermal expansion of the tissue samples. The imaging parameters for all 58 samples were constant (Table 1). After imaging, the samples were transferred into a 70% ethanol solution for storage and following histological analyses. Tomographic three-dimensional (3D) image reconstruction was performed with proprietary software installed in the instrument (Carl Zeiss, Xradia XMReconstructor, Pleasanton, CA, USA). The reconstructed image volume was imported into AVIZO image processing software (FEI Company, Hillsboro, OR, USA) for image processing and inspection. The anal sphincter complex was visualized through 3D volume rendering. It was used in combination with tomographic data exploration as freely selectable two-dimensional (2D) views of the 3D dataset to confirm the histological assessment of muscle continuity. Additionally, it was used to visualize in 3D the entire sample instead of selected 2D sections as in histology and to monitor the presence of any detectable PMP-50 labeled cells.

### Table 1. Technical details about the micro-computed tomography imaging

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
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<td>Voltage</td>
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</tr>
<tr>
<td>Current</td>
<td>166 μA</td>
</tr>
<tr>
<td>Source distance</td>
<td>58 mm</td>
</tr>
<tr>
<td>Detector distance</td>
<td>170 mm</td>
</tr>
<tr>
<td>Exposure time</td>
<td>4 seconds</td>
</tr>
<tr>
<td>Imaging time</td>
<td>2.5 hours</td>
</tr>
<tr>
<td>Pixel size</td>
<td>17.0354 μm</td>
</tr>
<tr>
<td>Image volume dimensions</td>
<td>1000 × 1000 × 1000 pixels</td>
</tr>
</tbody>
</table>

Fluorescence pictures were taken with Fluorescence Microscope (Olympus. IX51S18F-2 and camera DP71, Japan).

The phenotype of the hASCs was assessed with fluorescence-activated cell sorter (BD Biosciences, Franklin Lakes, NJ, USA, https://www.bdbiosciences.com). BD FACSAria Fusion Cell Sorter) at passage 7. Monoclonal antibodies against CD14-phycoerythrin-cyanine (PE-Cy7), CD19-PE-Cy7, CD45RO-allophycocyanin (APC), CD54-Fluorescein isothiocyanate (FITC), CD73-phycoerythrin (PE), CD90-APC (BD Biosciences, Franklin Lakes, NJ, USA), CD105-PE (R&D Systems Inc., Minneapolis, MN, USA, http://www.rndsystems.com), CD34-APC and HLA-DR-PE (Immunotools GmbH, Friesoythe, Germany, http://www.immunotools.de) were used. According to the International Society for Cellular Therapy standard criteria, cells positive for CD73, CD90 and CD105 but negative for CD14, CD34, CD45, and HLA-DR are considered as MSCs [32, 33].
Results

In Vitro Results

Live/dead staining demonstrated that after 3 hours of incubating the hASCs in Bulkamid, the hASCs were mostly alive. However, there were also some dead cells in the mixture (Fig. 2). The hASC phenotype was assessed at passage 7 using flow cytometry. The hASCs expressed surface markers CD73 (99.9%), CD90 (99.9%), and CD105 (99.6%). Expression of CD14 (1.4%), CD45RO (0.6%), and HLA-DR (0.3%) was very low, and expression of CD19 (6.0%), CD34 (8.1%), and CD54 (8.5%) was low. This confirms the mesenchymal origin of the hASCs.

Histology

The formalin fixed, paraffin embedded tissues were sectioned (4 μm) and stained with Hematoxylin & Eosin (HE) for morphological interpretation. Furthermore, picrosirius red staining was used to demonstrate collagen and Perls Prussian blue to demonstrate the presence of iron from tissue sections. Immunohistochemistry using anti-human Vimentin 1:200 (Clone:BS13, BSH-7100, Nordic Biosite, Täby, Sweden) and STEM121 1:500 (Clone: Stem121, Y40410, Takara Bio Inc., Shiga, Japan) antibodies was performed to detect hASCs from rat tissue sections. Smooth muscle actin antibody 1:200 (Clone BS66, BSH-7459, Nordic Biosite, Täby, Sweden) and Anti-Desmin mouse monoclonal antibody (Clone BS21, Nordic Biosite, Täby, Sweden) were used to detect rat muscle tissue. Finally, CD68 antibody 1:200 (Clone: ED1, ab31630, ABCAM, Cambridge, United Kingdom) was used to detect rat macrophages and verify the origin (PMP-50 containing hASCs or naturally iron containing rat macrophages) of positive Perls Prussian blue staining. All antibodies were produced in mouse and primary antibodies were detected using anti-mouse horseradish peroxidase polymer. Rat and human paraffin control multi-tissue sections were used as a negative and positive tissue controls for immunohistochemical analyses.

Olympus BX-60 microscope (BSH 747) and an integrated color digital camera (Scion) were used for the evaluation of the slides. The slides were also imaged and digitalized with a 3D Histec Panoramic MIDI instrument. The scoring of the inflammation was based on the inflammatory cell infiltration into the lesion site, edema, hemorrhage and necrosis of tissue [34]. The number of inflammatory cells was evaluated manually from the low power field (∼200) image frame using a “hot spot” selection. The inflammation was scored with following grades: 0 = no histological features of inflammation; 1 = diffuse inflammatory cell infiltration, <100 cells; 2 = diffuse inflammatory cell infiltration, 100–500 cells, mild edema and hemorrhage; 3 = inflammatory cell infiltration, >500 cells, edema and hemorrhage; 4 = inflammatory cell infiltration, >1,000 cells, edema, hemorrhage and necrosis. Fibrosis was evaluated separately with following grades: 0 = no fibrosis; 1 = mild fibrosis/collagen formation; 2 = strong fibrosis/collagen formation. The infiltration of the cells into the hydrogel was evaluated with samples containing Bulkamid.

Statistical Analysis

The statistical analysis was performed by using the IBM SPSS version 22 (IBM, Chicago, IL, USA). The four treatment groups were compared at baseline, at 2, and at 4 weeks. Statistical significance was tested by using chi-square or Fisher’s exact test with categorical and one-way analysis of variance (ANOVA) with continuous variables. The groups and the time points within the groups were compared using ANOVA for repeated measures.

In Vivo Results

The baseline rat characteristics (weight, ARM measurements) of each group are presented in Table 2. The rats in the 0.9% NaCl and Bulkamid control groups were slightly, though statistically significantly, heavier than the hASC treatment groups’ rats. Otherwise, there was no difference in baseline characteristics between the groups.

Anorectal Manometry Results

First, the four groups were compared based on the ARM results before injury and at 2 and 4 weeks. The measured variables were the median resting pressure and the peak pressure during spontaneous contraction of the anal sphincter complex. The median resting and the peak contraction pressures were higher in the hASC treatment groups at 2 and at 4 weeks (Table 2). Further analysis showed that the trend of the contraction pressure was significantly higher in the both hASC-groups compared with the saline and Bulkamid control groups (Fig. 3). The difference between the groups remained statistically significant when adjusted for baseline measurement.

Histology

In the histological analysis, no hASCs were recognized in the preparations neither at 2 nor 4 weeks according the Vimentin, STEM121, or Perls Prussian blue staining. This was confirmed with CD68 staining of rat macrophages. Rat endogenous macrophages containing iron stained positively with Perls Prussian blue and rat specific CD68 immunoperoxidase reaction located into the same cells (Fig. 4). Vimentin and STEM121 were positive in a cytoblock section prepared from the same cells that were injected into the rats (data not shown).

There was no statistical difference in sphincter muscle continuity, fibrosis, or collagen formation between the four groups. The Bulkamid-hydrogel was well integrated in the tissue with minor foreign body reaction according to the HE staining. There was more inflammation in the hASC-groups, especially in the 0.9% NaCl + hASC-group (Table 2; Fig. 4, Supporting Information Fig. S2).

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The mCT image datasets were used to confirm independently the continuity of the sphincter muscle shown in the histology. By viewing the image data in multiple orthogonal views, greater confidence could be attested to the histological assessment. (Fig. 5, Supporting Information video). There was total agreement between histology and CT interpretation in 76% of the samples. There was minor disagreement in 11 samples and serious disagreement in muscle continuity in 3 samples (5%). This did not affect the statistical difference between the groups. Thus, the ability to conduct nondestructive histomorphometric analysis on samples provides valuable image data that can be used to perform robust 3D analyses when necessary. In some samples with Bulkmid, small regions with high x-ray attenuation regions were observed that could indicate cells or remains of PMP-50 particles (data not shown). However, it was not possible to confirm whether these regions indicate the presence of PMP-50 particles or whether they were the result of local aggregation of iodine. As iron specific staining of the histological samples failed to detect PMP-50 in any of the samples, the presence of PMP-50 particles could not be confirmed.

**DISCUSSION**

Our aim is to develop an effective, mini-invasive treatment for AI, which is a highly distressing condition and all the more lacking an efficient treatment method. Studies about the existing treatment methods are heterogeneous and long-term results are mostly missing. Tissue engineering and cell therapy have been considered to be compelling alternatives and hold a great deal of promise and excitement. The advantage of using ASCs compared with the other stem cell sources is that the tissue can easily be harvested and it is readily available in large quantities. ASCs can be easily expanded in vitro and have an extensive self-renewal capacity [35, 36].

In our in vivo-study, we found significant improvement in anal sphincter resting and contraction pressures in hASC treatment groups after acute sphincter injury compared with the control groups. Previous functional anal sphincter assessments have mostly been performed in vitro using anal sphincter muscle samples from euthanized animals to measure the contractility after electrical stimulation [10, 12, 17, 37]. Salcedo et al. measured in

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**Table 2. The rat baseline characteristics, functional results from anorectal manometry and histology**

<table>
<thead>
<tr>
<th></th>
<th>0.9%NaCl + hASC</th>
<th>Bulkmid + hASC</th>
<th>0.9%NaCl</th>
<th>Bulkmid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight (g)</td>
<td>Mean/median</td>
<td>SD/Q1–Q3</td>
<td>Mean/median</td>
<td>SD/Q1–Q3</td>
</tr>
<tr>
<td></td>
<td>268.0/270.0</td>
<td>12.1/260.0–280.0</td>
<td>263.6/270.0</td>
<td>18.6/250.0–280.0</td>
</tr>
<tr>
<td>PreopARM (n)</td>
<td>15</td>
<td>14</td>
<td>14</td>
<td>15</td>
</tr>
<tr>
<td>Rest med</td>
<td>9.1</td>
<td>3.4</td>
<td>8.4</td>
<td>2.8</td>
</tr>
<tr>
<td>Peak contr</td>
<td>97.5</td>
<td>25.2</td>
<td>88.5</td>
<td>29.6</td>
</tr>
<tr>
<td>ARM 2 wk (n)</td>
<td>15</td>
<td>14</td>
<td>14</td>
<td>15</td>
</tr>
<tr>
<td>Rest med</td>
<td>9.3</td>
<td>2.6</td>
<td>9.6</td>
<td>2.1</td>
</tr>
<tr>
<td>Peak contr</td>
<td>74.6</td>
<td>16.0</td>
<td>74.0</td>
<td>13.6</td>
</tr>
<tr>
<td>ARM 4 wk (n)</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>Rest med</td>
<td>10.1</td>
<td>3.0</td>
<td>9.2</td>
<td>2.3</td>
</tr>
<tr>
<td>Peak contr</td>
<td>79.5</td>
<td>16.4</td>
<td>76.2</td>
<td>21.3</td>
</tr>
</tbody>
</table>

| Inflammation        |                 |                |          |        |
| gr 0 (%)            | 0.0             | 21.4           | 28.6     | 33.3   |
| gr 1 (%)            | 20.0            | 28.6           | 50.0     | 46.7   |
| gr 2 (%)            | 40.0            | 42.9           | 21.4     | 20.0   |
| gr 3 (%)            | 13.3            | 7.1            | 0.0      | 0.0    |
| gr 4 (%)            | 26.7            | 0.0            | 0.0      | 0.0    |

Abbreviations: ARM 2 wk, anorectal manometry at 2-week time point; ARM 4 wk, anorectal manometry at 4-week time point; hASC, human adipose stem cells; NaCl, sodium chloride; Peak contr, peak pressure during spontaneous contraction; Preop ARM, preoperative anorectal manometry; Q1–Q3, 25 and 75 percentiles; Rest med, median resting anal pressure; Weight, preoperative weight.

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**Figure 3.** The trends of the four groups showed a significantly higher contraction pressures in both hASC treatment groups. Abbreviations: hASCs, human adipose stem cells; NaCl, sodium chloride.
vivo function of the anal sphincter after a sphincterotomy or pudendal nerve crush followed by either rat bone-marrow derived MSC injection or saline [18]. In their further investigations, the same research group found that both serial i.v. infusion and i.m. injections of MSCs after partial sphincter excision resulted in increased anal pressures [19]. Pathi et al., on the other hand, did not find significant advantage in the iv-administration of rat bone-marrow derived MSCs compared with the PBS controls, but the local injections of MSCs were effective according to in vitro assessment [17]. Recently Fitzwater et al. noticed that the administration of rat myogenic stem cells enhanced the contractile function of the sphincter without significant changes in histologic morphology, which addresses the paracrine processes in stem cell therapy [14]. To simplify the cell isolation and proliferation process, Mazanti et al. used freshly isolated minimally manipulated bone-marrow derived mononuclear cells without expansion and found them to be as effective as in vitro expanded BM-MSCs in the recovery of iatrogenic anal sphincter rupture [38]. Muscle-derived stem cell injections have been used in small pilot studies in women and the results are promising [39–41]. Frudinger et al. treated 10 patients with autologous myogenic stem cells and found a significant improvement in the AI symptoms at 1- and 5-year controls. However, there was no significant change in anal manometry measurements [39, 40]. In a small pilot study, where hASCs therapy was used in combination with surgical sphincteroplasty in treatment of AI, there was no difference in AI symptoms.

Figure 4. The histology stainings and examples of the inflammatory grading. (A): Picosirius red-staining, 0.9% sodium chloride (NaCl) at 2 weeks; (B): Anti-Desmin, Bulkamid at 4 weeks; (C): Immunohistochemistry staining CD68, Bulkamid+hASC at 2 weeks; (D): HE-staining, Bulkamid+hASC at 2 weeks, inflammation grade 1; (E): HE-staining, Bulkamid+hASC at 4 weeks, inflammation grade 2; (F): HE-staining, 0.9%NaCl+hASC at 2 weeks, inflammation grade 4; (G): Perls Prussian blue-staining for iron particles, Bulkamid+hASC at 2 weeks; (H): Combination of CD68 and Perls Prussian blue of the sample G showing that the iron particles localize at the rat endogenous macrophages. Arrow = Bulkamid+hASC-injection. Scale bar Figures (A–F) 500 μm, Figures (G–H) 20 μm.
between the hASC and saline injection group. However, the number of patients was limited and heterogeneous and the follow-up time short. Despite these limitations, they discovered an increase in total muscle area in an endorectal sonography assessment in the hASC treatment group compared with the saline control group [20].

In our study, the anal pressures were comparable with the other animal studies although there are slight variations—possibly due to different techniques (balloon vs. double-triple-lumen catheter, electrical stimulation vs. spontaneous contraction) and different size of the anal canal in different species [13, 16, 19]. Salcedo et al. stated that rat anal pressures tend to return to baseline after sphincterotomy at 4 weeks even without intervention, but this does not occur after pudendal nerve transaction [42]. In our results, both resting and contraction anal pressures were significantly lower in both control groups at 2 and 4 weeks compared with the hASC treatment groups. There was also a rising pressure trend within the stem cell groups that might have resulted in even higher pressures and better recovery in longer follow-up.

We used ASCs from a single donor mainly because of the safety regulations. It is known that there is some heterogeneity between different donors; for example, age, sex, body mass index, and site of harvest are known to have an effect on the properties of the ASCs [43–47]. Xenotransplantation has been used before to study development, physiology, and pathophysiology of human tissues in animal models, for example, enteric nervous system [48] and rodent models have been tested in treatment of stroke [49]. However, there are differences in study development, physiology, and pathophysiology of human and animal physiology that have to be kept in mind when interpreting the results [51].

Different labeling systems of injected cells have been used in previous studies. Kang et al. found fluorescent dye PKH-26 labeled rat myogenic stem cells in fluorescent microscopy of rat anal sphincters at one week after the injection [10]. PKH-26 label in animal autologous myogenic stem cells were also used by Kajbafzadeh (rabbit) and Oh (dog): both groups found labeled cells at 2 months and 3 months control, respectively [15, 16]. Aghaeasghari et al. used bromodeoxyuridine labeled human umbilical cord stem cells as well as rabbit bone marrow stem cells (BMSCs) in a rabbit model, the labeled cells were found in immunohistochemistry at 2 weeks control [52]. On the other hand, Cruz et al. demonstrated that the green fluorescent protein (GFP) labeled BMSCs disappear in 10 days after i.v.-administration [53]. Salcedo et al. found no visible GFP-labels in the anal sphincter after administration of i.m. or i.v. MSCs derived from rat bone marrow [18]. We attempted to find the PMP-50 labeled hASCs in the tissue samples using μCT, histology and immunohistochemistry. Although some potential signs of PMP-50 were detected in μCT, we were not able to confirm the PMP-50 labeled cells either in the μCT scan or in the histology analysis. Histological sections stained positive with Perls Prussian blue but rat macrophage specific CD68 antibody stained the same cells indicating that the signal comes from hASCs ingested by rat macrophages or just iron that macrophages contain naturally. In our pilot study, there were tracks of the injected cells in hydrogel with the higher cell amount (5 × 10⁶ cells/100 μl) and with bigger injection volume 100 μl (Supporting Information Fig. S2). In the actual study, due to better cell viability and injection experience, we decided to use lower cell count and lower volume and were not able to verify the presence of the stem cells. The evanescence of the cells may be due to the rat immune defense destroying the human cells. This is supported by the lack of human specific STEM121 and Vimentin staining of the tissue sections.

There are also the safety considerations in cell therapy. Jacobs et al. found no evidence of myogenic stem cell migration to the liver or lung. However, they detected local ectopic foci of growth in two treated rat anal sphincters after 30 days; the small tumors were benign tumors with no nuclear abnormalities [54]. In our study, normal histology was confirmed both by 3D μCT scanning of the whole sphincter sample area, as well as by selected histological sections. The μCT is a valuable approach to visualize the entire treated area and the injection sites in 3D to target the histological slides. The ability to increase imaging resolution without proportionally decreasing sample size opens possibilities for performing analyses without destructive sample preparation. The μCT protocol applied in this study showcased this methodology by complementing the traditional histological analyses with increased amount of available data, which resulted in greater confidence in the analysis. Additionally, in combination with appropriate concentrations of labeling agents, it may also be feasible to monitor the distribution of introduced cells.

This animal model, like any other animal model, is not directly comparable with birth injury in humans. The trauma in the rat model was caused by a direct clean-cut wound whereas in
obstetric trauma the sphincter is torn, and the tissue has suffered from prolonged hypoxia and denervation. In an obstetric trauma, there is also presence of tissue edema and blood, which makes suturing the rupture more challenging in real life. In our animal model, the rupture was easy to identify with no delay, and the operative technique was consistent with only one operator. The small scale of the rat anatomy made it difficult to distinguish the inner and outer sphincter from each other and that is why the injections were aimed at the external sphincter rather than submucosa or between the sphincters. In humans, the intersphincteric injection might be better. However, our model applies to acute injury and is perhaps not applicable for chronic conditions with fibrous anal sphincters often seen in AI patients. Fibrosis is a histological finding often seen in scar tissue and in our model the follow up time was too short to analyze the possible impact on scar tissue formation. There is evidence that cell therapy might be able to prevent vocal fold scaring in laryngeal microsurgery and fibrous keloid formation after skin incision in acute trauma model [55, 56].

The possible mechanisms of AI stem cell therapy are direct stem cell integration and muscle regeneration, the bulking effect of the cells and the carrier, the trophic effect caused by the stem cell growth factor secretion or immune modulation, and the decreased inflammation that improves tissue healing [33, 57]. The bulking effect is probably not the mechanism according to our results, because there was no advantage in the bulking agent Bulkamid compared with the 0.9% NaCl-controls, and also because there were good results with hASCs in 0.9% NaCl as well. For human AI, the bulking effect in the injection therapy is perhaps not even desirable [58]. According to our results, lack of differences in muscle continuity between the groups refers to paracrine effect rather than to direct stem cell differentiation into muscle cells. The mechanism may be through cell fusion, endogenous activation of satellite cells or some inflammatory process [59]. From the inflammatory perspective, Bulkamid seems to have an advantage over 0.9% NaCl as a carrier, although it is not clear what stage of inflammation is desirable for the healing process. Bisson et al. showed that certain amount of inflammation and acute injury is required in cell therapy: injection of myoblasts on the opposite site of cryoinjured anal sphincter did not restore the anal sphincter function, whereas the injection into the lesion itself did [60]. On the other hand, Salcedo et al. showed that systemic injection also had a positive effect; however, this was not seen in the study of Pathi et al. [17, 19]. One advantage of hydrogel over 0.9% NaCl is that the cell sedimentation and aggregation are mostly avoided with Bulkamid. Bovine collagen gel (Contigen) has also been used in clinical trials experimenting stem cell injection therapy for female urinary incontinence [61, 62].

It has been suggested that stem cell transplantation may facilitate endogenous repair even in patients with advanced age and comorbidities who have compromised repair function [13]. Therefore, stem cell therapy would be especially useful in patients with compromised anal sphincter function. Bohl et al. have successfully implanted engineered biosphincters in rabbits with iatrogenic AI to restore anal sphincter function [63]. Sphincter transplantation is however a major surgical operation. Stem cell injection therapy can be performed under local anesthesia in an outpatient setting with no major complications. According to our preliminary results, the ASC treatment is effective in treatment of an acute anal sphincter injury. In the future, animal models for chronic injury are needed to develop an effective and clinically relevant treatment method.

**CONCLUSION**

ASCs combined with either saline or synthetic nondegradable hydrogel is a novel and attractive, mini-invasive treatment method for acute anal sphincter rupture and AI. Due to low inflammatory response and good tissue integration, Bulkamid hydrogel appears to be a suitable injection agent and carrier for the ASCs. The technique of anal sphincter injection therapy is simple in an animal setting and the use of human ASCs in rat AI model is feasible. The 3D μCT is a valuable addition to the traditional histology in analyzing soft tissue samples. Further studies with suitable chronic injury animal models with longer follow-up periods are needed to confirm the functional restoration and to develop a truly effective treatment method for AI patients.

**ACKNOWLEDGMENTS**

We thank Tampere University Vetlab animal attendants for their assistance, Sari Kalliokoski from BioMediTech for the preparation of the anal sphincter samples, and Dr. Jukka Kallio for lending the magnifying loupes. The histological samples were prepared and evaluated by BioSiteHisto/Teppo Haapaniemi as an outsourced service. This study was financially supported by the Competitive State Research Financing of the Expert Responsibility area of Tampere University Hospital and TEKES (the Finnish Funding Agency for Innovation) Human Spare Parts project. The expense of the animal acquisition and maintenance, including drugs used in the operation; the study equipment (anal manometry); the stem cell GMP-laboratory expense and the outsourced histology service was covered by a competitive research grant for associate professor Susanna Miettinen from Pirkkanmaa Hospital District. Kirsi Kuismanen has received a 1-month research grant for the study from Tampere University hospital/Pirkkanmaa Hospital District.

**AUTHOR CONTRIBUTIONS**

K.K.: conception and design, administrative support, provision of study material, collection and assembly of data, data analysis and interpretation, manuscript writing, final approval of manuscript; M.J. and N.N.G.: conception and design, provision of study material, collection and assembly of data, data analysis and interpretation, manuscript writing, final approval of manuscript; H.T.: conception and design, data analysis and interpretation, final approval of manuscript; H. H.: conception and design, data analysis and interpretation, manuscript writing, final approval of manuscript; K.N. and J.H.: conception and design, administrative support, data analysis and interpretation, manuscript writing, final approval of manuscript; S.M.: conception and design, financial support, administrative support, data analysis and interpretation, manuscript writing, final approval of manuscript.

**DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST**

The authors indicated no potential conflicts of interest.
REFERENCES


