Decellularization and Characterization of Porcine Superflexor Tendon: A Potential Anterior Cruciate Ligament Replacement

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The porcine superflexor tendon (SFT) was identified as having appropriate structure and properties for development of a decellularized device for use in anterior cruciate ligament reconstruction. SFTs were decellularized using a combination of freeze–thaw and washes in hypotonic buffer and 0.1% (w/v) sodium dodecyl sulfate in hypotonic buffer plus protease inhibitors, followed by nuclease treatment and sterilization using peracetic acid. The decellularized biological scaffold was devoid of cells and cell remnants and contained only 13 ng/mg (dry weight) residual total DNA. Immunohistochemistry showed retention of collagen type I and III and tenascin-C. Quantitative analysis of sulfated sugar and hydroxyproline content revealed a loss of glycosaminoglycans compared with native tissue, but no loss of collagen. The decellularized SFT was biocompatible in vitro and in vivo following implantation in a mouse subcutaneous model for 12 weeks. Uniaxial tensile testing to failure indicated that the gross material properties of decellularized SFTs were not significantly different to native tissue. Decellularized SFTs had an ultimate tensile strength of 61.8–10.3 MPa (±95% confidence limits), a failure strain of 0.29–0.04, and a Young’s modulus of the collagen phase of 294.1–61.9 MPa. Analysis of the presence of the α-Gal (galactose-α-1,3-galactose) epitope by immunohistochemistry, lectin binding, and antibody absorption assay indicated that the epitope was reduced, but still present post decellularization. This is discussed in light of the potential role of noncellular α-Gal in the acceleration of wound healing and tissue regeneration in the presence of antibodies to α-Gal.

Keywords: decellularization, tendon, anterior cruciate ligament

Introduction

The anterior cruciate ligament (ACL) is the main intra-articular ligament in the knee joint1 and the most commonly injured, with a reported 50,000–175,000 reconstructions taking place each year in the United States.2 Rupture or tearing of the ACL leads to instability of the knee and reduced function, which (if not treated effectively) can lead to degenerative joint disease.3 The ACL has limited vascular supply and capacity to heal and therefore if it is ruptured or torn, the only viable treatment is surgical reconstruction. The current gold standard for ACL reconstruction is autologous bone–patellar tendon–bone; however, the use of semitendinosus or gracilis tendon has become more popular in recent years.4,5 Autografts offer good results and the advantage of no disease transmission or adverse immune response6,7; however, limitations include donor site morbidity, increased operative time, and the potential for the harvested tissue to prove unsuitable. The major alternative is to use tendon allograft with advantages of shorter less complicated surgeries, no donor site morbidity, and the availability of larger grafts.8,9 The major disadvantage of ACL reconstruction using allograft is that the donor tissue may suffer an adverse immune response. Reconstruction of the ACL using allograft tissue has also been shown to have a higher failure rate in younger more active patients.5

Following ACL reconstruction surgery with autografts or allografts, the endogenous cells are unlikely to survive,
leading to initial graft deterioration. During the remodeling process, there is a marked decrease in biomechanical strength of tendon autografts, which can be attributed to initial cell necrosis within the tissue. Following implantation, the rate of tissue degradation is greater than host cell repopulation. Indeed, it may well be better to use a non-vital graft with a stable matrix, such as an acellular natural tendon scaffold.

Biological scaffolds produced by decellularization of mammalian tissues are used extensively in surgical applications. Mostly derived from porcine small intestinal sub-mucosa, human or animal dermis, or pericardium, their use is typically limited by inadequate biomechanical properties. The innovation in our approach has been to develop processes for decellularization of the tissue type to be replaced, preserving the physical properties, thus delivering the natural multiscale hierarchical matrix architecture for appropriate tissue-specific biomechanical and biological functions. We have produced acellular biological scaffolds for use in cardiovascular, musculoskeletal and wound care applications. In this study, we describe the decellularization and characterization of porcine superflexor tendon (SFT), which was identified as having appropriate tissue-specific biomechanical and biological properties for potential use in ACL reconstruction.

Materials and Methods

Tissue procurement

The hind legs of female, 70 kg, 4-month-old, large White pigs were obtained from an abattoir (J. Penny, Leeds) within 24 h of slaughter. The SFT is located in the porcine foot running from the toe to the ankle. The SFT was identified after removal of the skin and subcutaneous tissue and cleaned of connective tissue before being aseptically dissected. The porcine SFT divides into two bands approximately halfway along its length and one band was carefully removed, taking care not to damage the remaining tissue. The SFTs were stored at −80°C on filter paper moistened with phosphate-buffered saline (PBS; Oxoid) until required.

Decellularization of porcine SFTs

SFTs were decellularized in batches of 12. SFTs were subjected to three freeze–thaw cycles at −80°C. The first freeze–thaw was conducted on dry tissue. SFTs were then immersed in hypotonic buffer plus proteinase inhibitors (10 mM Tris [Sigma Aldrich]; 2.7 mM disodium ethylene-diamine-tetra acetic acid [EDTA; Thermo Fisher Scientific]; 10 KIU/mL aprotinin [Nordic Pharma], pH 8.0), and frozen and thawed at room temperature (RT, 22–25°C) twice. SFTs were washed in acetone (WVR International) three times for 1 h (RT) to remove fat, then washed five times in PBS plus aprotinin (10 KIU/mL) to remove the aceton at RT. All subsequent steps in the process were conducted on tendons in individual pots containing 100 mL of solution at 37°C with agitation, unless otherwise stated. Tendons were disinfected in a solution of 0.5 mg/mL gentamicin sulfate (VWR International), 0.05 mg/mL vancomycin hydrochloride (Sigma), and 0.2 mg/mL polymyxin B (Sigma) in PBS for 1 h. Tendons were washed in hypotonic buffer for 24 h and then hypotonic buffer plus 0.1% (w/v) sodium dodecyl sulfate (SDS; Sigma) for 24 h. The washes in hypotonic buffer and hypotonic buffer plus SDS were repeated twice. SFTs were washed in PBS plus aprotinin three times for 30 min, 70 h, and 30 min before treatment with nuclease (1 U/mL benzonase [Merck] in 60 mL 50 mM Tris plus 1 mM MgCl2 [Thermo Fisher Scientific] at pH 7.6) for 2 h, three times. SFTs were washed three times in PBS plus 2.7 mM EDTA before an 18-h wash in hypertonic buffer (50 mM Tris; 1.5 M NaCl [Sigma]; pH 7.6). After three 30-min PBS washes, SFTs were sterilized using peracetic acid (0.1% v/v, pH 6; Sigma) for 3 h. Finally the SFTs were washed in PBS three times for 30 min, twice for 60 h, and once for 120 h and stored refrigerated in PBS.

Histology and immunohistochemistry

Six decellularized SFTs were analyzed and compared with native tissue (n = 6). Transverse and longitudinal samples were taken from the toe insertion point (toe region), middle, and ankle insertion point (ankle region) of each tendon and fixed in 10% (v/v) neutral buffered formalin (NBF; Gentamedical) for histochemical staining and zinc fixative (0.1 M tris; 3.2 mM calcium acetate [Thermo Fisher Scientific], 27 mM zinc acetate [Sigma Aldrich], and 37 mM zinc chloride [Fluka] pH 7.2) for 4 days for immunohistochemical analysis. The samples were processed automatically (Leica 11020 Tissue processor) and embedded in paraffin wax using standard techniques. For histochemical staining, sections (6 and 10 μm) were taken from the surface through to the central region of each block and stained with hematoxylin (Biostain ready reagent) and eosin (Merck) (H&E), 4’,6-diamidino-2-phenylindole (DAPI; Sigma), Alcian blue (Atom Scientific), and Sirius red (WVR International)/Millers elastin stains (Raymond A Lamb).

For immunohistochemistry, sections were taken at 6 μm. The Ultra Vision system-HRP kit (Thermo Scientific) was used for antibody detection. Sections were analyzed using monoclonal antibodies to collagen I, collagen II, collagen III (all at 1:50 dilution; Chemicon), and tenasin-C (1:100; Leica Microsystems) and polyclonal antibodies to galactose-α,1,3-galactose (α-Gal; 0.37 mg/mL; Enzo Life Sciences). Omission of the primary antibody and isotype control antibodies (mouse IgG1; rabbit IgM) were used as controls. Sections of native and decellularized SFTs were also treated with α-galactosidase for 20–24 h at 37°C (5 U/mL; Sigma) to remove the α-Gal epitope from the tissue as an additional negative control for determination of α-Gal. Sections were blocked with dual endogenous enzyme block (Ultra Vision). Antibodies were diluted in Tris-buffered saline (TBS) containing 0.1% (w/v) bovine serum albumin (Sigma). The sections were incubated with the diluted antibodies, isotype controls, or antibody diluent only for 1 h, TBS and TBS containing 0.05% (v/v) Tween (Sigma) were used as wash buffers throughout. The sections were labeled with polymer horseradish peroxidase (HRP), followed by DAB chromagen from the Ultra Vision kit.

The presence of the α-Gal epitope in tissue sections was also analyzed using biotinylated GSL-1 lectin (Vector Labs). Sections were blocked with streptavidin/biotin and carbo-free blocking solution (Vector Labs). Lectin was used at 5 mg/mL in TBS. The streptavidin polymer and IMMpact DAB chromagen (Vector Labs) were used to visualize lectin.
binding. Lectin (5 mg/mL) absorbed with 200 mM galactose, omission of the lectin, and \(\alpha\)-galactosidase-treated sections were used as negative controls. All sections were viewed using an Olympus BX51 microscope and imaged using an Olympus XC50 digital camera controlled through Cell B software (Olympus, United Kingdom).

**Determination of total DNA content of native and decellularized SFTs**

The total DNA content of six decellularized SFTs was determined and compared with the DNA content of native tissue. Tissue samples were taken from the toe, middle, and ankle regions of each tendon, combined, and macerated using a scalpel blade. The samples were weighed (circa 25 mg fresh tissue and 250 mg decellularized tissue) and freeze-dried to a constant weight. The DNA was extracted using a DNeasy blood and tissue kit (Qiagen) following the manufacturer’s instructions. The extracted DNA was then quantified at 260 nm using a nanodrop spectrophotometer (ND-500 Labtech).

**Determination of collagen content of decellularized SFTs**

The total collagen content of native \((n = 6)\) and decellularized \((n = 6)\) SFTs was determined using the hydroxyproline assay. Samples of native and decellularized tendons were freeze-dried to a constant weight, hydrolyzed in 6 M HCl for 4 h at 120°C, and neutralized with 6 M NaOH. Test samples (50 \(\mu\)L) were added to 96-well plates with 100 \(\mu\)L of chloramine-T solution (chloramine-T hydrate; Sigma) for 5 min with agitation. Ehrlich’s reagent (100 \(\mu\)L; Sigma) was added to each well and the wells covered and incubated at 60°C for 45 min. The absorbance was read at 570 nm. The hydroxyproline content of test samples was determined by interpolation from a standard curve of trans-4-hydroxy-L-proline (Sigma).

**Determination of the glycosaminoglycan content of decellularized SFTs**

The glycosaminoglycan (GAG) content of native \((n = 6)\) and decellularized \((n = 6)\) SFTs was determined using the dimethyl blue assay. Native and decellularized tendon samples were freeze-dried to a constant weight before being digested in papain (1250 units [Sigma] in 25 mL of 5 mM L-cysteine hydrochloride plus 5 mM Na2EDTA [WVR International]) for 26 h at 60°C with agitation. Standard or test solutions (40 \(\mu\)L) were added to wells of 96-well plates with 250 \(\mu\)L 1,9-dimethyl blue solution (DMB). Absorbance was read at 525 nm after 2 min of incubation. The sulfated sugar content of the test samples was determined by interpolation from the standard curve of chondroitin sulfate B (\(\mu\)g/mL; Sigma) versus OD 525 nm.

**Semiquantitative analysis of \(\alpha\)-Gal content of decellularized SFT**

Tissue samples (100 mg) from five different native and decellularized SFTs were macerated and added to individual 1.5-mL microcentrifuge tubes. \(\alpha\)-Galactosidase (5 U/mL in 50 mM Tris pH 6.0) was added to the samples of native and decellularized tissue and rotated at 37°C for 24 h (negative control tissue). Native, decellularized, and negative control tissues were placed into individual tubes and purified monoclonal antibody to \(\alpha\)-Gal diluted to 0.37 mg/mL in TBS was added to all samples. Antibody only (no tissue) was included as a no tissue control. The mixtures were rotated at 4°C for 16 h. Following incubation, the samples were centrifuged at 600 g for 15 min and the supernatant (100 \(\mu\)L) was added to the wells of MaxiSorp plates (Nunc) that had been coated with 50 \(\mu\)L \(\alpha\)-Gal-BSA (10 \(\mu\)g/mL in PBS; Dextra) overnight at 4°C and incubated for 3 h at RT. Following this, 50 \(\mu\)L secondary rabbit anti-mouse HRP antibody (1:1000; Dako) was added to the wells and incubated for 1 h at RT. The plates were washed with TBS before adding 100 \(\mu\)L \(\alpha\)-phenylenediamine dihydrochloride (OPD) solution (Sigma) and incubating for 10 min at RT in the dark. The reaction was stopped with 50 \(\mu\)L 3 M sulfuric acid and the absorbance measured at 490 nm with a reference filter of 630 nm.

**Determination of the biocompatibility of decellularized SFT tissue**

The *in vitro* biocompatibility of decellularized SFTs \((n = 6)\) was determined using contact and extract biocompatibility assays.

**Cell culture.** Murine 3T3 fibroblastic cells (European Collection of Cell Cultures; ECCAC) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) plus 10% (v/v) fetal calf serum (FCS), 2 mM L-glutamine, and 100 U/mL penicillin/streptomycin (all from Lonza). Baby hamster kidney (BHK; ECCAC) cells were cultured in Glasgow’s minimum essential medium plus 5% (v/v) FCS, 10% (v/v) tryptose phosphate broth (29.5 g/L; Sigma), and 100 U/mL penicillin/streptomycin. The cells were cultured at 37°C in 5% (v/v) CO2 in air in a humidified atmosphere.

**Contact biocompatibility assay.** Samples from the toe, middle, and ankle regions of decellularized SFTs were attached to the center of wells of a six-well plate (Nunc) using Steri-Strips (3 M; Medisave). Cyanoacrylate (RS Systems) was used as the positive control and Steri-Strip alone as the negative control. BHK and 3T3 cells were seeded at 5 \times 10^5 cells in 2 mL of appropriate culture medium into the sample and control wells and cultured for 48 h at 37°C in 5% (v/v) CO2 in air in a humidified atmosphere. The cell monolayers were washed with PBS, fixed with 10% (v/v) NBF, and stained with Giemsa solution (WVR International) before being washed with tap water and allowed to dry. The plates were then imaged by light microscopy.

**Extract cytotoxicity assay.** Samples from each region of the decellularized SFTs (toe, middle, and ankle) were macerated and incubated in DMEM (300 mg tissue in 3 mL DMEM) for 72 h at 37°C with agitation. BHK \((1 \times 10^5)\) or 3T3 \((2.5 \times 10^5)\) cells in 200 \(\mu\)L of appropriate culture medium were seeded into the wells of 96-well plates (Nunc) and cultured for 24 h at 37°C in 5% (v/v) CO2 in air. The culture medium was aspirated from the wells and 100 \(\mu\)L of the tissue extract plus 100 \(\mu\)L of appropriate culture medium (with double strength supplements) was added. The positive control was 40% (v/v) dimethyl sulfoxide (DMSO; Sigma) in culture medium (200 \(\mu\)L) and cells cultured in medium alone served as negative controls for cytotoxicity. The cells
were cultured for 48 h before assessing cell viability by measuring cellular ATP. ATP was measured using the ATPLite™ assay kit (PerkinElmer) following the manufacturer’s instructions. The luminescence (counts per second) in each well was measured using a TopCount (C9902 PerkinElmer) luminescence plate reader.

In vivo biocompatibility of decellularized SFT tissue

The host response to fresh (n = 3) and decellularized (n = 6) SFTs was assessed in a mouse subcutaneous implant model. Female C3H mice, 6–12 weeks old (Harlan), were anesthetized, the mid-dorsum of the mice was shaved and aseptically prepared and a small incision was made in the dorsal skin. Tissue samples (3 mm³) taken from the central region of the midportion of each tendon were implanted subcutaneously along the mid-dorsal line distant from the skin incision. Mice were fed food and water ad libitum and left for a period of 3 months. Mice were sacrificed according to schedule (1) and the sample and surrounding skin harvested for analysis. All animal procedures were carried out in accordance with UK Home Office regulations and with appropriate Home Office licenses.

The explanted samples were fixed in NBF for 48 h before being processed and embedded in paraffin wax. Sections were taken at 6 μm, retaining one section in every 200 throughout the full thickness of the tissues. Sections were stained with H&E and viewed as described above.

Tensile testing of native and decellularized SFTs

Six native and acellular SFTs were snap-frozen by immersion in dry ice and processed into dumbbell shapes of consistent cross-sectional area of 3.5 × 5 mm and gauge length of 30 mm. Specimens were then wrapped in PBS-soaked paper and allowed to thaw and equilibrate at RT for at least 2 h before biomechanical tensile testing.

Specimens were mounted through bespoke cryogrips to an Instron 3366 (Instron, Bucks, United Kingdom) material testing machine equipped with a 1 kN load cell. The grips were manufactured with a void into which dry ice was placed to reduce the primary gripping surface to subzero temperatures. Once secured in the cryogrips, specimens were tensioned to a preload of 0.5 N to ensure tautness was achieved. This load was subsequently tared and tensile testing performed. This consisted of 10× preconditioning cycles between 0% and 5% strain at a rate of 15 mm/min, followed by an extension ramp to failure at a rate of 30 mm/min. Load and extension data were recorded at a frequency of 10 Hz, from which the ultimate tensile strength (UTS), failure strain (FAIL), and Young’s modulus of the collagen phase (ECOLL) of the specimens were calculated.

Analysis of data

Statistical analysis was performed using Student’s t-test (native vs. acellular tissue) or one-way analysis of variance (ANOVA), followed by the T-test, to determine minimum significant differences (MSD) between more than two group means. A p-value of less than 0.05 was deemed to be significant.

Results

Gross structure of the porcine SFT

The SFTs harvested from pigs (n = 6) had a length of 19.7 ± 1.1 cm (±95% confidence limits [CL]), a width of 11.2 ± 1.3 mm, and a depth of 3.9 ± 0.28 mm. Images of the porcine SFT before and after decellularization are shown in Figure 1.

Histological and immunohistochemical analysis of native and decellularized SFT

Sections of native tendons stained with H&E and DAPI (Fig. 2A, C) revealed the presence of cells neatly aligned along the crimp of the collagenous tissue matrix. Blood vessels were apparent in the endotenon regions where there were large aggregations of cells. Sections stained with Sirius red/Millers viewed under polarized light (Fig. 2E) showed the aligned collagen fibers and also demonstrated the crimp formation of the fibers. Under bright-field illumination, Millers stain did not reveal the presence of any elastin in the tissue sections when the stain was combined with Sirius red. Millers staining alone showed that there was elastin present in the endotenon regions surrounding blood vessels. Alcian blue-stained tissue sections (Fig. 2G) showed GAGs distributed around the vascularized endotenon regions; however, the GAGs were more concentrated around the vascularized endotenon regions.

Sections of decellularized tendons stained with H&E (Fig. 2B) and DAPI (Fig. 2D) revealed the absence of any cell nuclei or double-stranded DNA. The histoarchitecture of the matrix appeared similar to native tissue with evidence of some loss of collagen fiber crimp. Sections stained with Sirius red/Millers (Fig. 2F) demonstrated the retention of the collagen fiber alignment. Alcian blue-stained sections (Fig. 2H) showed a decrease in the GAG content throughout the decellularized tissue compared with the native tissue. Scanned images of the decellularized porcine SFT stained with
Native tendon, positive control tissues, and isotype controls allowed verification of the specificity of antibodies used for detection of collagens I, II, III, and tenascin-C. The native tendon sections stained strongly for both collagens I (Fig. 3A) and III (Fig. 3C) throughout the tissue. Tenascin-C stained positively mainly around the endotenon regions (Fig. 3E). There was no positive staining for collagen II.

Sections of decellularized SFT showed similar patterns of staining, with no apparent difference in collagen I (Fig. 3B), collagen III (Fig. 3D), or tenascin-C (Fig. 3F) staining. Labeling of the α-Gal epitope in the tissue sections using both GSL-1 lectin and monoclonal antibody to the α-Gal epitope showed very strong staining in the endotenon regions of the native tendon. Sections of the decellularized tissue also showed positive staining with both the monoclonal antibody to α-Gal and the GSL-1 lectin, although this was reduced compared with native tissues. Controls of α-galactosidase-treated tissue sections and use of GSL-lectin absorbed with galactose confirmed the specificity of lectin to α-Gal. Images of sections stained with the GSL-1 lectin and controls are shown in Figure 4. This indicated that α-Gal was still present in the tissue following decellularization.

FIG. 2. Images of histological sections of native (A, C, E, G) and decellularized (B, D, F, H) porcine SFTs. Stained with hematoxylin and eosin (H&E; A, B) showing an absence of cells postdecellularization. Stained with 4,6-diamidino-2-phenylindole (DAPI) showing the blue-stained nuclei in (C), but not (D). Stained with Sirius red/Millers (E, F) and captured under polarized light showing the crimped nature of the collagenous matrix. Stained with Alcian blue (G, H) with blue-stained glycosaminoglycans (GAGs) mainly in the endotenon (G), but less prevalent in (H). All images captured at 10× magnification. Scale bars 200 μm.
Total DNA content of native and decellularized SFTs

Native porcine SFT had a mean total DNA content of 303 ng/mg per tissue dry weight. Following decellularization, the mean total DNA content of the tissue was 13.4 ± 2.2 (95% CL) ng/mg per tissue dry weight tissue (p < 0.0001; Student’s t-test).

In vitro biocompatibility of decellularized SFT

Contact cytotoxicity assays. BHK and 3T3 cells grew to and in contact with the decellularized tissue isolated from all regions (toe, middle, and ankle) of the tendon (representative images in Fig. 5). The cells did not show any change in morphology compared with the cells cultured alone or with Steri-Strip (negative controls). Cyanoacrylate glue (positive control) was highly toxic to the cells, resulting in cell lysis.

Extract cytotoxicity assays. Murine 3T3 cells cultured with tissue extracts from each of the six tendons showed no significant differences in ATP levels compared with the negative control (cells grown in culture medium; Fig. 5). The BHK cells cultured with the majority of tissue extracts did not show any significant difference in ATP levels compared with the negative control; however, there was a small but significant decrease in ATP levels in BHK cell culture in the presence of 4/18 extracts (Fig. 6).

Collagen and GAG content of native and decellularized SFTs

Native SFT had a hydroxyproline content of 138 ± 21 μg/mg (mean [n=6]± 95% CL) of dry weight, indicating that the extracellular matrix of the SFT mainly comprised collagen. Following decellularization, the SFT had a mean hydroxyproline content of 163 ± 21 μg/mg, which was not significantly different to the native tissue (Student’s t-test). The total sulfated sugar content of the native SFT was very low at 6.85 ± 1.7 μg/mg (mean [n=6]± 95% CL). Following decellularization, the total sulfated sugar content was reduced significantly to 1.79 ± 1.48 μg/mg (p < 0.001; Student’s t-test).

Semiquantitative determination of the α-Gal epitope in native and decellularized SFTs

Following adsorption of antibody to α-Gal present in samples of native, decellularized, and α-galactosidase-treated tissues, the relative levels of nonadsorbed antibody were determined by ELISA using α-Gal-BSA (Fig. 7). There was no significant difference in the amount of anti-α-Gal
antibody in the supernatant from the no tissue control and the α-galactosidase-treated native and decellularized tissues, indicating that α-galactosidase treatment effectively removed the α-Gal epitope from the tissues. The antibody levels in the supernatants from the native and decellularized tissues were significantly ($p < 0.05; \text{ANOVA}$) lower than the no tissue control and the α-galactosidase-treated native and decellularized tissues (indicated by the lower absorption values). The decellularized SFT tissue absorbed fewer antibodies than the native SFT tissue, indicating that the decellularization process had removed some α-Gal; however, the difference was not significant.

**FIG. 4.** Lectin binding in sections of native and decellularized porcine SFTs. (A) Native SFT stained with GSL-1 lectin, showing positive binding of lectin mainly in the endotenon regions; (B) native SFT treated with α-galactosidase to remove α-Gal and stained with GSL-1 lectin, showing an absence of binding of lectin following the enzyme treatment; (C) native SFT stained with GSL-1 lectin that had been absorbed with galactose, showing an absence of binding of lectin indicating the specificity of lectin to galactose moieties in the tissue; (D) decellularized SFT stained with GSL-1 lectin, showing reduced, but positive, binding of lectin mainly in the endotenon regions; (E) decellularized SFT stained with GSL-1 lectin that had been absorbed with galactose, showing an absence of binding of lectin indicating the specificity of the lectin to galactose moieties in the decellularized tissue; and (F) decellularized SFT nonlectin control treated with streptavidin polymer and chromagen only, showing an absence of background staining due to lectin detection reagents.

**FIG. 5.** Images of contact cytotoxicity assays to determine the biocompatibility of decellularized porcine SFT. *Top row:* baby hamster kidney (BHK) cells cultured in the presence of samples of decellularized SFT (A, B), Steri-Strip alone, negative control (C), and cyanoacrylate, glue positive control (D). **Bottom row:** 3T3 cells cultured in the presence of samples of decellularized SFT (E, F), Steri-Strip alone, negative control (G), and cyanoacrylate glue, positive control (H). The cells have been stained with Giemsa. The images show that both cell types grew up to and in contact with the decellularized SFT with no change in morphology compared with the negative control (Steri-Strip), while the positive control caused cell lysis. Images captured at 10× magnification. Scale bars 200 μm.
Biocompatibility of decellularized SFT in vivo

The native and decellularized tissue samples were explanted from mice and analyzed by histology. Sections of native tissue stained with H&E revealed a fibrous capsule around the tissue with little evidence of cellular infiltration into the depth of the tissue. Moreover, the native tissue explants were calcified and degenerating in the central regions indicated by dark staining (Fig. 8A, B). Histological sections of explanted decellularized SFT samples revealed minimal capsule formation with evidence of cellular infiltration with cells of a fibroblastic and macrophage morphology into the depth of tissues and an absence of calcification (Fig. 8C, D). Scanned images of the tissues explanted from mice at 12 weeks are provided in Supplementary Figure S2.

Material properties of native and decellularized SFTs

When subjected to uniaxial tensile testing to failure, midsubstance failure of all native and decellularized tissues was achieved. There were no significant differences in the UTS, failure strain, or Young’s modulus of the collagen phase of the native compared with decellularized tissue specimens (Student’s t-tests; Table 1).

Discussion

The porcine SFT was chosen as the source tissue for the development of a decellularized biological scaffold for ACL replacement because it could be routinely harvested in dimensions that are currently used for ACL replacement using hamstring tendon autografts. Grafts greater than 7 mm in diameter are preferred with a length of at least 9 cm for a single-stranded graft or at least 18 cm for a double-bundle graft and the porcine SFT met these specifications. The aim of this study to develop and evaluate a bioprocess for removal of immunogenic cells and cellular remnants from the porcine SFT that has minimal effect on the extracellular matrix and mechanical properties of the tissue was achieved.

The residual total DNA of the decellularized SFT was circa 13 ng/mg dry weight of tissue. This was well below the levels of double-stranded DNA of 50 ng/mg reported to be indicative of a decellularized tissue that would undergo constructive tissue remodeling in vivo. Immunohistochemical analysis showed retention of structural collagens and tenascin-C post decellularization. Tenascin-C is an extracellular matrix glycoprotein found in all musculoskeletal tissues. It is important in regulating cellular differentiation and proliferation and is believed to have a role in the healing process following musculoskeletal injury. Proteoglycans with associated GAGs account for less than 1% of tendon dry weight. The native SFT was found to contain circa 0.7% GAGs, which were located mainly in the endotenon regions. GAG depletion following decellularization was confirmed by quantitative analysis. The reduction of GAG content was not unexpected since previous studies have shown that decellularization of tissues using SDS leads to a loss of GAGs.

FIG. 6. ATP levels in 3T3 (A) and BHK (B) cells cultured in the presence of extracts of decellularized SFT. Data are presented as the mean (n = 3) counts per second × 10⁻⁶ ± 95% confidence limits following ATP Lite™ assay of cells following 48 h of culture in the presence of decellularized tissue extracts. One to six indicate different decellularized SFTs; A (ankle), M (mid-tendon), and T (toe) region of the tissue sample used to prepare the extract. Con represents cells cultured in the absence of extract with culture medium alone and dimethyl sulfoxide (DMSO) the positive control for toxicity. Data for each cell line were analyzed by one-way analysis of variance (ANOVA), followed by the T-test to determine any significant differences between group means. * Indicates significantly lower than the control (Con; p < 0.05).

FIG. 7. Semiquantitative evaluation of α-Gal in native and decellularized SFTs. Absorption at 490–630 (reference) in ELISA for detection of antibodies to α-Gal following adsorption with native SFT (NSFT), native SFT treated with α-galactosidase (aNSFT), decellularized SFT (DSFT), and α-galactosidase-treated decellularized SFT (aDSFT). NTC is the no tissue control (one group for each ELISA plate). Data are presented as the means (n = 5) ± 95% confidence intervals. Data were analyzed by one-way ANOVA, followed by determination of the minimum significant differences using the T-method. * Indicates significantly lower (p < 0.05) than the respective NTC and α-galactosidase-treated tissue.
The α-Gal epitope is present on cell membrane glycolipids and glycoproteins of lower mammals, but is not expressed in humans. Humans naturally produce antibodies, which recognize the α-Gal epitope, and these antibodies have a role to play in the hyperacute rejection of vascularized whole-organ xenografts. It was therefore of interest to determine the presence of the α-Gal epitope in the native and decellularized SFTs. The α-Gal epitope was mainly located in the endotenon regions of the native SFT, consistent with reported localization to the basement membrane. Qualitative and semiquantitative analysis of α-Gal showed that although the levels of α-Gal were reduced in the decellularized SFT compared with native porcine tissue, the epitope was still present. Whether the presence of residual α-Gal in acellular connective tissue biological scaffolds is of concern or indeed an attribute for future clinical translation requires discussion.

It has been well documented that acellular biological scaffolds derived from porcine tissues (such as the porcine intestinal submucosa) that have been used clinically in millions of patients stimulate a constructive remodeling response orchestrated by macrophages that have a regenerative wound healing phenotype (M2; macrophages; Brown et al. 28). It is also recognized that these biological scaffolds retain the α-Gal epitope. Recently, studies conducted in 1,3-galactosyltransferase knockout (GTKO) mice (which lack the α-Gal epitope and can produce anti α-Gal antibodies) have shown that application of α-Gal liposomes on wounds accelerated healing compared with control wounds by rapid recruitment and activation of macrophages. Moreover, scar formation was reduced. Similar studies in GTKO pigs have led to the concept that anti-Gal/α-Gal nanoparticles may have utility in tissue regeneration and prevention of fibrosis. Hence, although the expression of α-Gal by cells in xenogeneic organs and tissues may lead to antibody-dependent complement activation, cell lysis, and a proinflammatory response when transplanted in humans, the presence of the α-Gal epitope in the absence of cells in decellularized biological scaffolds may indeed contribute to rapid recruitment of macrophages, which are polarized to the wound-healing M2 phenotype, inducing tissue regeneration.

The decellularized SFT showed good biocompatibility in in vitro assays. The decellularized SFT was therefore tested for biocompatibility in a subcutaneous implant model in mice in comparison with native porcine SFT. The native

| Material Properties of Native and Decellularized Superflexor Tendons |
|--------------------------|--------------------------|--------------------------|
|                          | UTS (MPa)                | εFAIL (mm/mm)            | ECOLL (MPa)               |
| Native                   | 52.5 ± 5.9               | 0.33 ± 0.05              | 234.2 ± 51.3              |
| Acellular                | 61.8 ± 10.3              | 0.29 ± 0.04              | 294.1 ± 61.9              |
| p                        | 0.07                     | 0.15                     | 0.08                      |

The material properties were determined by uniaxial tensile testing. Data are presented as the means (n = 6) ± 95% confidence limits. The data for native and decellularized SFT samples were compared by Student’s t-test, which showed no significant differences between the two groups.

UTS, ultimate tensile strength.
tissue stimulated a foreign body-type response and the cellular tissue was heavily calcified at the center and showed signs of degeneration, most likely due to the presence of dead cells within the tissue. In contrast, the decellularized SFT tissue was well accepted in the mouse model with no evidence of an overt inflammatory or specific immune response; the tissue was not degenerated and calcified. Cells of a fibroblastic and macrophage morphology were observed penetrating the implanted decellularized tissue.

With regard to the potential of the decellularized porcine SFT for future clinical translation as a device for ACL reconstruction, the biomechanical properties of the scaffold are of utmost consideration. The UTS of the natural human ACL has been reported to be circa 36 MPa, while the native porcine SFT was found to have a UTS of 52.5 MPa and, following decellularization, the UTS was not significantly different (62 MPa). Hence, the strength of the decellularized porcine SFT is well within the range needed to meet surgical requirements. The porcine SFT also has similar dimensions to the human hamstring tendon, which is commonly used for ACL reconstruction.

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Disclosure Statement

E.I. and J.F. are shareholders in and consultants to Tissue Regenix. E.I. and J.F. are consultants to DePuy Synthes. J.F. is consultant to Invibio and Simulation Solutions. All other authors have no competing financial interests.

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